

This is the author's manuscript



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Resource driven community dynamics of NH4+ assimilating and N2O reducing archaea in a temperate paddy soil

Original Citation:	
Availability:	
This version is available http://hdl.handle.net/2318/1633796	since 2017-08-09T18:05:56Z
Published version:	
DOI:10.1016/j.pedobi.2017.02.001	
Terms of use:	
Open Access Anyone can freely access the full text of works made available as under a Creative Commons license can be used according to the tof all other works requires consent of the right holder (author or protection by the applicable law.	erms and conditions of said license. Use

(Article begins on next page)





This Accepted Author Manuscript (AAM) is copyrighted and published by Elsevier. It is posted here by agreement between Elsevier and the University of Turin. Changes resulting from the publishing process - such as editing, corrections, structural formatting, and other quality control mechanisms - may not be reflected in this version of the text. The definitive version of the text was subsequently published in PEDOBIOLOGIA, 62, 2017, 10.1016/j.pedobi.2017.02.001.

You may download, copy and otherwise use the AAM for non-commercial purposes provided that your license is limited by the following restrictions:

- (1) You may use this AAM for non-commercial purposes only under the terms of the CC-BY-NC-ND license.
- (2) The integrity of the work and identification of the author, copyright owner, and publisher must be preserved in any copy.
- (3) You must attribute this AAM in the following format: Creative Commons BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/deed.en), 10.1016/j.pedobi.2017.02.001

The publisher's version is available at: http://linkinghub.elsevier.com/retrieve/pii/S0031405616300762

When citing, please refer to the published version.

Link to this full text: http://hdl.handle.net/2318/1633796

This full text was downloaded from iris - AperTO: https://iris.unito.it/

Type of article: original article Resource driven community dynamics of NH₄⁺ assimilating and N₂O reducing archaea in a temperate paddy soil Maria Alexandra Cucu^{1,2,3*}, Sven Marhan², Daniel Said-Pullicino¹, Luisella Celi¹, Ellen Kandeler² and Frank Rasche³ ¹Rice Agro-ecosystem and Environmental Research Group, Department of Agricultural, Forest and Food Sciences, University of Turin, Italy ²Institute of Soil Science and Land Evaluation, Soil Biology Section, University of Hohenheim, Stuttgart, Germany ³Institute of Plant Production and Agroecology in the Tropics and Subtropics, University of Hohenheim, Stuttgart, Germany Running title: Archaeal N cycling in temperate paddy soils *Corresponding author: Dr. Maria Alexandra Cucu Phone: +39 011 670 8544 Fax: +39 011 670 8692 Email: mariaalexandra.cucu@unito.it

Abstract.

In fertilized paddy soils, the role of resource availability on ammonium (NH₄⁺) assimilation and immobilization by archaea requires advanced understanding as this may have considerable implications on subsequent catalytic steps in the soil N cycle including archaeal nitrous oxide (N₂O) reduction. To gain a deeper understanding about these process links, we incubated a temperate paddy soil under submerged conditions with or without straw and fertilized with either ¹⁵N-enriched (99 atom% ¹⁵N) or non-enriched (NH₄)₂SO₄. Notably, a variation in community structure and a higher abundance of archaeal N₂O reductase (*arc-nosZ*) genes in the no straw treatment than in the straw one was observed. This was attributed to NH₄⁺ assimilation by N₂O reducing archaea as was further corroborated by a considerable ¹⁵N-enrichment of archaeal glutamate dehydrogenase (*gdhA*) genes. Moreover, indications were found that denitrifying archaea controlled their chemotrophic and heterotrophic metabolisms in response to different availabilities of inorganic and organic N and C resources. However, in the presence of straw, bacterial *nosZ* genes may have also contributed to the completion of denitrification. Our results suggested that N assimilation contributed to the last step of archaeal denitrification. Therefore, archaea may play a key role in regulating major N fluxes in fertilized paddy soils, especially in the absence of rice straw.

- **Keywords:** arc-nosZ gene; total prokaryotic nosZ clade I gene; gdhA gene; ¹⁵N-DNA-based stable
- 45 isotope probing.

1. Introduction

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

Fertilized rice paddy soils generally exhibit lower nitrogen (N) recovery rates (30-40%) with respect to upland soils (Cassman et al., 2002). Incorporation of crop residues such as rice straw generally leads to N immobilization (assimilation to sustain their own metabolism) which reduces fertilizer use efficiency to a large extent (Reddy 1982; Eagle et al., 2000; Bird et al., 2001). In previous studies we observed that addition of rice straw induced, in comparison to no straw addition, an 80% increase of immobilization of applied ammonium (NH₄⁺) along with a 65% decrease in gaseous N losses (Cucu et al., 2014; Said-Pullicino et al., 2014). On the other hand, incorporation of rice straw in paddies was shown to stimulate N gaseous losses (i.e., N oxide gases and N₂) from soil-plant systems since up to 70% of applied N may be lost through coupled nitrification-denitrification (Cassman et al., 1998; Ghosh and Bhat, 1998; Majumdar, 2013). Significant amounts of nitrous oxide (N₂O), an important greenhouse gas, have been shown to be emitted under aerobic conditions (Zou et al., 2007; Ishii et al., 2011a). To mitigate this climate variation promoting process, it was acknowledged that in continuously submerged paddies complete denitrification via microbial nitrous oxide reduction can act as a considerable sink for N2O which favors N₂ losses (Davidson et al., 1986; Conrad 1995, 1996; Ferrè et al., 2012). Both N assimilation and denitrification are mediated by specific microbial communities which are well adapted to the anaerobic conditions of paddy environments (Liesack et al., 2000; Kögel-Knabner et al., 2010). These ecosystems are characterized by a dynamic microbial community in which archaea may predominate over bacteria (Cabello et al., 2004, Rush, 2016), and may also exhibit different metabolic pathways and resource utilization. Denitrification sensu latu, as the pathway of nitrate (NO₃-) respiration, can be associated with dissimilatory and assimilatory branches of microbial metabolism (Zumft, 1997). Under anaerobic conditions, bacterial denitrifiers typically adopt a dissimilatory metabolism for their energy gain (Zumft, 1992), while their potential assimilatory NO₃- reduction pathway is suppressed when NH₄+ is the predominant inorganic N form (Rice and Tiedje, 1989; Cabello et al., 2004).

Archaeal denitrifiers can simultaneously perform dissimilatory and assimilatory reactions (Rusch, 2013) by using N not only for energy but also electron sink and detoxification (Zumft, 1997). The gained N is also utilized to build up nitrogenous compounds (e.g., DNA, amino sugars, proteins) (Devêvre and Horwáth, 2001; Nannipieri and Paul, 2009), which ultimately contribute to the soil organic N pool (Kögel-Knabner et al., 2010). However, assimilatory NO₃- reduction is less frequent than respiratory NO₃- reduction in the archaeal domain (Martínez-Espinosa et al., 2001). Instead, Cabello et al. (2004) highlighted the importance of the archaeal NH₄⁺ assimilation pathway catalyzed by glutamate dehydrogenase (GDH). Encoded by the gdhA gene, GDH generally shows a high activity under non-limiting NH₄⁺ soil conditions (Bonete et al., 2008). This may be the case for fertilized rice agro-ecosystems characterized by high NH₄⁺ availability as a result of mineral N addition and mineralization of organic matter (OM) including freshly applied rice straw (Sahrawat, 2004; Cucu et al., 2014). It needs to be pointed out that NH₄⁺ assimilation by archaea in paddy soils requires explicit understanding as this may have considerable implications on subsequent catalytic steps in the soil N cycle including the last step of denitrification (i.e., N₂O reduction). Although N fertilization is recognized as the primary factor influencing denitrification under reduced conditions (Garcia and Tiedje 1982; Nogales et al., 2002; Prieme et al., 2002), the role of rice straw and the interactive effects of available N and C resources on archaeal $\mathrm{NH_4^+}$ assimilation and $\mathrm{N_2O}$ reduction remains uncertain. Compared to bacterial denitrification (Philippot et al., 2002; Henry et al., 2006; Ishii et al., 2011b), only limited knowledge exists about this process in archaea. Likewise, only a few archaeal denitrification genes and related enzymes have been investigated so far (de Vries and Schroder, 2002; Cabello et al., 2004). Moreover, information on the reduction of N2O to N2 is only available for very few hypertermophilic and halophilic archaeal species including Haloferax denitrificans and Pyrobaculum aerophilum (Tomlinson et al., 1986; de Vries and Schroder, 2002). Jones et al. (2013) assigned several archaeal species to nosZ groups I and II covering the bacterial and archaeal domains. Recently, Rusch (2013) has introduced a complementary marker for the arc-nosZ gene

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

that targets the archaeal N₂O reducing community. The *arc-nosZ* gene is part of the total prokaryotic *nosZ* gene clades I and II. Although a phylogenetic bias based on hypertermophilic and halophilic archaeal sequences has to be taken into account due to limited database information, Rusch (2013) considered the *arc-nosZ* gene marker suitable for the full range of applications targeting archaeal denitrifiers. While the crucial role of archaeal species in methanogenesis and methanotrophy in rice paddies (Fazli et al 2013) is well known, their involvement in denitrification remain less understood.

The primary objective of this study was to provide advanced understanding about the role of archaea involved in N assimilation coupled with the final step of denitrification (i.e., N₂O reduction) as a function of C and N resource availability in a temperate paddy soil with no NH₄⁺ limitation. To achieve this goal, we tested the following hypotheses: i) archaeal N assimilation is controlled by the availability of labile organic C substrates (i.e., rice straw addition); ii) a fraction of the assimilated N is used for proliferation of archaeal community members harboring the *arc-nosZ* gene; iii) in absence of rice straw, the relative contribution of archaea to denitrification is more

2. Materials and methods

important than that of their bacterial counterparts.

115 2.1. Incubation experiment

Soil was collected from the Ap horizon (0-15 cm) of a long-term field experiment in Vercelli, Northwest Italy (45°17′47″N, 8°25′51″E) in February 2012 before rice straw incorporation. The paddy soil, classified as a Haplic Gleysol (WRB, 2007), has been under continuous single-cropped rice cultivation for the last 30 years. The soil had a pH around 6, a sandy loam texture (7% clay, 41% silt, 52% sand) and relatively low contents of organic C and total N (11.6 and 1.1 g kg⁻¹, respectively). The amount of clay in conjunction with soil organic matter content accounted for the low cation exchange capacity (CEC = 6.7 cmol (+) kg⁻¹). Further details on soil properties have been previously reported in Cucu et al. (2014) and Said-Pullicino et al. (2014). After manual removal of

124 remaining vegetal residues, field moist soil was homogenized after passing through a 2 mm sieve.

