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- 2 MICROBIAL MATS
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- 9 Running head: Denitrification in a coastal microbial mat.
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18 Abstract

Denitrification was measured in three structurally different coastal microbial mats by 19 using the stable isotope technique. The composition of the denitrifying community was 20 21 determined by analyzing the nitrite reductase (nirS and nirK) genes using clone libraries and the GeoChip. The highest potential rate of denitrification  $(7.0\pm1.0 \text{ mmol N m}^{-2} \text{ d}^{-1})$ 22 was observed during summer at station 1 (supra-littoral). The rates of denitrification were 23 24 much lower in the stations 2 (marine) and 3 (intermediate) (respectively  $0.1 \pm 0.05$  and  $0.7\pm0.2$  mmol N m<sup>-2</sup> d<sup>-1</sup>) and showed less seasonality when compared to station 1. The 25 denitrifying community at station 1 was also more diverse than that at station 2 and 3, 26 which were more similar to each other than either of these stations to station 1. In all 27 three stations, the diversity of both *nirS*- and *nirK*-denitrifiers was higher in summer 28 29 when compared to winter. The location along the tidal gradient seems to determine the composition, diversity and activity of the denitrifier community, which may be driven by 30 salinity, nitrate/nitrite and organic carbon. Both *nirS* and *nirK* denitrifiers are equally 31 32 present and therefore they are likely to play a role in the denitrification of the microbial mats studied. 33

34

36 Introduction

Denitrification is a bacterial process during which nitrate or nitrite is stepwise reduced 37 through a few intermediate gaseous nitrogen compounds to dinitrogen (Zumft, 1997). 38 Nitrite reductase is present in all denitrifying bacteria and mediates the reduction of 39 nitrite to nitric oxide and is considered as the key enzyme of denitrification. There are 40 41 two functional equivalent but structurally distinct nitrite reductases known in denitrifying bacteria (Zumft, 1997). These are Cytochrome cd1 (Cd-Nir), encoded by nirS and a 42 copper nitrite reductase (Cu-Nir), encoded by nirK. There are no organisms known that 43 44 carry both genes and therefore these two enzymes are thought to be mutually exclusive (Zumft, 1997). Nitrite reductase genes have been used as functional molecular markers 45 for denitrification in natural environments and have revealed the diversity of denitrifying 46 bacteria in a variety of habitats such as soil (Prieme et al., 2002; Throbäck et al., 2007), 47 estuarine sediments (Santoro et al., 2006), marine sediments (Braker et al., 2000; Hannig 48 et al., 2006), and seawater (Castro-Gonzalez et al., 2005; Jayakumar et al., 2004; Oakley 49 et al., 2007). Environmental factors were identified that shaped the denitrifier community 50 composition (Braker et al., 2000; Hallin et al., 2009). Moreover, the type of habitat 51 52 determined the presence or dominance of *nirS*- and *nirK*- type denitrifiers (Hannig *et al.*, 53 2006; Oakley et al., 2007).

54

Coastal microbial mats are compact, highly structured, small-scale ecosystems (Stal,
2001). These mats are built by cyanobacteria, oxygenic phototrophic bacteria, which
through primary production enrich the sediment with organic matter. This organic matter
forms the basis of a complex, multi-layered microbial ecosystem. Previous studies of

59 nitrogen cycling in microbial mats have focused mainly on the nitrogen fixation and only a few studies documented denitrification in microbial mats. Joye & Paerl (1994) studied 60 denitrification in microbial mats of Tomales Bay (California) and found that it removed 61 only 15% of the  $N_2$  that was fixed on an annual basis in these mats. In a hypersaline 62 microbial mat, denitrification was lower than N<sub>2</sub> fixation in summer, but exceeded N<sub>2</sub> 63 64 fixation in winter when it turned the mat into a sink for nitrogen (Bonin & Michotey, 2006). Desnues et al. (2007) reported spatio-temporal distribution of denitrifying bacteria 65 in a hypersaline microbial mat. These studies focused on the rates of denitrification and 66 67 did not give information on the diversity and the temporal and spatial distribution of the denitrifying bacteria and their activities and therefore provided only an incomplete view 68 69 on this process in microbial mats.

70

We investigated microbial mats that proliferate at the North Sea coast of the Dutch 71 barrier island Schiermonnikoog. Based on morphological, microscopic, and molecular 72 genetic differences we distinguish three major types of microbial mats that develop along 73 the tidal gradient. The bacterial, archaeal and eukaryal community composition and 74 75 microbial diversity were intrinsic of the mat type and depended on the location along the tidal salinity gradient (Bolhuis & Stal, 2011; Bolhuis et al., 2013). Previously it was 76 shown that N<sub>2</sub> fixation and the diazotrophic community varied along the same lines at 77 78 these three stations (Severin & Stal, 2010). The variation of  $N_2$  fixation may be attributed to different environmental conditions in microbial mats, which changes 79 80 spatially (location along the tidal gradient) and temporally (tide, day-night, and seasonal). 81 We expect that the same factors exert also a selective force on the denitrifying

82	community and its activity. At the different positions along the tidal gradient the mats
83	would allow the development of different community compositions, which would also
84	alter the potential rate of denitrification that can be achieved (Philippot & Hallin, 2005).
85	In this study we measured the potential rates of denitrification in the three different mat
86	types during different seasons. Alongside, we identified the denitrifying communities and
87	measured relevant environmental variables in order to elucidate: 1) whether mats along
88	the tidal gradient contain different types of denitrifying bacteria; 2) whether a relationship
89	exists between the denitrifier community and the potential rate of denitrification; 3)
90	which environmental factors affect denitrification and the composition of the denitrifier
91	communities.
92	
93	Material and methods
94	Study area and sampling
95	The study site was located on the North Sea coast of the Dutch barrier island
96	Schiermonnikoog. The geographical locations and descriptions of the three types of
97	microbial mats (stations) that were sampled for this study are given in Table 1. The
98	sample stations were situated along a transect perpendicular to the beach covering the
99	tidal gradient. Sampling was done at four different seasons during 2010 and 2011.
100	Samples were taken from the top 2.5-3 cm of the mat using custom-made transparent
101	Lexan cylinder cores of 50 mm inner diameter and 60 mm height. The cores were
102	transported back to the laboratory within 4 h of sampling and kept at ambient temperature
103	and light. The incubations started within 24 h after sampling. Additional samples were

taken for nucleic acid extraction. These samples were taken from the top 1 cm of the mat

by using a 10 ml truncated syringe as a corer. These mat samples were divided into fourequal parts, put into cryo-vials, and immediately frozen in the field in liquid nitrogen.

107

108 Chemical analyses

For nutrient analyses 5 g mat sample (top 1 cm) was extracted with 40 ml 2 *M* KCl. The
extracts were filtered through Whatman GF/F filters and the filtrates were kept at -20 °C
until analysis (within a month). Nutrient concentrations were measured on an automated
Segmented Flow Analyzer using standard analytical procedures. Other mat samples were
freeze-dried for the determination of total nitrogen (TN), total organic carbon (TOC) and
C/N ratio by EA-IRMS (DELTA V Advantage; Thermo Fisher Scientific, Bremen,
Germany).

116

117 Measurement of potential denitrification

Subsamples of  $1.2 \text{ cm}^2$  (10 mm thickness) of the cores of the mats were placed into 12.5

119 ml Exetainers (Labco Limit, Buckinghamshire, England) by using a 5-ml syringe as a

120 corer. The measurements were carried out according to Thamdrup & Dalsgaard (2002)

121 with some modifications. Briefly, the Exetainers were completely filled with anoxic

artificial seawater (NaCl 20.5 g,  $Na_2SO_4$  3.4 g, KCl 0.58 g, KBr 0.084 g and  $H_3BO_3$ 

123 0.022 g, MgCl<sub>2</sub>.6H<sub>2</sub>O 0.05 mol, CaCl<sub>2</sub>.2H<sub>2</sub>O 0.01 mol in 1000 ml Milli-Q water).

124 Addition of 100  $\mu$ M Na<sup>15</sup>NO<sub>3</sub> (98.5%, <sup>15</sup>N atom%; Sigma-Aldrich, 100  $\mu$ M <sup>15</sup>NH<sub>4</sub>Cl

125 (99.2%, <sup>15</sup>N atom%; Sigma-Aldrich) and 100  $\mu$ M <sup>15</sup>NH<sub>4</sub>Cl +100 $\mu$ M Na<sup>14</sup>NO<sub>3</sub> were

applied to the Exetainers, respectively. Before the addition of the <sup>15</sup>N tracer, the

127 Exetainers were placed in dark for 2 h to allow the depletion of  $NO_x^-$  and any residual

128	oxygen. All Exetainers, including one set without any addition, were incubated for 24 h
129	in the dark at ambient temperature (Table 2). At 4-h intervals during 24 h two replicate
130	Exetainers from each treatment were fixed by injecting 200 $\mu l$ 50% (w/v) ZnCl_2. The
131	Exetainers were stored at 4°C in the dark upside down until analysis (within a week). The
132	isotopic composition of the dinitrogen of He-equilibrated headspace (2 ml in the 12.5-ml
133	Exetainer vials) was determined by an EA-IRMS (DELTA V Advantage; Thermo Fisher
134	Scientific, Bremen, Germany) equipped with a Haysep Q column. The potential rate of
135	denitrification was calculated from the linear production of excess $^{29}\text{N}_2$ and $^{30}\text{N}_2$
136	according to Thamdrup & Dalsgaard (2002).
137	
138	DNA extraction, PCR, cloning and sequencing
139	DNA was extracted using the MoBio UltraCLEAN soil DNA kit (MoBio Laboratories,
140	Inc., Carlsbad, CA, USA) according to the manufacturer's instructions. Fragments of the
141	genes <i>nirS</i> and <i>nirK</i> were amplified using the primer pairs cd3aF-R3cd for <i>nirS</i> and
142	F1aCu-R3Cu for nirK (Throbäck et al., 2004). PCR conditions for the two sets of primer
143	pairs were 2 min at 95°C, 35 cycles of 50 sec 95°C, 50 sec 53°C, and 50 sec at 72°C,
144	followed by a final extension of 10 min at 72°C. PCR products were checked on a 1%
145	agarose gel. PCR products were cloned using the TOPO-TA cloning kit with the pcR2.1
146	vector and TOP10 competent cells (Invitrogen, Carlsbad, CA, USA) following the
147	manufacturer's instructions. Transformants were randomly picked from each clone
148	library and screened by PCR using T3 and T7 vector primers following the recommended
149	PCR conditions (Invitrogen, Carlsbad, CA, USA). Forty-eight clones were randomly
150	selected for sequencing with the T7 vector primer using ABI PRISM 3130 Genetic

151	Analyzer (Applied Biosystems, Foster City, CA, USA). The total number of sequences
152	obtained from each clone library varied (28-70 sequences) due to the variable quality of
153	the sequencing reads. Sequences have been submitted to NCBI (accession numbers
154	KJ738332 - KJ739305).
155	

156 Sequence analysis

157 Sequences were edited, aligned and translated using MEGA 5 (Molecular Evolutionary

158 Genetics Analysis, http://www.megasoftware.net/mega5/mega.html) and manually

159 checked. Neighbor-joining trees were produced and the reliability of the phylogenetic

160 reconstructions was evaluated by bootstrapping (1000 replicates). The program Mothur

161 (http://www.mothur.org/wiki/Main\_Page) was used to calculate the non-parametric

162 richness estimators and the Shannon diversity index and to determine the differences in

nucleic acid sequences. Operational Taxonomic Units (OTUs) were defined by a 5%

164 difference in nucleic acid sequence for the purpose of community analysis. Based on the

165 OTUs from each library, sequence data were transformed into binary data

166 (presence/absence) for community composition analysis.