Rice straw (Oryza sativa L. cv. Sirio CL) was sampled from the same experimental site in October

2011 after grain harvest, dried at 40°C and cut into 1 cm segments. Total C and N contents of the

rice straw were 400 and 6.6 g kg⁻¹ respectively, resulting in a C/N ratio of 61. A laboratory incubation experiment was carried out adopting a setup similar to that described by Cucu et al. (2014). The experimental design consisted of a completely randomized arrangement of soil microcosms incubated under submerged conditions with or without the addition of rice straw (application dose of 4.3 g straw kg⁻¹ soil, equivalent to 10 Mg dry weight ha⁻¹). After a preincubation period with or without straw addition for 14 days at 50% soil water holding capacity to reestablish the microbial equilibrium, inorganic N (application dose of 56 mg N kg⁻¹ soil, equivalent to 130 kg N ha⁻¹, generally applied in the field) was added to the soil samples and immediately submerged under 5 cm of deionized water. Inorganic N was added as ¹⁵N-labelled (NH₄)₂SO₄ (99 atom% ¹⁵N) to assess active N assimilatory prokaryotes by stable isotope probing (SIP, see below). A second set of soil samples was treated similarly with non-labeled (NH₄)₂SO₄ as control for SIP studies (Buckley et al., 2007b; Neufeld et al., 2007). The experiment was run with six replicates per treatment and time point. Incubation under submerged conditions was carried out for 60 days at 25°C in the dark. After 1, 5, 10, 20, 30 and 60 days from fertilizer application, soil microcosms were destructively sampled. The fresh soil sample was immediately used for DNA extraction as well as for inorganic N determination (see bellow). Submergence water was decanted, total volume

145

146

147

148

149

125

126

127

128

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

2.2. Chemical analyses

N₂O emissions were monitored daily during the first 4 weeks of incubation. On each measurement day, soil microcosm headspace was sampled twice: immediately and 30 min after closure of the lids. Afterwards the gas sample was transferred into pre-evacuated exetainers (5.9 ml, Labco Ltd.,

recorded, filtered at 0.45 µm and used for inorganic N determination. Afterwards, both soil and

water samples have been frozen at -20 °C for further analysis.

- Lampeter, UK), and analyzed for N₂O using a gas chromatograph equipped with an electron capture
- detector (Agilent 7890, Santa Clara, CA, USA). Three standard gases with known concentrations
- were used for gas flux calibration and calculation. N₂O fluxes were calculated by linear regression
- as described by Hutchinson and Mossier (1981).
- Soil pH was measured potentiometrically in H₂O. Submergence water samples as well as 0.5 M
- 155 K₂SO₄ soil extracts (1:4 w/v soil-to-extractant ratio) were analyzed for inorganic N (NH₄⁺, NO₃⁻)
- 156 (Auto-analyzer 3, Bran & Luebbe, Norderstedt, Germany), total dissolved nitrogen (TDN) and
- dissolved organic carbon (DOC) after sample acidification (TOC/TN analyzer Multi NC 2100S,
- 158 Analytic Jena GmbH, Jena, Germany).
- 159
- 160 2.3. Molecular analyses
- 161 2.3.1. DNA extraction
- 162 At each time point, total DNA was extracted from each fresh individual replicate soil sample using
- 163 the FastDNA® Spin kit for soil (MP Biomedicals, Solon, Ohio, USA) according to the
- manufacturer's instructions. DNA quantification was performed with a NanoDrop ND-2000
- 165 (NanoDrop Technologies, Wilmington, DE, USA).
- 166
- 167 2.3.2. Stable isotope probing
- 168 Measurement of ¹⁵N-enrichment in DNA extracts
- To measure the ¹⁵N-enrichment of soil DNA extracts according to España et al. (2011), a tin capsule
- 170 (Hekatech, Wegberg, Germany) was filled with 10 mg sorbsil (May & Baker, Dagenham, United
- Kingdom) plus 2 µg of the DNA extracts. Twenty µg N of dissolved unlabeled ammonium sulphate
- 172 (Roth, Karlsruhe, Germany) were then added to the capsule and dried at 50°C overnight. All DNA
- samples were analyzed with an elemental analyzer (Euro EA 3000; Hekatech) coupled with an
- isotope ratio mass spectrometer (DeltaPlus XP, Thermo Scientific, Waltham, USA). ¹⁵N-enrichment

175 of DNA extracts was subsequently calculated by taking into account the isotopic signature of the ammonium sulphate spike. 176 ¹⁵N-DNA-based stable isotope probing (SIP) was carried out on samples incubated for 30 days, 177 corresponding to distinct alteration of microbial activity (i.e., ¹⁵N enrichment of soil DNA extracts) 178 between "straw" and "no straw" treatments. To reach a high concentration and volume for the 179 180 further centrifugation, two replicates per DNA samples were pooled together. Finally, four DNA 181 samples were selected for SIP: two with highest ¹⁵N label (straw: 6.83 atom% ¹⁵N, no straw: 5.02 182 atom% ¹⁵N) and two respective ¹⁴N controls. Centrifugation mixtures were prepared by adding 5 mg of DNA diluted in 1 mL gradient buffer (0.1 M Tris, 0.1 M KCl, 1 mM EDTA) to 4.9 ml of a 183 184 7.163 M CsCl (Calbiochem/Merck, Darmstadt, Germany) solution. Isopycnic fractionation was 185 performed in 5.1 mL polyallomer centrifuge tube (13×51mm) placed in a Vti 65.2 vertical rotor 186 (both Beckman Coulter, Krefeld, Germany). Tubes were then centrifuged in an Optima™ L-90K ultracentrifuge (Beckman Coulter) was performed with 140,000 × g at 20°C for 69 h (España et al., 187 2011). After centrifugation, each gradient was fractionated into 16 individual fractions (312 µl 188 189 each) using a syringe pump (NE-1000, New Era Pump Systems, New York, NY, USA). Buoyant

195

196

197

198

199

200

190

191

192

193

194

2.3.3. Microbial abundance

Abundance of total bacterial and archaeal communities were done by quantifying the respective 16S rRNA genes (Rasche et al., 2011). Description of primer sets and amplification details used for quantitative PCR are specified in Table 1. Quantification of *nosZ* clade I was performed as described by Henry et al. (2006). The quantification of *arc-nosZ* genes was performed using

density was adjusted prior to centrifugation (AR200 digital refractometer, Reichert, New York,

USA) and calculated for each fraction according to Buckley et al. (2007b). SIP fractions were

precipitated with 20 µg of glycogen (Roche Diagnostics GmbH, Penzbeg, Germany) and 1 ml of a

30% polyethylene glycol 6000 solution (Carl Roth GmbH, Karlsruhe, Germany), washed (70%)

ethanol) and suspended in 30 µl of Tris-EDTA buffer (pH 8.0) (Neufeld et al., 2007).

primers published by Rusch, (2013). Primer accuracy was confirmed (Rusch, personal communication) and preliminary tests (data not shown) were carried out to assess the specificity of the primer set for the studied soil. To confirm the targeted archaeal NH₄⁺ assimilation process and the implied labeling success, primers targeting N assimilation in *Methanobrevibacter smithii* by means of the GDH system encoded by the gdhA gene were used. Methanobrevibacter sp. have been earlier found in paddy soils (Fetzer et al., 1993) and bioreactors (i.e., Methanobrevibacter related species), here with potential denitrifying abilities (Chuang et al., 2014) Abundance of bacterial and archaeal 16S rRNA genes and of the three functional genes (i.e., total prokaryotic nosZ clade I, arc-nosZ, archaeal gdhA) was determined by quantitative PCR (qPCR) using a StepOnePlusTM Real-Time PCR System (Applied Biosystems, Foster City, CA, USA) for bulk soil DNA as well as for gdhA genes in SIP fractions (Table 1). For standard preparation, amplicons from each target gene were generated, purified (Invisorb Fragment CleanUp, Stratec Molecular GmbH, Berlin, Germany), ligated into the Strata-Clone PCR cloning vector pSC-A (Strataclone PCR Cloning Kit, Agilent Technologies Inc.) and ligation products were transformed into StrataClone SoloPack competent cells (Stratagene). Specificity of clones used as quantitative PCR (qPCR) standards were checked via sequencing at LGC Genomics GmbH (Berlin, Germany) and BLAST analysis. Plasmid DNA was isolated (GenElute™ Plasmid Miniprep Kit, Sigma-Aldrich, St. Louis, MO, USA) from standard clones and quantified as described above. As assessed in preliminary tests and qPCR reaction optimization, each 25 µl qPCR cocktail contained 5 ng DNA (16S rRNA genes) or 10 ng DNA (total prokaryotic nosZ clade I and archaeal arc-nosZ gene), 1x Power SYBR green master mix (Applied Biosystems), 0.15 µM of each primer (Table 1), 0.25 μl of T4 gene 32 protein (500 μg ml⁻¹, MP Biomedicals). For gdhA gene, the 25 μl qPCR cocktail contained 1x FastStart Universal SYBR Green Master (ROX) (Roche), 0.2 μM of each primer (Table 1), 0.625 U Uracil-DNA Glycosylase (Roche) and the DNA template (10 ng from extracted soil DNA, 10 µl DNA from SIP fractions). Each sample was quantified in triplicate across plates, while standards in 10-fold serial dilutions were run in duplicate. The optimal dilution

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

of DNA extracts was tested to compensate any reaction inhibition by organic compounds (e.g., humic acids) co-extracted during DNA isolations (data not shown). Melting curves of amplicons were generated to ensure that fluorescence signals originated from specific amplicons and not from primer dimers or other artifacts. This was confirmed by checking the amplification products on 1% agarose gel. For quality control, melting curve analyses were performed. Amplification efficiency ranged from 80% (archaeal 16S rRNA, total prokaryotic nosZ clade I and arc-nosZ genes) to 103% (bacterial 16S rRNA and gdhA genes). The slopes were between -3.890 and -3.235 and $R^2 \ge 0.98$. Gene copy numbers were calculated with StepOneTM software version 2.2 (Applied Biosystems).

The data were normalized and presented in figures as copies g⁻¹ dry soil.

To determine the relative abundance of arc-nosZ gene with respect to the total prokaryotic nosZ gene clade I, the arc-nosZ to nosZ gene abundance ratio was calculated. Values > 0.5 indicate a higher relative abundance of archaea with respect to their bacterial counterparts within total prokaryotic nosZ clade I.