167

168 GeoChip analysis

169 The GeoChip is a high-throughput functional gene array covering 289 functional gene

170 families involved in the biogeochemical cycling of carbon, nitrogen, phosphorus and

sulfur (He *et al.*, 2010). For the analysis using the GeoChip we used DNA extracted in

triplicate from the three types of microbial mats sampled in July and in November. The

173 DNA was purified using UltraClean 15 DNA purification Kit (MoBio Laboratories, Inc.,

174	Carlsbad, CA, USA) in order to achieve the quality necessary for hybridization on the
175	chip. The DNA quantity was measured by an ND-1000 spectrophotometer (Nanodrop
176	Inc., Wilmington, DE). The procedures for DNA labeling and microarray hybridization
177	followed the previously established protocols (Wu et al., 2006). Briefly, 800 ng of
178	environmental DNA was labeled with fluorescent dye Cy-5 by random priming. The
179	labeled DNA was re-suspended in 50 $\mu$ l hybridization solution [40% formamide, 5 x
180	SSC, 5 $\mu$ g of unlabeled herring sperm DNA (Promega, Madison, WI), and 0.1% SDS]
181	and 2 $\mu$ l universal standard DNA (0.2 pmol $\mu$ l <sup>-1</sup> ) labeled with the fluorescent dye Cy-3
182	(Liang et al., 2010), denatured at 95°C for 5 min and maintained at 50°C until loaded
183	onto the microarray slides. Arrays were hybridized on a MAUI Hybridization Station
184	(Roche, South San Francisco, CA) for 12 h at 42°C. Hybridized microarrays were
185	scanned by a ScanArray Express Microarray scanner (Perkin-Elmer, Wellesley, MA) at
186	95% laser power and 85% photomultiplier tube gain. The resulting images were analyzed
187	by ImaGene with signals processed as SN>2.0 (signal to noise ratio).
188	

189 Statistical analysis

In order to summarize the gene overlap at station and season level, the detected genes from the GeoChip in the three replicates of each station from July and November were deployed as one pool (mean value from the three replicates). The analyses of overlapping genes, unique genes, and diversity indices were performed using an online pipeline (http://ieg.ou.edu/). The proportion of overlapping genes was calculated as the number of overlapping genes divided by the total number of genes detected in both stations. The 196 proportion of unique genes at each station was calculated as the number of unique genes 197 at each station divided by the total number of genes detected at that station. Statistical analyses of the multi-response permutation procedure (MRPP) and canonical 198 199 correspondence analysis (CCA) (see below) were performed based on community data from the clone libraries as well as from the GeoChip data. MRPP using Bray-Curtis 200 distance was used to test for significant differences in community composition. The 201 MRPP A-statistics describes the within and between group relatedness relative to what is 202 expected by chance. A p-value <0.05 and an A-statistics >0.1 is considered as significant 203 204 difference between groups (McCune et al., 2002). To test the relationship between the 205 denitrifier community and the environmental variables, CCA was carried out. The significance of the whole canonical model was tested by 999 permutations. All statistical 206 analyses were carried out in the open source-software R (Team, 2011), using the vegan 207 package (Oksanen, 2011). Stepwise regression was carried out to test the influence of the 208 environmental factors and denitrifier community on potential denitrification rates using 209 210 SigmaPlot (SigmaPlot, Version 12). Denitrifier communities were converted into univariate variables based on the sample scores for the first two CCA axes. 211 212

213 Results

214 Physicochemical characteristics

Table 2 lists the seasonal and annual mean values of the physicochemical parameters of 215 216 the sample sites and the potential denitrification rates in the three mats. The physicochemical parameters fluctuated seasonally and some parameters showed 217 differences between the three mat types. Ammonium concentration was lowest at Station 218 1 (ST1) and highest at ST3, except in April. Nitrate/nitrite concentrations were in the 219 same range at all stations, albeit with slightly higher concentrations in July and April and 220 221 slightly lower concentrations in September and January. Phosphate concentration was highest in April and lowest in September at all stations. TOC and TN were similar at ST2 222 and ST3 but always lowest at ST1. 223

224

225 Potential denitrification rates

<sup>29</sup>N<sub>2</sub> and <sup>30</sup>N<sub>2</sub> were produced at the expected ratio for denitrification given the addition of

227 99.2-atom% enriched  ${}^{15}NO_{3}^{-}$ . Denitrification rates (N<sub>2</sub> production) showed remarkable

differences between the three stations and varied also seasonally. For ST1 (supratidal,

close to the dunes) the potential denitrification rates ranged from  $0.1\pm0.05$  -  $7.0\pm1.0$ 

mmol N m<sup>-2</sup>d<sup>-1</sup> (Table 2). The denitrification rate was highest in July (7.0 $\pm$ 1.0 mmol N m<sup>-</sup>

<sup>2</sup>d<sup>-1</sup>) and much lower (1.6 $\pm$ 0.3 mmol N m<sup>-2</sup>d<sup>-1</sup>) in September. Denitrification was lowest

232  $(0.1\pm0.05 \text{ mmol N m}^{-2}\text{d}^{-1})$  in April. The seasonal trend of denitrification at the littoral site

233 (ST3) was slightly different from that at ST1. At ST3, the highest denitrification rate

234  $(0.7\pm0.2 \text{ mmol N m}^{-2}\text{d}^{-1})$  was in July and was lowest in September  $(0.1\pm0.05 \text{ mmol N m}^{-1})$ 

<sup>2</sup>d<sup>-1</sup>). A higher rate (0.5 $\pm$ 0.2 mmol N m<sup>-2</sup>d<sup>-1</sup>) was again observed during January. Unlike

the other two sites, the highest rate of denitrification at ST2 (low water mark) ( $1.6\pm0.4$ mmol N m<sup>-2</sup>d<sup>-1</sup>) was observed in January and the lowest rate ( $0.1\pm0.05$  mmol N m<sup>-2</sup>d<sup>-1</sup>) occurred in July and September. The annual average denitrification rate was highest at the supra-littoral (near the dunes, ST1) (2.8 mmol N m<sup>-2</sup>d<sup>-1</sup>) and significantly higher (P<0.05) than at the other two stations, which were low and not significantly different from each other (0.5 mmol N m<sup>-2</sup>d<sup>-1</sup> at ST2 and 0.4 mmol N m<sup>-2</sup>d<sup>-1</sup> at ST3).

243 *nirS* and *nirK* diversity and composition

*nirS* and *nirK* sequences from the three stations were analyzed. Combining the clone

libraries, 76 unique *nirS* operational taxonomic units (OTUs) and 74 unique *nirK* OTUs

246 (at a 5% distance cut-off) were retrieved. The richness and diversity estimators showed

247 different *nirS* and *nirK* gene richness at the three stations. *nirS*-denitrifier community was

richest at ST3, while the other two stations showed a similar richness. *nirK*-denitrifier

community was richest at ST1 and poorest at ST3.

250

251 Phylogenetic analyses of deduced amino acid sequences for *nirS* and *nirK* gene fragments

are shown in Fig. 1. The *nirS* sequences clustered into three distinct groups (Fig. 1A).

Group I contained 53%, 36% and 68% of the total sequences from ST1, ST2 and ST3,

respectively. The sequences in this group were closely related to the cultivated denitrifier

255 Marinobacter sp. U31 (CAF25138) as well as to environmental clones from a

hypersaline microbial mat (e.g. CAL69009, CAL69007), an estuarine sediment (e.g.

AEK77712, ABY52470) and from the Baltic Sea (e.g. CAJ87449). Group II contained

258 12%, 64% and 30% sequences from ST1, ST2 and ST3, respectively. Sequences in

Group II clustered closely with those of a variety of cultivated denitrifiers including *Roseobacter denitrificans*, *Paracoccus denitrificans*, and *Silicibacter pomeroyl*. The third
group contained 35% and 2% of the sequences of ST1 and ST3. These sequences were
closely related to *Pseudomonas stutzeri*, *Azospirillum brasilense* and *Ralstonia eutropha*.

*nirK* sequences clustered into four groups (Fig. 1B). Group I contained 2%, 81% and

265 100% of the total *nirK* sequences from ST1, ST2 and ST3, respectively. The sequences

belonging to group I were most closely related to environmental clones from San

267 Francisco Bay estuarine sediment (ADM93883) and from the Arabian Sea oxygen

268 minimum zone (ACT98741). Three OTUs from ST1 fell into Group II, showing the best

269 hit of 87% nucleotide sequence similarity with a sequence retrieved from a Chinese

agricultural soil (HM628810). Group III contained sequences from ST1 (7%; 17%; 68%,

Fig. 1B) and were related to a variety of cultivated denitrifiers, such as *Rhodobacter* 

272 sphaeroides (CCA12211) and Rhodopseudomonas palustris (NP949481). Group IV

comprised only sequences (19%) from ST2. These sequences are closely related to

environmental clones from San Francisco Bay estuarine sediment (ADM93844,

ADM93870) and remotely related to the cultivated *Alcaligenes* sp. (75% similarity and
48% sequence coverage).

277

278 Diversity of *nirS* and *nirK* genes based on GeoChip analysis is summarized in Table 4. A

total of 264 *nirS* sequences showed a hybridization signal in at least one of mat. The

average number of *nirS* sequences (richness) at ST1 (July), ST2 (July), ST3 (July), ST1

281 (January), ST2 (January) and ST3 (January) was 253, 206, 146, 215, 161, and 125,

282	respectively (Table 4). Thirty-three sequences were unique and detected only at one of
283	the stations and during one season. In July, ST1 harbored 26 unique sequences, which
284	was 10.3% (26/253) of the total number detected. ST2 and ST3 harbored 2.4% (5/206)
285	and 0.7% (1/146) unique sequences, respectively. In January, the number of unique
286	sequences in ST1 dropped to 3, which was only 1.4% (3/215) of the total number
287	detected. In January no unique sequences were observed at ST2 and ST3. Pairwise
288	comparison of <i>nirS</i> sequences showed a high number of overlapping <i>nirS</i> sequences
289	between summer and winter as well as between the stations: 81% (ST1 July & January),
290	72% (ST2 July & January), 70% (ST3 July & January), 74-77% (ST1&ST2), 57%
291	(ST1&ST3) and 68-70% (ST2&ST3).
292	
292 293	Similar results were obtained for <i>nirK</i> (Table 4). We detected 264 <i>nirK</i> sequences in the
	Similar results were obtained for <i>nirK</i> (Table 4). We detected 264 <i>nirK</i> sequences in the mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141,
293	
293 294	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141,
293 294 295	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141, 214, 150 and 127 <i>nirK</i> sequences at ST2 (July), ST3 (July) ST1 (January), ST2 (January)
293 294 295 296	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141, 214, 150 and 127 <i>nirK</i> sequences at ST2 (July), ST3 (July) ST1 (January), ST2 (January) and ST3 (January), respectively (Table 4). In July, ST1 harbored 26 unique sequences,
293 294 295 296 297	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141, 214, 150 and 127 <i>nirK</i> sequences at ST2 (July), ST3 (July) ST1 (January), ST2 (January) and ST3 (January), respectively (Table 4). In July, ST1 harbored 26 unique sequences, which was 10.2% (26/256) of the total detected sequences. ST2 and ST3 harbored 1.5%
293 294 295 296 297 298	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141, 214, 150 and 127 <i>nirK</i> sequences at ST2 (July), ST3 (July) ST1 (January), ST2 (January) and ST3 (January), respectively (Table 4). In July, ST1 harbored 26 unique sequences, which was 10.2% (26/256) of the total detected sequences. ST2 and ST3 harbored 1.5% (3/204) and 0.7% (1/141) unique sequences of the total detected number, respectively. In
293 294 295 296 297 298 299	mats. Most <i>nirK</i> sequences were detected in summer at ST1 (256). We detected 204, 141, 214, 150 and 127 <i>nirK</i> sequences at ST2 (July), ST3 (July) ST1 (January), ST2 (January) and ST3 (January), respectively (Table 4). In July, ST1 harbored 26 unique sequences, which was 10.2% (26/256) of the total detected sequences. ST2 and ST3 harbored 1.5% (3/204) and 0.7% (1/141) unique sequences of the total detected number, respectively. In January, only 1 unique sequence was detected both at ST1 and ST2. No unique sequence

The diversity indices for *nirS* and *nirK* were assessed by richness and Shannon-Weaver index (Table 4). At all stations the values for both diversity estimators were higher in summer. The highest richness was observed in summer at ST1 and the lowest value was found in summer and winter at ST3 (p<0.01). The highest abundance for both *nirS* and *nirK* were observed in summer at ST1. The lowest abundance for *nirS* was found at ST3 and for *nirK* was found in summer at ST2 and winter at ST3.