241 2.3.4. Microbial community structure

The community structure of archaeal 16S rRNA, total prokaryotic *nosZ* clade I, and archaeal *arc-nosZ* genes in bulk DNA soil, as well as in SIP fractions (total prokaryotic *nosZ* clade I and *arc-nosZ* genes) was studied by terminal restriction fragment length polymorphism (T-RFLP) analysis. Description of primer sets, PCR ingredients and amplification details used for T-RFLP analysis are specified in Table 2. Prior to digestion, all genes were amplified using PCR cocktails and amplification according to the details described in Table 2. For T-RFLP analysis, each forward primer was labeled with the fluorescent dye 6-FAM. Replicate amplicons were pooled, purified (Sephadex G-50, GE Healthcare Biosciences, Waukesha, WI, USA) according to Rasche et al. (2006), and 200 ng of each amplicon were digested with 5 U *Alu*I (archaeal 16S rRNA; New England Biolabs (NEB) Inc., Ipswich, MA, USA), overnight at 37°C (Rasche et al., 2011). For the digestion of *nosZ* and *arc-nosZ* gene amplicons, several restriction enzymes and their combinations

were tested (data not shown). Finally, 10 U HpyCH4IV (total prokaryotic nosZ clade I; NEB) and a 10 U cocktail of AluI, RsaI, and HpyCH4IV (arc-nosZ; NEB) revealed representative T-RFLP fingerprints of both genes. Prior to T-RFLP analysis, digests were purified with SephadexTM G-50 (Rasche et al., 2006) and a 2 µl aliquot was mixed with 17.75 µl HiDi formamide (Applied Biosystems) and 0.25 µl internal size standard (500 ROX Size Standard, Applied Biosystems). Labeled terminal-restriction fragments (T-RFs) were denatured at 95°C for 3 min, chilled on ice and detected on an ABI 3130 automatic DNA sequencer (Applied Biosystems). Electropherograms were compiled into numeric data using the Peak Scanner software (version 1.0, Applied Biosystems) and fragment length with peak height >50 fluorescence units were used for profile comparison. Raw data were normalized prior to statistical analysis (Dunbar et al., 2000).

2.4. Statistical data analysis

Data on abundance of the studied genes and soil chemical properties were subjected to ANOVA using Statistical Analysis Software program (SAS V 9.2, SAS Institute Inc., North Carolina, USA). For statistical analysis of the gene abundance data, a generalized linear mixed model with a negative binomial distribution error and a log link function was used. Effects of straw addition (factor "straw") and incubation time (factor "time") as well as the interaction between both factors on abundance of all studied genes and on chemical properties were tested by two-way analysis of variance (ANOVA). The data were checked for normality and homoscedasticity on model residuals using quantile–quantile (Q–Q) plots, histograms and studentized residual plots.

Pearson linear correlation analyses were conducted and visualized for linearity in the SAS COR

procedure (data not shown) to relate the abundance of the target genes (dependent variables) to the soil physico-chemical properties (independent variables).

Co-occurrence/co-exclusion analysis between the abundances of *arc-nosZ* and *gdh* genes and

chemical properties (NH₄⁺, DON) was carried out by using the "corrplot" package of R (http://www.r-project.org). The analysis was performed controlling the "False Discovery Rate"

279 (FDR), as described by Benjamini and Hochberg (1995). The adjustment methods include the 280 Bonferroni correction ("bonferroni") in which the p-values are multiplied by the number of 281 comparisons. 282 TRFLP data sets were analyzed using Bray-Curtis similarity coefficients (Bray and Curtis, 1957). A similarity matrix was generated for 16S archaea, total prokaryotic nosZ (nosZ) and archaeal nosZ 283 284 (arc-nosZ) target genes This similarity matrix was used for one-way analysis of similarity 285 (ANOSIM) statistics (Clarke, 1993) to test if the composition of target communities was altered by 286 factors "straw", "time" and by "straw x time" interaction. ANOSIM is based on rank similarities 287 between the sample matrix and produces a test statistic 'R' (Rees et al., 2005). A 'global' R was 288 first calculated in ANOSIM, which evaluated the overall effect of a factor in the data set. This step 289 was followed by a pair wise comparison, whereby the magnitude of R indicated the degree of 290 separation between two tested communities. An R score of 1 indicated a complete separation, while 291 0 indicated no separation (Rees et al., 2005). 292 Statistical analysis of T-RFLP profiles was performed with Primer V6.1.13 software (Primer E, 293 Plymouth, UK) according to Rasche et al. (2011). 294 Additionally, to test the influence of NH₄ on the structure of nosZ and arc-nosZ genes among straw 295 and no straw treatments over the incubation time, principal component analysis (PCA) was 296 performed using the "dudi.pca" function of in R-package ade4 (Thioulouse et al. 1997; Dray et al. 297 2007). Thus, to indicate a redundancy in the data, a correlation matrix was constructed, where the 298 presence or absence as well as relative height of T-RFs were used as distinct data, whereas NH₄⁺ 299 data were included in the analysis as 'environmental' variable. The data were grouped by straw and 300 no straw treatments over the incubation time. The graph of the PCA represents the differences 301 induced by NH₄⁺ in the community structures of nosZ and arc-nosZ genes across the treatments at 302 each sampling time: 1, 5, 10, 20, 30, and 60 days, according to the axes 'x' and 'y', which represent 303 eigen vectors with the greater variance.

305 3. Results

- 306 3.1. Soil chemical properties
- After submergence, soil pH showed a progressive increase from 5.8 to 7.0 (p<0.01) after 60 days of
- incubation for both treatments with or without straw. This was in line with our previous results
- 309 confirming the onset of anaerobic conditions with a decrease in redox potential (Eh) from +293 mV
- 310 to -180 and +110 mV over the first 30 days of incubation, in the presence or absence of straw,
- respectively (Cucu et al., 2014).
- N₂O fluxes from soils receiving straw were negligible throughout the incubation period, while soils
- 313 not receiving straw showed a peak in N₂O emissions on the first three days immediately after
- mineral N addition, followed by negligible emissions for the remaining incubation period (Fig.1).
- 315 Under submerged conditions, NH₄⁺ was the main inorganic N form (Figure 2a). In the presence of
- straw, NH₄⁺ decreased from 57 mg N kg⁻¹ to 41 mg N kg⁻¹ in the first 20 days and subsequently
- increased to values similar to the initial concentration by the end of the incubation (time × straw
- interaction, p<0.001). In contrast, in the absence of straw, NH₄⁺ concentrations decreased steadily to
- 319 23 mg N kg⁻¹ by day 60 (p<0.001).
- 320 Soils receiving straw showed NO₃- concentrations below detection limit throughout the incubation
- period (Fig. 2b). In the absence of straw, NO₃ present at the beginning of the incubation (5 mg N
- 322 kg⁻¹) rapidly disappeared within the first 5 days, followed by an increase with time reaching 4 mg N
- kg^{-1} at day 30 and 3 mg N kg^{-1} at day 60 (p<0.001).
- TDN concentrations in soils receiving straw increased from 66 to 76 mg N kg⁻¹ with time (p<0.001;
- Fig. 2c), and DON, calculated as the difference between TDN and mineral N (i.e., NH₄⁺-N + NO₃⁻-
- N), increased from 10 to 20 mg N kg⁻¹ at day 60. On the other hand, in soils without straw addition,
- 327 TDN decreased from 66 to 32 mg N kg⁻¹ by 60 days (time × straw interaction, p<0.001) mainly due
- to a loss of inorganic N-forms, since DON only varied from 10 to 6 mg kg⁻¹ over the same time
- 329 period.

During the entire incubation, soils with straw showed higher DOC concentrations with respect to soils without straw (p<0.01), with initial concentrations of approximately 155 mg and 79 mg C kg⁻¹, respectively (Fig. 2d). In both treatments, DOC contents decreased rapidly within the first 5 days of the incubation and increased thereafter reaching highest values by day 60 (192 and 110 mg C kg⁻¹ in straw and no straw treatments, respectively). Increase of DOC was more pronounced for soils receiving straw (time × straw interaction, p<0.01).

336

337

338

339

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

330

331

332

333

334

335

3.2. Microbial abundance

Abundance of bacterial 16S rRNA genes generally decreased with time, particularly for soils receiving straw (time × straw interaction; p<0.001), with values ranging between 1.0×10¹⁰ and 2.7×10¹⁰ copy numbers g⁻¹ soil (Fig. S1a). Variations in archaeal 16S rRNA gene abundance with time generally followed a trend similar to that observed for bacterial 16S rRNA gene abundance (time × straw interaction; p<0.001), also exhibiting a higher gene copy number in soils without straw (8.5×10⁹) with respect to those with straw (6.5×10⁹9. However, soil samples receiving straw showed a more pronounced decrease in archaeal 16S rRNA gene copy numbers from day 5 onwards with respect to bacterial 16S rRNA genes. After 60 days of incubation, archaeal 16S rRNA gene abundance declined to 1.1×10⁹ and 5.5×10⁸ copy numbers g⁻¹ soil for the no straw and straw treatments, respectively (Fig. S1b). Abundance of gdhA genes ranged from 1.3×109 to 2.0×109 copy numbers g-1 soil in the straw treatment, and from 1.1×10^9 to 1.8×10^9 copy numbers g^{-1} soil in the no straw treatment (Fig. S2). In both treatments, gdhA gene abundance decreased during the first 5 days of incubation, but subsequently increased with time, generally showing higher values for soils receiving straw than those not receiving straw by the end of the incubation time (time × straw interaction; p<0.001). Abundance of total prokaryotic *nosZ* clade I genes ranged from 2.3×10^7 to 8.6×10^7 copy numbers g^{-1} soil in the straw treatment and from 2.5×10^7 to 6.2×10^7 copy numbers g^{-1} soil in the no straw treatment (Fig. 3a), while the abundance of arc-nosZ genes ranged from 7×106 to 6.3×107 copy

- numbers g⁻¹ soil and from 1.7×10⁷ to 6.4×10⁷ copy numbers g⁻¹ soil for the straw and no straw
- 357 treatments, respectively (Fig. 3b). For both genes, the abundance tended to decrease during the first
- 358 10 days of incubation, particularly for the arc-nosZ genes in the straw treatment (time × straw
- interaction; p<0.001) and increased again during the later phase of incubation (between 20-60
- 360 days).
- 361 The relative abundance of *arc-nosZ* genes with respect to total prokaryotic *nosZ* clade I genes,
- expressed as *arc-nosZ/nosZ* ratio, varied during the incubation time from 0.6 to 1 and from 1 to 0.2
- in the no straw and straw treatments, respectively (data not shown).
- 364 Soils not receiving straw showed stronger positive correlations between gene abundance and
- 365 chemical properties with correlation coefficients ranging from r = 0.216 (p<0.01) to 0.879
- 366 (p<0.001) (Table 3). On the other hand, soils receiving straw showed correlation coefficients
- ranging between r = 0.153 (p<0.01) and 0.574 (p<0.001).
- The co-occurrence and co-exclusion between arc-nosZ and gdhA gene abundance and NH₄⁺ and
- 369 DON were investigated by considering the significant correlations at False Discovery Rate (FDR) <
- 370 0.05 (Fig. S3). Abundance of *arc-nosZ* genes showed positive correlations with the abundance of
- 371 the *gdhA* gene and NH₄⁺ content, while a negative correlation was found for DON contents (FDR <
- 372 0.05).