310

Multi-response permutation procedure (MRPP) statistics was carried out to test the 311 312 differences of the composition of the microbial community between the stations based on nirS and nirK OTUs obtained from the clone libraries and the GeoChip. MRPP testing of 313 these two data sets gave consistent results. Distinctly different denitrifier communities 314 were found in ST1 when compared to the other two stations (p < 0.05) from the two data 315 sets. ST2 and ST3 did not contain significantly different communities (p>0.05) when the 316 analysis was based on data from the GeoChip but were significantly different when using 317 the data from the clone libraries (p>0.05). This was the case for both *nirS* and *nirK* (Table 318 5). There were no seasonal differences in the denitrifier communities in any of the 319 320 stations (data not shown). These results were confirmed by CCA analyses (Fig. 2).

321

322 Environmental control of denitrifying mat community and activity

323 No relationship between potential denitrification rates, environmental factors, and the

denitrifier community were revealed based on stepwise regression analysis. Canonical

325 correspondence analysis (CCA) was applied in order to discover patterns in the

326 composition of the denitrifying community. Using CCA we analyzed the *nirS* and *nirK* 

sequence data obtained from the clone libraries and from the GeoChip with the relatedenvironmental factors (Table 2).

330	Figure 2A shows the results of the CCA from the <i>nirS</i> sequences obtained from clone
331	libraries. In the diagram, denitrifiers were distinctly grouped according to sample station.
332	There was no effect of the season. The community composition was significantly
333	correlated with all selected variables in the adapted CCA model (p=0.02) (based on 999
334	permutations test). In the CCA diagram (Fig. 2A), the first two axes explained 71% of
335	relationship between the total <i>nirS</i> containing community and the environmental factors.
336	The first canonical axis explained 42.5% of total variations (p=0.008) and was dominated
337	by the environmental variables TOC (p<0.05), TN, and ammonium concentration
338	(p<0.05). The second canonical axis explained an additional 28.5% of the constrained
339	variations and was dominated by phosphate (p<0.05) and the nitrate+nitrite
340	concentration. The nirS containing community in ST1 was distinctly different from those
341	in ST2 and ST3 along the first canonical axis (Axis 1), while the <i>nirS</i> containing
342	communities in ST2 and ST3 separated along the second canonical axis (Axis 2). ST2
343	and ST3 were influenced by both the first and second canonical axes and positively
344	correlated with TOC (p<0.05), TN, phosphate (p<0.05) and ammonium concentrations
345	(p<0.05), but negatively correlated with nitrate+nitrite concentration (p<0.05). ST1 was
346	primarily influenced by the first canonical axis, reflecting the role of TOC (p<0.05), TN,
347	and ammonium concentrations (p<0.05). For each individual variable, a significant
348	correlation was found between community and TOC ( $r^2=0.89$ , $p=0.003$ ) and TN ( $r^2=0.87$ ,
349	p=0.003), which is indicated by the length of the arrows in the CCA diagram.

351	Figure 2B shows the CCA profiles based on the seasonal nirS community data as
352	obtained from the GeoChip. GeoChip analyses were only performed on the summer and
353	winter samples. Spatially, nirS communities from different stations were separated along
354	the first axis of the CCA diagram. Temporally, nirS communities of the summer samples
355	were separated from those of the winter samples along the second axis. In general, the
356	community composition was significantly correlated with all selected variables in the
357	adapted CCA model (p=0.001) (based on 999 permutations test). Due to
358	multicollinearity, the C:N ratio, salinity and the potential rate of denitrification were
359	removed. Therefore, five environmental factors were selected and are depicted in the
360	diagram. The first two axes explained 47.7% of the relationship between the total nirS
361	community composition and the environment. The first canonical axis explained 33.4%
362	of the total variation (p=0.001). The first axis was dominated by the environmental
363	variables TOC (p<0.001), TN (p<0.001) and ammonium concentration (p<0.001). The
364	second axis explained the rest 14.4% of the total variation ( $p = 0.001$ ) and was dominated
365	by phosphate (p< $0.001$ ) and the nitrate+nitrite concentration (p< $0.001$ ). For ST1, the
366	nirS community was influenced by all factors taken into account and showed negative
367	correlation with TOC, TN and ammonium concentration. The nirS containing community
368	in the summer samples correlated positively with nitrate+nitrite and phosphate
369	concentrations, while the community in the winter samples was negatively correlated
370	with these factors. For ST2, <i>nirS</i> in the summer samples was only influenced by the
371	factors reflected by second axis. The nirS community in the winter samples was
372	influenced by all the selected factors and was positively correlated with TOC, TN and

373	ammonium concentrations but negatively correlated with phosphate (p<0.001) and
374	nitrate+nitrite concentrations. For ST3, the nirS community was positively correlated
375	with TOC, TN and ammonium concentrations. The <i>nirS</i> in the summer samples also
376	showed a positive correlation with phosphate (p<0.001) and nitrate+nitrite concentrations
377	(although this was not significant), while <i>nirS</i> in the winter samples was not strongly
378	influenced by any of the environmental variables on the second axis.
379	
380	Figure 2C and 2D depict the results of the CCA from the <i>nirK</i> containing communities.
381	For both the cloning and GeoChip data, the first two axes explained the community
382	composition better than the observations of the <i>nirS</i> containing community. The axes 1
383	and 2 explained 72.1 and 51.7% of the total variation of the <i>nirK</i> containing community,
384	respectively. The nirS and nirK containing communities responded similarly to spatial
385	and temporal variation of the environmental factors.
386	
387	

389 Discussion

390 The few existing studies on denitrification in microbial mats used the acetylene inhibition technique (AI). The published rates of denitrification in microbial mats ranged from 0 to 391 3.14 mmol N m<sup>-2</sup> d<sup>-1</sup> (Bonin & Michotev, 2006; Desnues *et al.*, 2007; Jove & Paerl, 392 1994). In the present study, we measured potential denitrification rates by the isotope 393 pairing (IP) technique using small cores of the mat from 0.06 to 7.00 mmol N m<sup>-2</sup> d<sup>-1</sup> and, 394 hence, were in the same range. We do realize that such comparisons fall short because of 395 differences that are inherent of the technique as well as different incubations (i.e. intact 396 397 cores versus slurries). E.g. Lohse et al. (1996) concluded that the AI technique underestimated the rate of denitrification by a factor of two when compared to the IP 398 technique. Also Bonin & Michotey (2006) measured denitrification in a microbial mat in 399 the Camargue using both the AI and IP techniques and found that the latter gave 10-fold 400 higher rates. However, the AI technique is not adequate when denitrification depends on 401 nitrification in the sediment, because acetylene blocks the latter. Our measurements did 402 403 not depend on nitrification since we added ample nitrate.

404

A previous study on the same mats revealed that in summer nitrogen fixation was 2.0, 0.5 and 2.1 mmol N m<sup>-2</sup> d<sup>-1</sup> for the stations 1, 2 and 3, respectively (Severin & Stal, 2008). Similar mats on the German Wadden Sea barrier island Mellum fixed 3.2, 0.2 and 1.0 mmol N m<sup>-2</sup> d<sup>-1</sup> for the stations that were representative for those studied here (station 1, 2 and 3, respectively) (Stal *et al.*, 1984). Hence, these values indicate that denitrification and N<sub>2</sub> fixation are in the same range in the coastal mats studied here. Joye & Paerl (1994) measured denitrification by the AI technique and showed that denitrification was only 15% of N<sub>2</sub> fixation on an annual basis in the mats of Tomales Bay. However, given
that denitrification is underestimated by the AI technique, denitrification may have been
responsible for a much higher proportion of the loss of the fixed N<sub>2</sub>. Also, Bonin &
Michotey (2006) found that denitrification exceeded N<sub>2</sub> fixation in winter in the
hypersaline mat in Camargue. Hence, we conclude that denitrification can be an
important sink for the fixed nitrogen in microbial mats.

418

Spatial and temporal heterogeneity of potential denitrification rates were observed in the 419 420 present study and have also been documented for other microbial mats (Bonin & Michotey, 2006; Joye & Paerl, 1994). ST1 and ST3 showed the highest potential 421 denitrification rates in summer (July), which was consistent with what has been reported 422 for the hypersaline mats in the Camargue (Bonin & Michotey, 2006) and for the mudflat 423 mats in Tomales Bay (Joye & Paerl, 1994). These results suggest that the nitrogen cycle 424 in different phototrophic microbial mats behaves in a similar way. This might be due to 425 426 the fact that microbial processes in phototrophic microbial mats are fundamentally the same and driven by the physicochemical gradients typically existing in these ecosystems 427 428 (Stal, 2012).

429

The potential rate of denitrification in each of the mats can most likely be attributed to the
dissimilar denitrifier communities. Denitrifiers are phylogenetically diverse and therefore
it is expected that the physiology and enzyme affinities may vary considerably (Philippot
& Hallin, 2005). Consequently, shifts in community composition would lead to changes

in the potential rate of denitrification and this has actually been shown in several cases

435 (Cavigelli&Robertson, 2000; Jayakumar et al., 2004; Rich et al., 2003).

436 We used GeoChip and clone libraries to investigate the denitrifier community in 437 microbial mats. Clone libraries offer the possibility to discover novel species of denitrifiers (assessed by evaluating and analyzing the *nirS* and *nirK* genes) in the 438 microbial mats. However, the limited numbers of clones that were sequenced and the bias 439 of the PCR approach targeting mainly dominant groups could have underestimated the 440 441 rare types. Therefore we used in addition the GeoChip. This chip provides a high 442 coverage of the *nirS* and *nirK* genes that are not sufficient abundant to be retrieved by clone libraries, provided that their probes were included on the chip. The combination of 443 444 these two approaches allowed us to obtain a comprehensive diversity of the *nirS*- and 445 *nirK*-denitrifiers in the microbial mats. The results from both analyses were in agreement 446 with each other.

447

The phylogenetic analysis of the denitrifier community using *nirS* and *nirK* revealed that 448 denitrifiers inhabited all three types of microbial mats. Most of the *nirS* and *nirK* genes 449 retrieved in this study were unrelated to known denitrifying bacteria but shared 450 considerable phylogenetic similarity with sequences from diverse environments including 451 452 estuarine (Santoro et al., 2006), marine habitats (Castro-Gonzalez et al., 2005) as well as 453 soil (Throbäck et al., 2007) and sludge (Osaka et al., 2006). This suggests that a large 454 number of the denitrifiers in these microbial mats have not yet been cultivated. The high 455 diversity of the denitrifier community may be due to a variety of potential environmental niches present in the microbial mats, which would allow diverse denitrifiers to 456

457 proliferate. The deduced amino acid sequences of *nirS* and *nirK* fragments retrieved from clone libraries made from each of the sample stations were more similar to each other 458 than to those of the other stations and hardly overlapped with the sequences from other 459 stations. Although the *nirS* phylogenetic tree did not show a clear division of the clones 460 according to the stations from which they originated, as was the case for the *nirK* tree, 461 462 CCA analyses confirmed that both *nirS*- and *nirK*-denitrifier communities partitioned according to the different mat types. This shows that the conditions that prevail in a 463 certain mat type selects for the type of denitrifier. This is in agreement with a clone 464 465 library based study of a denitrifier community along a salinity and nitrate gradient in a coastal aquifer. Santoro and coworkers found that both NirS and NirK were distinct for 466 certain communities, exhibiting little overlap between stations (Santoro et al., 2006). This 467 habitat specificity of *nirS*- and *nirK*- denitrifier communities was observed in various 468 other environments such as the Baltic Sea and freshwater lakes (Kim et al., 2011) or soils 469 (Prieme et al., 2002). 470

471

Diversity estimates (Shannon-Weaver) based on the clone library GeoChip indicated that 472 473 ST1 harbored a more diverse *nirK*-type denitrifying community than the other two stations. With respect to *nirS* diversity, estimates made by clone libraries and the 474 GeoChip were not consistent and we are therefore unable to draw a conclusion for this 475 476 gene. The stringency of hybridization was optimized (Wu et al., 2006). The groupspecific probes matched perfectly with their targets and the false positive signal was 477 negligible. Most unique *nirS* and *nirK* were detected at ST1 suggesting that this station is 478 479 different from the other two. Moreover, the phylogenetic trees of translated *nirS* and *nirK*  480 from ST2 and ST3 show that a large number of sequences fell into the same clusters, suggesting that the denitrifying communities of these two stations were similar. The *nirS* 481 and *nirK* sequences from ST1 formed distinct clusters. Multi-response permutation 482 procedure analysis based on clone libraries and GeoChip of *nirS* and *nirK* confirmed that 483 the similarity was higher between ST2 and ST3 than either of these stations to ST1. 484 485 Hence, in this respect the dissimilarity of the denitrifying communities in the three stations did not follow the pattern of the whole microbial community (Bolhuis & Stal, 486 2011). These authors found that the microbial community of ST2 was more dissimilar 487 488 from that of ST1 and ST3. The *nirS*- and *nirK* denitrifiers showed higher diversity during summer and a lower diversity during winter. This is in agreement with the development 489 of a mat, which grows to maturity during summer and growth stops during winter when 490 the mat is degraded (Stal et al., 1985). 491