- 3.74 3.3. Microbial community structure
- 375 ANOSIM was used to compare effects of "straw" and "time" on the community structure of
- archaeal 16S rRNA gene, nosZ clade I genes and arc-nosZ genes (Table 4). Factor "time" had a
- 377 greater influence than factor "straw" on all studied genes (p<0.001). ANOSIM detected significant
- interactions between both factors (archaeal 16S rRNA (R=0.724), total prokaryotic nosZ clade I
- 379 (R=0.691), arc-nosZ (R=0.735) genes; p<0.001)).
- 380 NH₄⁺ had a distinct effect on the determined shifts in the community structures of nosZ and arc-
- 381 nosZ genes across the treatments during the incubation time (Fig. 4 and 5). The PCA clearly showed

- a clustering of nosZ straw treated samples (Fig. 4a) from 10, 20 and 30 days which were well
- separated from samples belonging to 1 and 5 and 60 days. A different pattern was revealed by the
- PCA of *nosZ* no straw treated samples (Fig. 4b) which clustered in two different groups: samples
- from 1 and 5 days and samples belonging to 10, 20, 30 and 60 days.
- The PCA of arc-nosZ straw samples (Fig. 5a) showed a separation of samples from 10, 20, 30 and
- 387 60 days from samples belonging to 1 and 5 days, while the PCA of *arc-nosZ* no straw samples (Fig.
- 388 5b) revealed community differences between samples from 20, 30 and 60 days, separated from
- samples belonging to 1, 5 and 10 days.
- 390
- 391 3.4. DNA ¹⁵N-enrichment
- 392 Addition of labeled (99 atom% ¹⁵N) (NH₄)₂SO₄ resulted in a progressive ¹⁵N-enrichment of DNA
- with respect to natural ¹⁵N abundance (0.3663 atom% ¹⁵N) (Fig. S4). The highest enrichment was
- obtained after 30 days of incubation, corresponding to 6.8 and 5.4 atom% ¹⁵N excess in the straw
- and no straw treatments, respectively (time × straw interaction; p<0.001).
- 396
- 397 3.5. ¹⁵N-SIP analysis
- 398 T-RFLP fingerprints of *arc-nosZ* genes were generated from heaviest SIP gradient fractions (1.70 to
- 399 1.76 g ml⁻¹) to identify those community members which most efficiently incorporated the ¹⁵N
- 400 tracer (Fig. 6). As expected, community structure in fractions showed a lower diversity as compared
- 401 to the bulk samples, since only a part of the community was active in ¹⁵N assimilation. In all
- fractions, with or without straw, the arc-nosZ gene community was dominated by two T-RFs (i.e.,
- 403 191 and 273 bp), but also by other T-RFs (e.g., 107, 117, 164, 185 bp) which were detected in
- several fractions. T-RFLP fingerprinting revealed a clear distinction in T-RF allocation between ¹⁴N
- 405 (Fig. 6a and 6c) and ¹⁵N SIP gradients (Fig. 6b and 6d) suggesting a successful ¹⁵N enrichment in
- 406 the heaviest fractions.

Conversely, total *nosZ* gene clade I T-RFLP fingerprinting analysis showed no ¹⁵N-enrichment in all fractions of both the straw and no straw treatments (data not shown).

Incorporation of the ¹⁵N tracer in *arc-nosZ* genes was further corroborated by quantification of

Incorporation of the ¹⁵N tracer in *arc-nosZ* genes was further corroborated by quantification of archaeal *gdhA* genes performed on heavy CsCl gradient fractions in both treatments, with or without straw. In the straw treatment (Fig. 7a), label incorporation was evident in the ¹⁵N labeled gradient, where the gene abundance peak shifted to a buoyant density (BD) of 1.7594 g ml⁻¹ compared to that of the ¹⁴N control (1.7570 g ml⁻¹). This was similar in the no straw treatment, where the gene abundance peak in the ¹⁵N gradient shifted to a BD of 1.7157 g ml⁻¹ compared to the ¹⁴N control with 1.6987 g ml⁻¹ (Fig. 7b). The shift in BD of approximately 0.02 g ml⁻¹ corresponded to 50% of the density shift of 0.04 g ml⁻¹ expected for 100% label incorporation (Lueders et al., 2004). In addition, the presence of ¹⁵N labeled DNA into even heavier fractions (BD of 1.7365 to 1.7560 g ml⁻¹) indicated an overall higher label incorporation in the no straw with respect to straw ¹⁵N-gradient.

4. Discussion

422 4.1. Archaeal N assimilation

Rice straw addition to submerged rice paddy soils represented the key factor for observed diverse responses of total bacterial and archaeal communities. This suggested that soil N immobilization and gaseous losses – as quantified in the previous experiment as a function of straw application (Cucu et al., 2014) – were controlled by both bacteria and archaea. However, under our experimental conditions, the archaeal community showed a clear domination over bacteria, especially in the absence of straw. In contrast to denitrifying bacteria, which usually use a dissimilatory metabolism under non-limiting NH₄⁺ conditions, archaeal denitrifiers rely on an assimilatory pathway to acquire N for formation of nucleic and amino acids as well as proteins (Cabello et al., 2004; Nannipieri and Paul, 2009; Rusch, 2013). This was confirmed by the high abundance of *gdhA* gene and progressive ¹⁵N-enrichment of DNA that displayed the effective biotic

immobilization of N during incubation. These observations suggested that independently of straw addition, archaeal N assimilation was an important process in the studied soil. Moreover, the similar increase in gdhA gene abundance in both straw-treated and non-treated soils suggested that archaeal assimilation under non-NH₄⁺ limiting environments was hardly influenced by the presence of labile organic C. However, the negative correlation between gdhA gene abundance and DOC indicated a stronger influence of soil derived C sources with respect to that of added straw. Moreover, although gdhA gene abundance was positively correlated with NH₄⁺ content in both treatments, this relationship was stronger for soils not receiving straw with respect to straw treated soils. On the other hand, the stronger correlation between gdhA gene abundance and TDN (i.e., the sum of inorganic and organic N) with respect to NH₄⁺ for soils receiving straw suggested that, although archaeal N assimilation was driven by NH₄⁺ availability, archaea could have modified their metabolism with time to obtain N from other easily accessible sources (e.g., dissolved organic N (DON) from labile OM) (Offre et al., 2013). Support for this was given by the negative correlation between gdhA gene abundance and DON which was in line with a rapid consumption and turnover of this labile N pool and the consequent increase in inorganic available N forms (Cucu et al., 2014). In addition, the slightly higher *gdhA* gene abundance in the presence of labile organic matter (OM) could be most likely attributed to an enhanced supply of NH₄⁺ resulting from the mineralization of added straw and dissolved OM released under anoxic conditions (Cucu et al., 2014).

451

452

453

454

455

456

457

458

433

434

435

436

437

438

439

440

441

442

443

444

445

446

447

448

449

450

4.2. N assimilation as support for archaeal denitrification

¹⁵N-DNA-based SIP revealed specific archaeal community members involved in both N assimilation and denitrification processes. The ¹⁵N-labeled NH₄⁺ was actively assimilated via *gdhA* gene for subsequent utilization (e.g., N₂O reductase synthesis), N immobilization and community growth. Although labeling success of DNA was apparently low in comparison to other studies (e.g., Buckley et al., 2007a; España et al., 2011), a clear labeling effect was observed in *arc-nosZ* gene community fingerprints of 'heavy' fractions in ¹⁵N-enriched samples compared to control

treatments. Moreover, NH_4^+ assimilation was confirmed by a higher abundance of gdhA gene in heavy ^{15}N labeled fractions, especially in the absence of straw.

Total prokaryotic nosZ clade I gene fingerprinting in the SIP fractions showed no evident 15N assimilation (data not shown). This result indicated especially in the straw treatment the shift in the total prokaryotic nosZ community towards bacterial members and suggested that bacterial denitrifiers were most probably using only N as an electron sink and for energy conservation (Zumft 1997). Archaeal N assimilation resulted in an increase in the abundance of assayed N₂O reduction genes (arc-nosZ), indicating the predominance of denitrifying archaea compared to their bacterial counterparts, especially in soils not receiving straw. This was confirmed by the high relative abundance of arc-nosZ with respect to total prokaryotic nosZ clade I genes, as evidenced by arc-nosZ/nosZ ratios. Since the abundance of total prokaryotic nosZ clade II genes is in the same range of clade I (Jones et al., 2013), quantification of clade II may result in an underestimation of the archaeal proportion of the total N₂O reducing community. However, archaeal members may still have a significant role in completing the denitrification process. The abundance of total prokaryotic nosZ clade I and specifically arc-nosZ genes was reflected in negligible N₂O emission fluxes over the whole incubation period in both treatments. This confirmed that the gaseous losses of applied N (up to 60%) in the absence of straw (c.f. only 20% lost in the presence of straw), reported by Cucu et al (2014) could be mainly attributed to N₂ emissions. These findings were in line with previous studies reporting high N₂O reduction under continuous rice soil flooding (DeDatta 1995; Ussiri and Lal, 2013; Peyron et al., 2016).

479

480

481

482

483

478

459

460

461

462

463

464

465

466

467

468

469

470

471

472

473

474

475

476

477

4.3. Resource driven archaeal NH_4^+ assimilation and denitrification

It is generally assumed that net ammonification and consequently N availability are greater under anaerobic soil conditions due to low metabolic N requirements of anaerobic microorganisms (Reddy and DeLaune, 2008). However, our findings indicated that archaeal denitrifiers were able to

484 assimilate N under non-limiting NH₄⁺ conditions and that this metabolic modification was most 485 likely a function of C and N availability. 486 Observed alterations of arc-nosZ gene community dynamics in soils without straw suggested a 487 greater efficiency in NH₄⁺ assimilation and N immobilization than in the presence of straw. This 488 was probably due to a more competitive and versatile metabolism of chemoautotrophic 489 microorganisms (e.g., archaea) being well-adapted to resource limitation. Although it is generally 490 accepted that increased C substrate availability increases microbial population size (Anderson and 491 Domsch 1978; Fontaine et al. 2003) and thus activity, our results clearly indicated that under C 492 limiting conditions archaeal denitrifiers were most likely equipped with a unique metabolic 493 flexibility to scavenge alternative nutrient sources (Müller et al., 2014). Accordingly, in the absence 494 of alternative labile organic C (e.g., rice straw), the microbial communities might utilized carbon 495 dioxide as their main C source (Bock et al., 1991), reduced compounds (e.g., ammonium, iron(II), 496 and sulfide) as electron donors (Liesack et al., 2000; Megonigal et al., 2004), or N₂O as sole 497 electron acceptor (Stres et al., 2004; Strohm et al., 2007; Braker and Conrad, 2011). This suggested 498 a contrast with the general assumption that prokaryotic denitrifiers are predominantly heterotrophic 499 (Parkin 1987). 500 Independent of the adopted agricultural management, our results might display that rice paddy soils 501 act as temporary sink for N₂O. Furthermore, abundance of both denitrifying genes was positively 502 correlated to NH₄⁺ and TDN concentrations highlighting the co-occurence of both complete 503 denitrification and ammonification in the same genome (Tiedje et al., 1988; Sanford et al., 2012). 504 Support for this was also given through the negative correlation of arc-nosZ gene abundance with 505 DON contents. Likewise, mineralization promotes the recycling of immobilized N which may then 506 enter further processes including denitrification (Nannipieri et al., 2003). 507 As expected, the higher availability of labile C with straw addition under non-limiting N conditions 508 induced N assimilation but also a lower abundance of total nosZ clade I and arc-nosZ genes, as well 509 as distinct variation in the community structure of these genes. Lower abundance of nosZ genes in

the presence of straw was in contrast with other studies (Chen et al., 2012a,b). However, at the later stages of incubation, when most of added straw was probably decomposed, progressively higher abundance of *nosZ* and *arc-nosZ* genes was positively related to DOC content which was in line with the findings of Kandeler et al., (2006) and Philippot et al., (2009). Labile organic C served as resource for denitrifying bacteria that outcompeted archaeal counterparts, as evidenced by a decreasing *arc-nosZ/nosZ* ratio with incubation time. This was in agreement with Ishii et al. (2011b) who, by using functional single-cell (FSC) and DNA-based SIP with ¹³C-labeled succinate as electron donor and N₂O as electron acceptor showed that under their experimental conditions, most N₂O reducers are bacterial denitrifiers. Examining the N₂O reduction rates of the isolated strains, the authors confirmed the growth of putative bacterial denitrifiers reciprocally to N₂O reduction in rice field soils, although many bacteria have only partial pathways of denitrification (Shapleigh, 2013).

5. Conclusions and outlook

Nitrogen immobilization based on archaeal NH_4^+ assimilation was shown to represent an important step for proliferation and dynamics of those microbial community members harboring the *arc-nosZ* gene encoding N_2O reductase enzyme, irrespective of the presence or absence of labile OM. Under our experimental conditions, the relative abundance of *arc-nosZ* genes with respect to total prokaryotic *nosZ* from clade I genes was based on an adaptable metabolic portfolio. However, in the presence of straw, bacterial *nosZ* genes may also contribute to the completion of denitrification. Based on these considerations, we developed a conceptual model to represent the archaeal N assimilation and denitrification pathways driven by different resources under anaerobic conditions (Fig. 8). The response of *arc-nosZ* genes to different C availability under non-limiting N conditions may have important ecological implications in controlling the immobilization and loss of N from fertilized paddy soils. Furthermore, the high adaptability of archaea to drive denitrification to completion may be a key feature in mitigating N_2O emissions.

The present study suggested the potential contribution of archaea to the last step of denitrification when favorable environmental conditions are given as complement to the bacterial counterparts from *nosZ* clade I. However, although we have provided important insights, the full understanding of archaeal involvement in the denitrification process remains still incomplete. This accounts particularly for the currently limited phylogenetic information on *arc-nosZ* gene synthesizing archaea in soils (Rusch 2013). Hence, we strongly suggest the generation of *arc-nosZ* gene libraries to enhance phylogenetic knowledge on this particular gene. In addition, prospective research should also consider the unconsidered *nosZ* gene (e.g., clade II, atypical *nosZ*) to better understand the relative role of archaea in acting as N₂O sink.

As N assimilation processes are regulated by resource availability, we recommend evaluating the effects of different N fertilizer and organic residue types with contrasting C/N ratio. In this respect, ¹⁵N and ¹³C-label SIP-based studies may be appropriate to evaluate the metabolic adaptability of archaea for mineral and organic substrate consumption and their implication in mitigating emissions

Funding

- This work was partly supported by the Italian Ministry of Agriculture, Food and Forestry
- (MiPAAF) under the project "Sustaining the National Rice Industry through Research, Technology,
- Innovation and Formation (POLORISO)" and International Humic Substance Society (IHSS).

Acknowledgements

of climate relevant gases from paddy soils.

- We gratefully thank R. Gorra (Department of Agricultural, Forest and Food Sciences, University of
- 558 Turin, Italy) for fruitful discussions; J. Laso (Department of Bioinformatics, University of
- Hohenheim, Germany) and I. Ferrocino (Department of Agricultural, Forest and Food Sciences,
- University of Turin, Italy) for their support in statistical analysis. The authors are further grateful to
- L. Hölzle (Institute of Animal Science, University of Hohenheim) for access to his ABI 3130

- Genetic Analyzer and C. Röhl (Institute of Plant Production and Agroecology in the Tropics and
- 563 Subtropics, University of Hohenheim) for her technical support.

564	References
565	Anderson JPE, Domsch KH (1978) A physiological method for the quantitative measurement of
566	microbial biomass in soils. Soil Biol Biochem: 10: 215–221
567	Bird JA, Horwáth WR, Eagle AJ, van Kessel (2001) Immobilization of fertilizer nitrogen in rice.
568	Soil Sci Soc Am J 65:1143-1152
569	Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful
570	approach to multiple testing. J R Stat Soc Series B 57: 289-300
571	Bray JR, Curtis JT (1957) An ordination of the upland forest communities of southern Wisconsin.
572	Ecol Monogr 27:325-349
573	Bock E, Koops HP, Harms H, Ahlers B (1991) The biochemistry of nitrifying organisms. In:
574	Shively JM, Barton LL (eds) Variations in autotrophic life, London, Academic Press, pp
575	171-200
576	Bonete MJ, Martínez-Espinosa RM, Pire C, Zafrilla B, Richardson DJ (2008) Nitrogen metabolism
577	in haloarchaea. Saline Systems 4:9
578	Braker G and Conrad R. Diversity, structure, and size of N ₂ O-producing microbial communities in
579	soils-what matters for their functioning? (2011) In: Laskin I, Sariaslani S, Gadd GM (eds)
580	Advances in Applied Microbiology, vol.75, SanDiego, Elsevier Academic Press Inc., pp 33-
581	70
582	Buckley DH, Huangyutitham V, Hsu SF, Nelson TA (2007a) Stable isotope probing with ¹⁵ N ₂
583	reveals novel non-cultivated diazotrophs in soil. Apppl Environ Microbiol 73:3196-3204
584	Buckley DH, Huangyutitham V, Hsu SF, Nelson TA (2007b) Stable isotope probing with ¹⁵ N
585	achieved by disentangling the effects of genome G+C content and isotope enrichment on
586	DNA density. Appl Environ Microbiol 73:3189-3195
587	Cabello P, Roldán MD, Moreno-Vivían C (2004) Nitrate reduction and the nitrogen cycle in
588	archaea. Microbiol 150:3527-3546

589	Cassman KG, Peng S, Olk DC, Ladha JK, Reichardt W, Dobermann A, Singh U (1998)
590	Opportunities for increased nitrogen-use efficiency from improved resource management in
591	irrigated rice systems. Field Crop Res 56:7-39
592	Cassman KG, Dobermann A, Walters DT (2002) Agro-ecosystems, nitrogen-use efficiency, and
593	nitrogen management. Ambio 31:132-140
594	Chen Z, Liu J, Wu M, Xie X, Wu J, Wei W (2012a) Differentiated response of denitrifying
595	communities to fertilization regime in paddy soil. Microb Ecol 63:446-459
596	Chen Z, Hou H, Zheng Y, Qin H, Zhu Y, Wu J, Wei W (2012b) Influence of fertilization regimes
597	on a nosZ-containing denitrifying community in a rice paddy soil. J Sci Food Agr 92:1064-
598	1072
599	Chuang H-P, Wu J-H, Ohashi A, Abe K, Hatamato M (2014) Potential of nitrous oxide conversion
600	in batch and down-flow hanging sponge bioreactor systems. Sustain Environ Res 24:117-
601	128
602	Clarke KR (1993) Non-parametric multivariate analyses of changes in community structure. Aust J
603	Ecol 18: 117-143
604	Conrad R (1995) Soil microbial processes involved in production and consumption of atmospheric
605	trace gasses. In: Jones JG (ed) Advances in Applied Microbiology, New York. Plenum
606	Press, pp 207-250
607	Conrad R (1996) Soil microorganisms as controllers of atmospheric trace gases (H2, CO, CH4,
608	OCS, N ₂ O, and NO). Microbiol Rev 60:609-640
609	Cucu MA, Said-Pullicino D, Maurino V, Bonifacio E, Romani M, Celi L (2014) Influence of redox
610	conditions and rice straw in corporation on nitrogen availability in fertilized paddy soils.
611	Biol Fertil Soil 50:755-764
612	Davidson EA, Swank WT, Perry TO (1986) Distinguishing between nitrification and denitrification
613	as sources of gaseous nitrogen production in soil. Appl Environ Microb 52:1280-1286
614	DeDatta SK (1995) Nitrogen transformations in wetland rice ecosystem. Fert Res 42:193-203

615	de Vries S, Schröder I (2002) Comparison between the nitric oxide reductase family and its aerobic
616	relatives, the cytochrome oxidases. Biochem SocTrans 30:662-667
617	Devêvre OC, Horwáth WR (2001) Stabilization of fertilizer nitrogen-15 into humic substances in
618	aerobic vs. waterlogged soil following straw incorporation. Soil Sci Soc Am J 65:499-510
619	Dray S, AB Dufour (2007) The ade4 package implementing the duality diagram for ecologist. J Stat
620	Softw 22:1-20
621	Dunbar J, Ticknor LO, Kuske CR (2000) Assessment of microbial diversity in four south western
622	United States soils by 16S rRNA gene terminal restriction fragment analysis. Appl Environ
623	Microb 66:2943-2950
624	Eagle AJ, Bird JA, Horwáth WR, Linquist BA, Brouder SM et al (2000) Rice yield and nitrogen
625	utilization efficiency under alternative straw management practices. Agronom J 92:1096-
626	1103
627	España M, Rasche F, Kandeler E, Brune T, Rodriguez B et al (2011) Identification of active
628	bacteria involved in decomposition of complex maize and soybean residues in a tropical
629	Vertisol using ¹⁵ N-DNA stable isotope probing. Pedobiologia 54:187-193
630	Fazli P, Hasfalina CM, UmiKalsom MS, Azni I (2013) Review Article: Characteristics of
631	methanogens and methanotrophs in rice fields. AsPac J Mol Biol Biotechnol 21: 3-17
632	Ferrè C, Zechmeister-Boltenstern S, Comolli R, Andersson M, Seufert G (2012) Soil microbial
633	community structure in a rice paddy field and its relationships to CH_4 and N_2O fluxes. Nutr
634	Cycl Agrosyst 93:35-50
635	Fetzer S, Bak F, Conrad R (1993) Sensitivity of methanogenic bacteria from paddy soil to oxygen
636	and desiccation. FEMS Microbiol Ecol 12:107-115
637	Fontaine S, Mariotti A, Abbadie L (2003) The priming effect of organic matter: a question of
638	microbial competition? Soil Biol Biochem 35:837-843