492

The spatial organization of the denitrifying community in the microbial mats was likely 493 the result of the different environmental conditions. Salinity has been proposed as the 494 major driver of the microbial community composition for these microbial mats (Bolhuis 495 496 et al., 2013). This might also apply to the denitrifier community. Jones & Hallin (2010) concluded that the global distribution pattern of *nirS* and *nirK* genes corresponded to 497 salinity. The lower salinity at ST1 may explain the higher diversity of the denitrifier 498 499 community. This has also been observed in a benthic denitrifier community along the estuarine gradient in Chesapeake Bay (Bulow et al., 2008). These authors found the 500 501 highest *nirS* diversity at a freshwater station and the lowest diversity at a station with 502 high salinity. Nitrite reductase genes in a wastewater treatment system showed that

503 salinity decreased the diversity of both *nirS* and *nirK* containing denitrifying bacteria (Yoshie *et al.*, 2004). Similar observations have been made for other functional genes of 504 the N-cycle. Severin et al. (2012) investigated the same mats as in this study and found 505 506 that the proportion of cyanobacterial *nifH* transcripts decreased with increasing salinity. 507 Likewise, Bernhard et al. (2010) found that the loss of diversity of ammonia-oxidizing 508 bacteria correlated with increasing salinity in the Plum Island Sound estuary. Possible factors for the denitrifier compositional changes were partly associated with but not 509 exclusively driven by salinity. 510

511

Canonical correspondence analysis of the *nirS* and *nirK* genes indicated that organic 512 substrates and nitrate/nitrite are also important environmental factors influencing the 513 denitrifier community composition but in opposite ways. Nitrite is the electron acceptor 514 in denitrification. The nitrate/nitrite concentration at ST1 was slightly higher than at the 515 516 other two stations and this might be the underlying reason for the different denitrifier community in this station (Table 2). This is in line with the observation of Liu et al. 517 (2003) who found that denitrifying communities were similar when the nitrate 518 519 concentrations were at the same level. Organic carbon is the primary electron donor for heterotrophic denitrifiers (Zumft, 1997). We showed that the highest diversity of the 520 denitrifier community was at the station with the lowest concentration of organic matter 521 522 (Table 2). Most of this organic matter is recalcitrant polymeric material (Stal, 2003). We conceive that this would increase the diversity of denitrifiers. When organic matter is 523 524 available, diversity will be low because of strong competition and out-competing of the 525 less adapted species.

527 The two dimensions of CCA explained only part of the total variance of the denitrifier community (Fig. 2). This implies that there are also other factors that contribute to the 528 529 composition of microbial community. For example, interaction and competition for 530 resources with other microorganisms could be additional factors. In the microbial mats, 531 denitrifiers compete for nitrate+nitrite with the primary producers such as cyanobacteria and diatoms that represent dominant groups in these mats (Bolhuis & Stal, 2011; Severin 532 et al., 2010). The diversity of the nifH gene in these mats varied in a similar way (Severin 533 534 & Stal, 2010). These authors found that the diazotrophic communities of ST2 and 3 were more similar to each other than to ST1. These results suggest that regardless different 535 functional genes (*nifH*, *nirS* and *nirK*), the structure of the mats and its position in the 536 littoral gradient overwhelmingly drive the diversity of the community, rather than single 537 geochemical factors. 538

539

540 NirS and NirK nitrite reductases are functionally equivalent, but there is a debate going on as to whether the two types of denitrifiers are ecologically distinct (Jones & Hallin, 541 542 2010). Smith & Ogram (2008) found that *nirS*- and *nirK*- denitrifiers responded differently to environmental gradients. In this study, we found that nirS- and nirK-543 denitrifiers were similarly affected by environmental variables (Fig. 2). However, 544 545 although this does not exclude the possibility that the two types of denitrifiers inhabit different niches, we have also no evidence for the opposite that they occupy the same 546 niche. Desnues et al. (2007) investigated the vertical zonation of nirS- and nirK-547 548 denitrifiers in a hypersaline mat. These authors found that *nirS* was mainly localized in

549 the permanent anoxic layer whereas *nirK* occurred throughout the whole mat and seemed 550 to be better adapted to environmental fluctuations. Shannon index based on *nirS* and *nirK* sequences indicated a higher diversity of *nirS* clones compared to *nirK* clones at station 551 552 3. This finding agrees with observations from another ecosystem (Mosier & Francis, 2010). However, the opposite was found for the stations 1 and 2. The seasonal changes of 553 abundance of nirS and nirK were not consistent and varied between stations. This would 554 imply that *nirS*- and *nirK*-denitrifiers adapt differently to the environment. As illustrated 555 in a previous study (Santoro et al., 2006), caution is needed because comparisons by 556 557 using richness estimates may vary according to sample size. The GeoChip results showed a similar diversity and richness for nirS and nirK. It has been shown that in some 558 ecosystems nirS-denitrifiers were more abundant than nirK-denitrifiers (Mosier & 559 560 Francis, 2010). In our study, *nirS*- and *nirK*-denitrifiers were equally abundant in the GeoChip analysis, suggesting that both types of denitrifiers play important roles in 561 denitrification in the microbial mats. 562 563

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741

- 743 Tables:
- Table 1. The geographical coordinates and description of the mats investigated in this
- 745 study.