639	Garcia JL, Tiedje JM (1982) Denitrification in rice soils. In: Dommergues YR, Diem HG (eds)
640	Microbiology of tropical soils and plant productivity. Springer Netherlands, Dordrecht, pp
641	187-208.
642	Ghosh BC, Bhat R (1998) Environmental hazards of nitrogen loading in wetland rice fields.
643	Environ Pollut 102:123-126
644	Henry S, Bru D, Stres B, Hallet S, Philippot L (2006) Quantitative detection of the nosZ gene,
645	encoding nitrous oxide reductase, and comparison of the abundances of 16S rRNA, narG,
646	nirK, and nosZ genes in soils. Appl Environ Microbiol 72:5181-5189
647	Hutchinson GL, Mosier AR (1981) Improved soil cover method for field measurement of nitrous
648	oxide fluxes. Soil Sci Soc Am J, 45: 311–316
649	Ishii S, Ikeda S, Minamisawa K, Senoo K (2011a) Nitrogen cycling in rice paddy environments:
650	past achievements and future challenges. Microbes Environ J 26:282-292
651	Ishii S, Ohno H, Tsuboi M, Otsuka S, Senno K (2011b) Identification and isolation of active N ₂ O
652	reducers in rice paddy soil. ISME J 5:1936-1945
653	Jones CM, Graf DRH, Bru D, Philippot L, Hallin S (2013) The unaccounted yet abundant nitrous
654	oxide-reducing microbial community: a potential nitrous oxide sink. ISME J 7:417-426
655	Kandeler E, Deiglmayr K, Tscherko D, Bru D, Philippot L (2006) Abundance of narG, nirS, nirK,
656	and nosZ genes of denitrifying bacteria during primary successions of a glacier foreland.
657	Appl Environ Microbiol 72:5957-5962
658	Kögel-Knabner I, Amelung W, Cao Z, Fiedler S, Frenzel P, Jahn R et al (2010) Biogeochemistry of
659	paddy soils. Geoderma 157:1-14
660	Lane D (1991) 16S/23S rRNA sequencing. In: Stackebrandt A and Goodfellow M (eds) Nucleic
661	Acid Techniques Systematics, UK, John Wiley, West Sussex, pp 115-175
662	Liesack W, Schnell S, Revsbech NP (2000) Microbiology of flooded rice paddies. FEMS Microbiol
663	Rev 24:625-645

664	Lueders T, Friedrich M (2000) Archaeal population dynamics during sequential reduction processes
665	in rice field soil. Appl Environ Microbiol 66:2732-2742
666	Lueders T, Manefield M, Friedrich MW (2004) Enhanced sensitivity of DNA and rRNA based
667	stable isotope probing by fractionation and quantitative analysis of isopycnic centrifugation
668	gradients. Environ Microbiol 6:73-78
669	Majumdar D (2013) Biogeochemistry of N ₂ O uptake and consumption in submerged soils and rice
670	fields and implications in climate change. Crit Rev Env Sci Technol 43:2653-2684
671	Martinez-Espinosa RM, Marhuenda-Egea FC, Bonete MJ (2001) Assimilatory nitrate reductase
672	from the haloarchaeon Haloferax mediterranei: purification and characterization. FEMS
673	Microbiol Lett 204:381–385
674	Megonigal JP, Hines ME, Visscher PT (2004) Anaerobic metabolism: linkages to trace gases and
675	aerobic processes. In: Schlesinger WH (ed) Biogeochemistry, Oxford, UK, Elsevier-
676	Pergamon, pp 317-424
677	Mulec MI, Ausec L, Danevčič T, Levičnik-Höfferle Š, Jerman V, Kraigher B (2014) Microbial
678	community structure and function in peat soil. Food Technol Biotechnol 52:180-187
679	Müller J, Hense BA, Marozava S, Kuttler Ch, Meckenstock RU (2014) Model selection for
680	microbial nutrient uptake using a cost-benefit approach. Math Biosci 255:52-70
681	Muyzer G, Dewaal EC, Uitterlinden AG (1993) Profiling of complex microbial populations by
682	denaturing gradient gel electrophoresis analysis of polymerase chain reaction-amplified
683	genes coding for 16S rRNA. Appl Environ Microbiol 59:695-700
684	Nannipieri P, Ascher J, Ceccherini MT, Landi L, Pietramellara G, Renella G (2003) Microbial
685	diversity and soil functions. Europ J Soil Sc 54:655-670
686	Nannipieri P, Paul EA (2009) The chemical and functional characterization of soil N and its biotic
687	components. Soil Biol Biochem 41:2357-2369
688	Neufeld JD, Vohra J, Dumont MG, Lueders T, Manefield M et al (2007) DNA stable-isotope
689	probing. Nat Protoc 2:860-866

690 Nogales B, Timmis KN, Nedwell DB, Osborn AM (2002) Detection and diversity of expressed 691 denitrification genes in estuarine sediments after reverse transcription-PCR amplification 692 from mRNA. Appl Environ Microbiol 68:5017-5025 693 Offre P, Spang A, Schleper C (2013) Archaea in biogeochemical cycles. Ann Rev Microbiol 67: 694 437-457 695 Parkin TB (1987) Soil microsites as a source of denitrification variability. Soil Sci Soc Am J 51: 696 1194-1199 697 Peyron M, Bertora C, Pelissetti S, Said-Pullicino D, Celi L, Miniotti E, Romani M, Sacco D (2016) 698 Greenhouse gas emissions as affected by different water management practices in temperate 699 rice paddies. Agric Ecosyst Environ 232:17-28 700 Philippot L (2002) Denitrifying genes in bacterial and archaeal genomes. Biochim Biophys Acta 701 1577:355-376 702 Philippot L, Čuhel J, Saby NPA, Chèneby D, Chroňáková A, Bru D et al (2009) Mapping field-703 scale spatial distribution patterns of size and activity of the denitrifiers community. Environ 704 Microbiol 11:1518-1526 705 Prieme A, Braker G, Tiedje JM (2002) Diversity of nitrite reductase (nirK and nirS) gene fragments 706 in forested upland and wetland soils. Appl Environ Microbiol 68:1893-1900 707 Rasche F, Hödl V, Poll C, Kandeler E, Gerzabek MH, Van Elsas JD et al (2006) Rhizosphere 708 bacteria affected by transgenic potatoes with antibacterial activities compared with the 709 effects of soil, wild type potatoes, vegetation stage and pathogen exposure. FEMS Microbiol 710 Ecol 56: 219-235 711 Rasche F, Knapp D, Kaiser C, Koranda M, Kitzler B, Zechmeister-Boltenstern S et al (2011) 712 Seasonality and resource availability control bacterial and archaeal communities in soils of a 713 temperate beech forest. ISME J 5:389-402 714 Rees GN, Baldwin DS, Watson GO, Perryman, S, Nielsen DL (2005) Ordination and significance

testing of microbial community composition derived from terminal restriction fragment

716 length polymorphisms: application of multivariate statistics. Ant Van Leeuwenhoek 86: 717 339-347. http://dx.doi.org/10.1007/s10482-005-0498-5 718 Reddy KR (1982) Nitrogen cycling in a flooded-soil ecosystem planted to rice (*Oryza sativa* L.). 719 Plant Soil J 67:209-220 720 Reddy KR, deLaune RD (2008) Biogeochemistry of wetlands: science and applications, USA, CRC 721 Press, Boca Raton, FL 722 Rice CW, Tiedje JM (1989) Regulation of nitrate assimilation by ammonium in soils and in isolated 723 soil microorganisms. Soil Biol Biochem 21:597-602 Rusch A (2013) Molecular tools for the detection of nitrogen cycling. Article ID 676450, Archaea 724 725 doi:10.1155/2013/676450 Rusch A (2016) Archaea. In Kennish MJ (ed) Encyclopedia of Estuaries, Chapter: 287, Springer 726 727 Science, Business Media Dordrecht, doi: 10.1007/978-94-017-8801-4 287, pp 35-37 728 Sahrawat KL (2004) Ammonium production in submerged soils and sediments: the role of 729 reducible iron. Commun Soil Sci Plant Anall 35:399-411 730 Said-Pullicino D, Cucu MA, Sodano M, Birk JJ, Glaser B, Celi L (2014) Nitrogen immobilization 731 in paddy soils as affected by redox conditions and rice straw incorporation. Geoderma 228-732 229:44-53 733 Samuel BS, Hansen EE, Manchester JK, Coutinho PM, Henrissat B, Fulton R et al (2007) Genomic 734 and metabolic adaptations of Methanobrevibacter smithii to the human gut. PNAS J 104: 735 10643-10648 736 Sanford RA, Wagner DD, Wu Q, Chee-Sanford JC, Thomas SH, Cruz-Garcia C et al (2012) 737 Unexpected nondenitrifier nitrous oxide reductase gene diversity and abundance in soils. PNAS J 48:19709-19714 738 739 Shapleigh JP. Denitrifying prokaryotes (2013). In: Rosenberg E et al (eds) Prokaryotes. Prokaryotic

Physiology and Biochemistry, Berlin Heidelberg, Springer-Verlag, pp 405-425

- 741 Stres B, Mahne I, Avgustin G, Tiedje JM (2004) Nitrous oxide reductase (nosZ) gene fragments 742 differ between native and cultivated Michigan soils. Appl Environ Microbiol 70:301–309 743 Strohm TO, Griffin B, Zumft WG, Schink B (2007) Growth yields in bacterial denitrification and 744 nitrate ammonification. Appl Environ Microbiol 73:1420–1424 Thioulouse J, Chessel D, Doledec S, Olivier JM (1997) ADE-4: a multivariate analysis and 745 746 graphical display software. Stat Comput 7:75–83 747 Tiedje JM (1988) Ecology of denitrification and dissimilatory nitrate reduction to ammonium. In: 748 Zehnder AJB (ed) Biology of Anaerobic Microorganisms, New York, John Wiley & Sons, 749 pp 179-244 750 Tomlinson GA, Jahnke LL, Hochstein LI (1986) *Halobacterium denitrificans* sp. nov., an extremely halophilic denitryfing bacterium. Int J Syst Bacteriol 36:66-70 751 752 Ussiri D, Lal R (2013) Nitrous oxide emissions from rice fields. In: Soil emission of nitrous oxide 753 and its mitigation, Netherlands, Springer, pp 213-242 754 WRB, IUSS Working Group WRB, 2007. World Reference Base for Soil Resources 2006, first 755 update 2007. World Soil Resources Reports No. 103. FAO, Rome, Italy 756 Zou J, Huang Y, Zheng X, Wang Y (2007) Quantifying direct N₂O emissions in paddy fields during 757 rice growing season in mainland China: dependence on water regime. Atm Environ 41: 8030-8042 758 759 Zumft WG (1992) The denitrifying prokaryotes. In: Balows A, Trüper HG, Dworkin M, Harder W, 760 Schleifer K-H (eds) The prokaryotes. A handbook on the biology of bacteria: ecophysiology, isolation, identification, application, New York, Berlin, Heidelberg, 761
- Zumft WG (1997) Cell biology and molecular basis of denitrification. Microbiol Mol Biol Rev 61:
 533-616

Springer-Verlag, 2nd edn, vol 1, pp 554-582

Figure Captions

767

- 768 **Fig. 1** Daily nitrous oxide (N₂O) flux over time in soils incubated with or without straw addition.
- 769 The error bars (n=3) represent the least significant difference at p=0.05

770

- 771 Fig. 2 Variations in concentrations of ammonium (sum of submergence water and soil extractable
- NH₄⁺) concentrations (a), nitrate (sum of submergence water and soil extractable NO₃⁻) (b), total
- dissolved N (sum of submergence water and soil water-extractable N) (c), and dissolved organic C
- (sum of submergence water and soil water-extractable C) (d) with time for soils incubated with or
 - without straw addition. The error bars (n=6) represent the least significant difference at p=0.05

776

775

- 777 Fig. 3 Abundance of total prokaryotic *nosZ* clade I (a) and *arc-nosZ* (b) genes over time in soils
- incubated with or without straw addition (n=6, means with standard errors). Different letters above
- bars indicate significant differences at p<0.05 between straw and no straw treatments at every
- sampling time (lowercase letters) as well as significant differences among different sampling times
- 781 (uppercase letters).

782

- 783 Fig. 4 Principal component analysis (PCA) of TRFLP profiles using nosZ gene T-RF relative
- abundance data obtained form straw (a) and no straw (b) treatments in function of NH₄⁺
- concentration. Sampling time are indicated with colors (T 1 day blue; T 5 days red; T 10 days
- 786 purple; T 20 days pink; T 30 days dark grey; T 60 days green)

- 788 Fig. 5 Principal component analysis (PCA) of TRFLP profiles using arc- nosZ gene T-RF relative
- abundance data obtained form straw (a) and no straw (b) treatments in function of NH₄⁺
- 790 concentration. Sampling time are indicated with colors (T 1 day blue; T 5 days red; T 10 days
- 791 purple; T 20 days pink; T 30 days dark grey; T 60 days green)

Fig. 6 T-RFLP fingerprints of arc-nosZ genes generated from density resolved SIP fractions of the straw $^{14}N-(NH_4)_2SO_4$ (a), $^{15}N-(NH_4)_2SO_4$ (b) and the no straw $^{14}N-(NH_4)_2SO_4$ (c), and $^{15}N-(NH_4)_2SO_4$ (c), and $^{15}N-(NH_4)_2SO_4$ (c), and $^{15}N-(NH_4)_2SO_4$ (d), $^{15}N-(NH_4)_2SO_4$ (e), and $^{15}N-(NH_4)_2SO_4$ (f), and $^{15}N-(NH_4)_2SO_4$ (g), and $^{15}N-(NH_4)_2SO_4$ (h) and the no straw $^{14}N-(NH_4)_2SO_4$ (g), and $^{15}N-(NH_4)_2SO_4$ (g), and $^{15}N-(NH_4)_2SO_4$ (NH₄)₂SO₄ (d) treatment. Buoyant densities (g ml⁻¹) of analyzed fractions are given in parentheses Fig. 7 Quantification of archaeal gdhA genes in comparative CsCl density fractions of DNA extracted from straw (a) and non-straw (b) treatments fertilized with either ¹⁴N- or ¹⁵N-¹⁵N- $(NH_4)_2SO_4$ Fig. 8 Schematic relationship between C and N resource availability and nitrogen assimilatory archaeal denitrification in fertilized paddy soils suggesting the ecological importance of this pathway

Graphical abstract

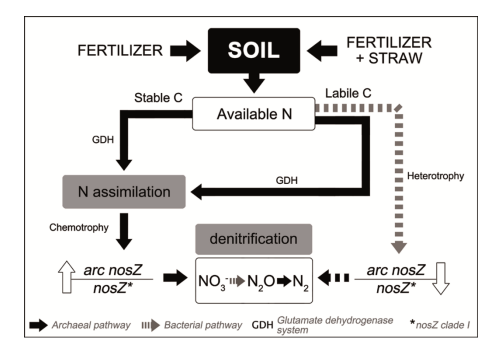


Figure 1

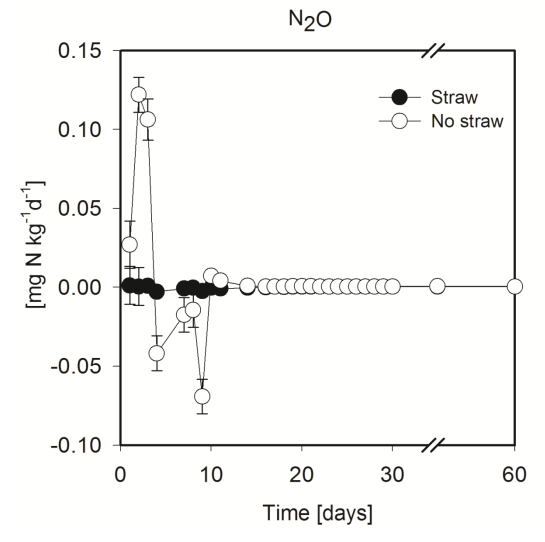


Figure 2

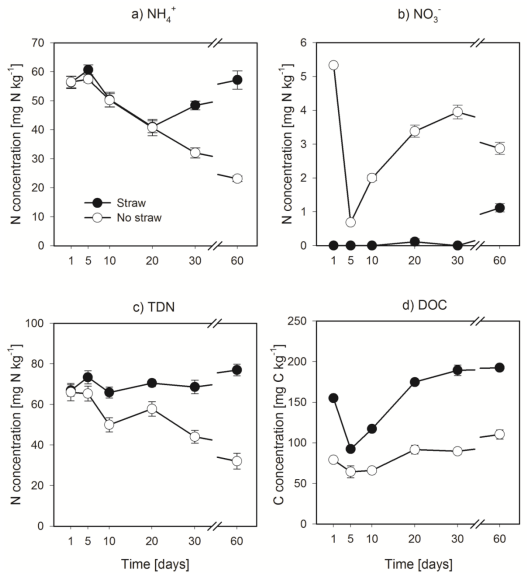


Figure 3

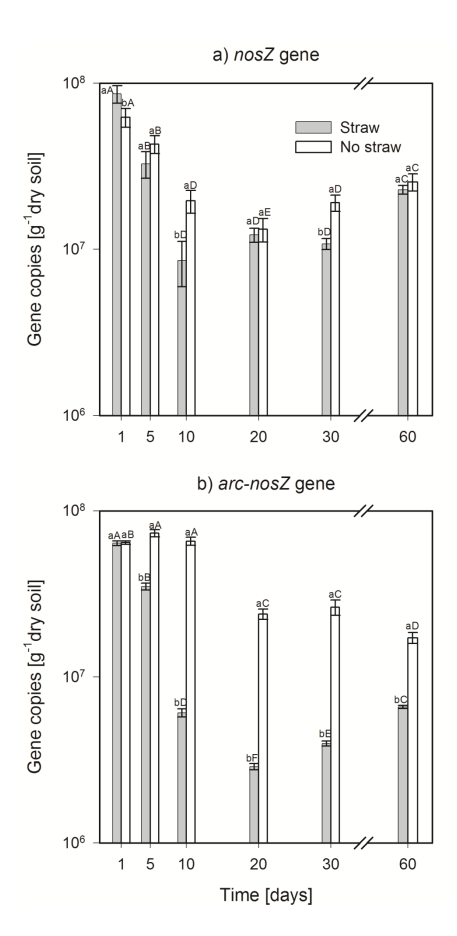


Figure 4

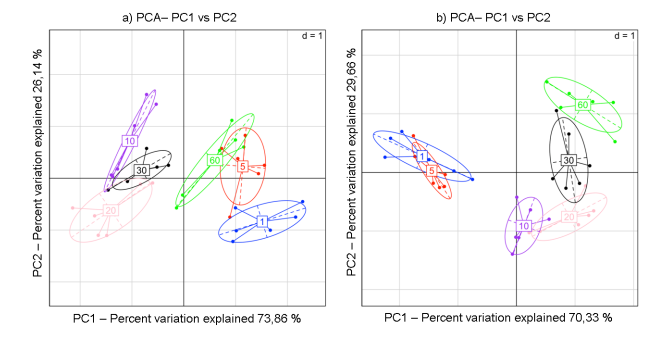


Figure 5

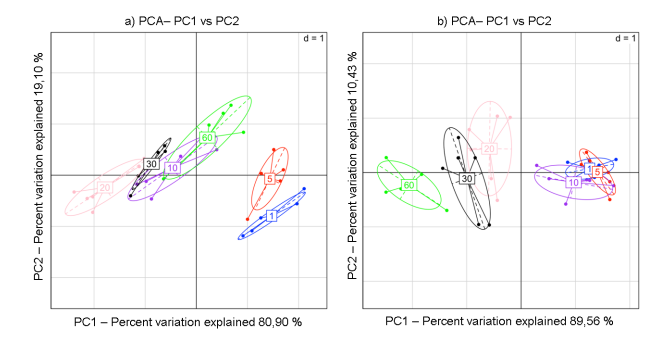
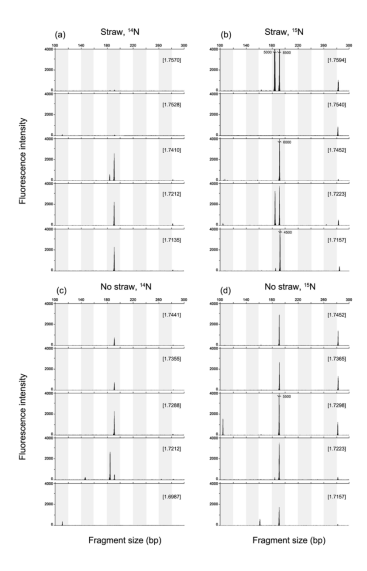
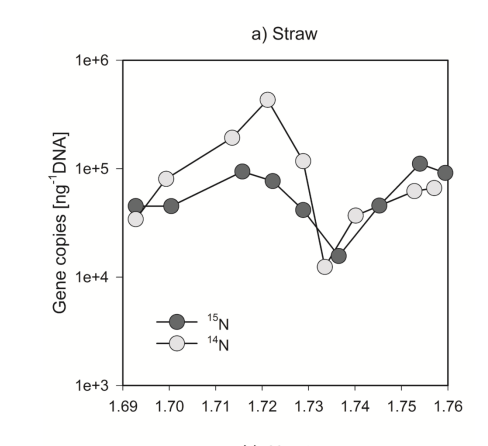


Figure 6





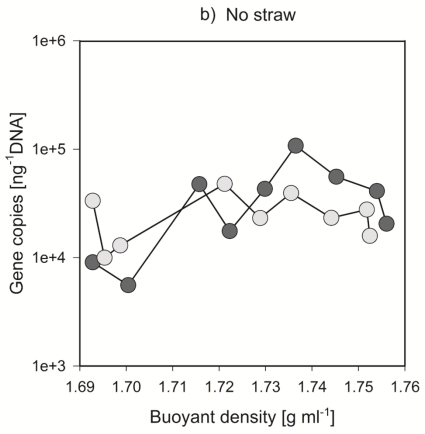


Figure 8

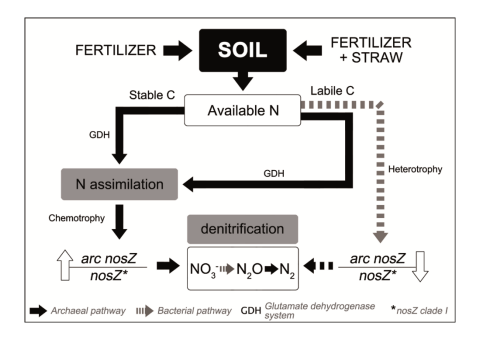


Table 1 Description of primer sets and amplification details used for quantitative PCR.

Target group	Primer (reference)	Amplification details
All bacteria	Eub338 (Lane 1991)	40 cycles
(16S rRNA gene)	Eub518 (Muyzer et al. 1993)	95°C 30s, 55°C 35s, 72°C 45s
All archaea	Ar109f (Lueders and Friedrich 2000)	40 cycles
(16S rRNA gene)	Ar912r (Lueders and Friedrich 2000)	95°C 30s, 52°C 35s, 72°C 45s, 78°C 20s
nosZ gene clade I	nosZ-2f (Henry et al. 2006)	6 touch down cycles
	nosZ-2r (Henry et al. 2006)	95°C 15s, 65°C 30s (-1), 72°C 30s
		40 cycles
		95°C 15s, 60°C 30s, 72°C 30s, 81°C 30s
arc-nosZ gene	arc-Nos-f (Rusch 2013)	6 touch down cycles
	arc-Nos-r (Rusch 2013)	95°C 15s, 65°C 30s (-1), 72°C 30s
		40 cycles
**		95°C 15s, 60°C 45s, 72°C 60s, 81°C 35s
gdhA gene	MSM0888f (Samuel et al. 2007)	40 cycles
	MSM0888r (Samuel et al. 2007)	95°C 30s, 58°C 35s, 74°C 30s

Table 2 Description of primer sets, PCR ingredients and amplification details used for T-RFLP analysis.

Target group	Primer (reference)	DNA (ng)	Taq (U)	MgCl ₂ (mM)	Primer (mM)	dNTPs (mM)	Amplification details
All archaea	Ar109f (Lueders and Friedrich 2000) Ar912r (Lueders and Friedrich 2000)	5	2	1.5	0.15	0.2	35 cycles 95°C 5 m, 95°C 60s, 52°C 30s, 72°C 60s,72°C10 m
nosZ gene clade I	nosZ-2f (Henry et al. 2006) nosZ-2r (Henry et al. 2006)	10	2	1.5	0.2	0.2	6 touch down cycles 94°C 30s, 65°C 30s (-1), 72°C 30s 40cycles 94°C 30s, 60°C 30s, 72°C 30s
arc-nosZ gene	arc-Nos-f (Rusch 2013) arc-Nos-r (Rusch 2013)	10	0.5	-	0.4	0.2	40 cycles 95°C 60s, 95°C 20s, 72°C 20s, 72°C 90s, 72°C 15 m

6 Table 3 Linear correlation coefficients (Pearson correlation coefficients, n=36) between microbial abundance and chemical data

Property	All bacter	ia	All archae	a	total nos	Z clade I	arc-nosZ		gdhA	
	straw	no straw	straw	no straw	straw	no straw	straw	no straw	straw	no straw
NH ₄ ⁺	0.351*	0.709***	0.469**	0.879***	0.414*	0.854***	0.457**	0.796***	0.313*	0.706***
TDN	ns	0.593***	ns	0.662***	ns	0.439**	ns	0.550***	0.574***	0.624***
NO_3^-	ns	ns	ns	ns	ns	0.319*	ns	0.272*	ns	ns
DOC	-0.464**	-0.450**	-0.470**	-0.742***	0.153*	0.216*	0.237*	0.246*	ns	-0.427**

Significance levels: not significant-ns: p>0.05; *p<0.05; **p<0.01; ***p<0.001.

Table 4 Analysis of similarity (ANOSIM) revealing the treatments effect and time on soil archaeal, total prokaryotic *nosZ* (*nosZ*) clade I and archaeal *nosZ* (arc-*nosZ*) denitrifying community structure

	Global R		
Factor	All archaea	nosZ	arc-nosZ
		clade I	
straw	ns	0.173***	0.472***
time	0.479***	0.358***	0.567***
	0.175	0.550	0.507

Levels of significance between two tested populations: not significant-ns: p>0.05; ***p<0.001.

Highlights

- Archaeal N assimilation was not affected by C availability
- N assimilation and immobilization preceded the archaeal denitrification
- In the presence of labile C denitrifying bacteria outcompeted archaeal counterparts
- Metabolic resilience induced a significant role of archaea in N₂O reduction
- A conceptual model of archaeal assimilatory denitrification was proposed

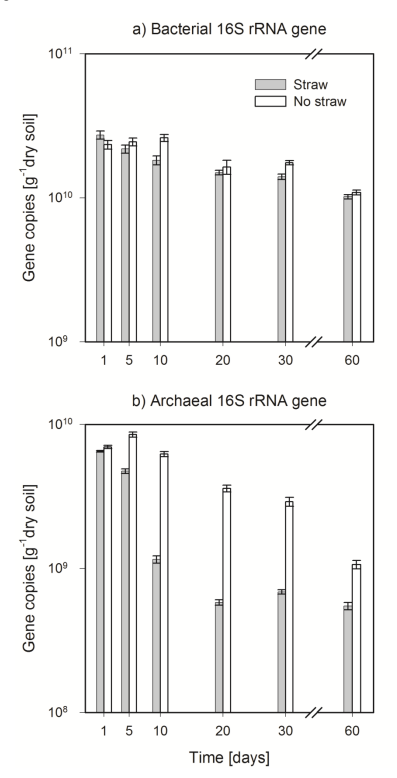


FIG. S1 Abundance of bacterial (a) and archaeal (b) 16S rRNA genes over time in soils incubated with or without straw addition (n=6, means with standard errors).

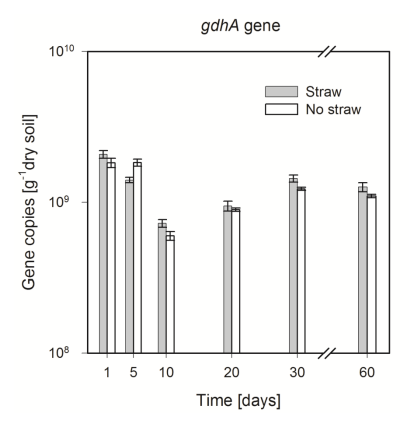


FIG. S2 Abundance of archaeal *gdhA* genes over time in soils incubated with or without straw addition (n=6, means with standard errors).

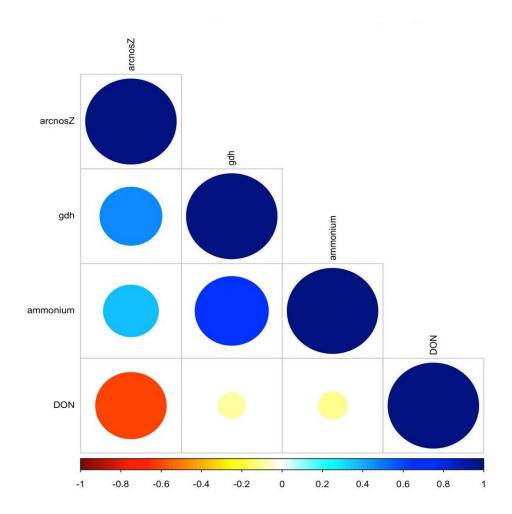


FIG. S3 Significant co-occurrence and co-exclusion relationships between arc-nosZ and gdhA genes abundances and chemical properties (NH₄⁺, DON). Spearman's rank correlation matrix of genes abundance and chemical properties values. Strong correlations are indicated by large circles, whereas weak correlations are indicated by small circles. The colours of the scale bar denote the nature of the correlation, with 1 indicating a perfectly positive correlation (dark blue) and -1 indicating a perfectly negative correlation (dark red). Only significant correlations (FDR < 0.05) are shown.

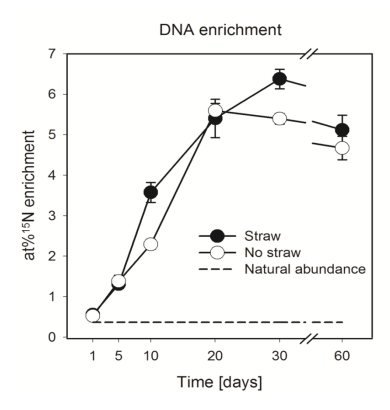


FIG. S4 Development of ¹⁵N enrichment of DNA (at% ¹⁵N) during incubation time and natural abundance of 0.336 at % ¹⁵N of ammonium sulphate (dashed line) (n=3, means with standard errors).