Station	Geographical coordinates	Description
Station 1 (ST1)	53°29.445′N, 6°8.718′E	Mainly freshwater influenced site, close to the dunes. Irregularly inundated.
Station 2 (ST2)	53°29.460'N, 6°8.309'E	Seawater influenced site, developing microbial mat. At the low water mark.
Station 3 (ST3)	53°29.445′N, 6°8.342′E	Seawater and freshwater influenced site, located between ST1 and ST2, at the edge of the salt marsh

	July (2010)	September (2010)	January (2010)	April (2011)
Station 1		/		
Temperature				
(°C, sediment)	17	10	0	8
$NH_4^+(\mu mol/l)$	128.9±3.0	83.7±17.3	$191.3 \pm 23.4$	233.1±23.7
NO <sub>X</sub> -(µmol/l)	23.3±4.6	8.4±1.1	9.6±4.6	25.9±3.4
$PO_4^{3-}(\mu mol/l)$	20.1±6.1	3.1±0.5	n.d.	25±3.8
TOC (%)	0.04	0.06	0.04	0.04
TN (%)	0.006	0.01	0.007	0.007
C/N	6.7	6.0	5.7	5.7
Salinity (psu)	18	19	15	17
Denitrification				
$(\text{mmol N m}^{-2}\text{d}^{-1})$	7.0±1.0	1.6±0.3	2.4±0.3	0.1±0.05
Station 2				
Temperature (°C, sediment)	17	10	0	8
$NH_4 + (\mu mol/l)$	587.9±41.2	216.2±69.8	736.8±199.6	486.1±61.3
$NO_{X}(\mu mol/l)$	22.4±14.7	8.7±1.4	6.2±1.1	20.6±4.5
$PO_4^{3}(\mu mol/l)$	19.7±2.0	3.2±0.9	n.d.	58.6±26.3
TOC (%)	0.19±0.01	0.20±0.03	0.20±0.03	0.17±0.02
TN (%)	0.03	0.04	0.03	0.03
C/N	6.3	5.0	6.7	5.7
Salinity (psu)	28	28	30	28
Denitrification	01007	0.1.0.07	1 < 6 /	
$(\text{mmol N m}^{-2}\text{d}^{-1})$	0.1±0.05	0.1±0.05	1.6±0.4	0.2±0.1
Station 3 Temperature				
(°C, sediment)	17	10	0	8
$NH_4^+$ (µmol/l)	217.3±102.3	255.9±68.4	510.2±62.2	475.8±3.5
$NO_{X}$ (µmol/l)	16.0±3.7	7.6±1.9	2.8±0.6	19.6±4.0
$PO_4^{3-}$ (µmol/l)	28.5±15.2	3.9±1.2	n.d.	48.5±10.6
TOC (%)	0.11±0.03	0.15±0.02	0.15±0.02	0.13±0.02
TN (%)	0.02	0.03	0.03	0.02
C/N	5.5	5.0	5.0	6.5
Salinity (psu)	25	25	22	23
Denitrification				
$(\text{mmol N m}^{-2}\text{d}^{-1})$	0.7±0.2	0.1±0.05	0.5±0.2	0.1±0.05

747 Table 2. Physicochemical parameters and potential denitrification rates in the microbial748 mats during the 2010 sampling period.

749 Abbreviations: n.d., no data; TOC, total organic carbon; TN, Total nitrogen.

750	Table 3. Richness and diversity statistics of <i>nirS</i> and <i>nirK</i> clone libraries based on 95%
751	cutoffs.

	No. of clones	No. of OTUs	ACE	Chao1	Shannon	Simpson
nirS						
ST1	55	20	56	42	2.1	0.25
ST2	48	19	131	49	2.2	0.18
ST3	66	32	48	47	3.2	0.03
nirK						
ST1	70	35	481	194	3.3	0.05
ST2	69	27	122	84	2.6	0.18
ST3	52	17	41	30	2.1	0.13

Table 4. Summary of *nirS* and *nirK* genes detected by GeoChip, including the number
and percentage of overlapping (italic) and unique (bold) sequences, the diversity indices
and abundance for each station.

	ST1_July	ST2_July	ST3_July	ST1_Jan.	ST2_Jan.	ST3_Jan.
nirS						
ST1_July	26(10.3%)	200(77.2%)	144(56.5%)	210(81.4%)	159(62.4%)	124(48.8%)
ST2_July		3(1.5%)	145(70.1%)	185(78.4%)	154(72.3%)	123(59.1%)
ST3_July			1(0.7%)	139(62.6%)	123(66.9%)	112(70.4%)
ST1_Jan.				3(1.4%)	161(74.9%)	124(57.4%)
ST2_Jan.					0(0.00%)	116(68.2%)
ST3_Jan.						0(0.00%)
Richness*	253	206	146	215	161	125
Shannon-Weaver (H)	5.5	5.3	5	5.4	5.1	4.8
Abundance (%)**	8.2	7.7	7.4	7.9	7.8	7.4
nirK						
ST1_July	26(10.2%)	200(76.9%)	139(53.9%)	212(82.2%)	147(56.8%)	124(47.9%)
ST2_July		1(0.5%)	139(67.5%)	184(78.6%)	144(68.6%)	126(61.5%)
ST3_July			1(0.7%)	134(60.6%)	117(67.2%)	116(76.3%)
ST1_Jan.				1(0.5%)	145(66.2%)	122(55.7%)
ST2_Jan.					1(0.7%)	116(72.1%)
ST3_Jan.						0(0.00%)
Richness *	256	204	141	214	150	127
Shannon-Weaver (H)	5.5	5.3	4.9	5.4	5	4.8
Abundance (%)**	8.3	7.6	7.3	7.9	7.3	7.6

759 \* richness was determined as probe numbers detected.

\*\*abundance was determined by dividing the hybridization intensity of *nirS* or *nirK* on the GeoChip by the

total signal of all nitrogen cycling genes detected on the array.

762

Spatial differences	A-value (clone library)	A-value (GeoChip)
nirS		
ST1 vs ST2	0.753 (p=0.024)*	0.190 (p=0.002)*
ST1 vs ST3	0.749 (p=0.018)*	0.400 (p=0.001)*
ST2 vs ST3	0.667 (p=0.027)*	0.059 (p=0.108)
irK		
T1 vs ST2	0.604 (p=0.029)*	0.242 (p=0.004)*
ST1 vs ST3	0.721 (p=0.045)*	0.454 (p=0.002)*
ST2 vs ST3	0.501 (p=0.020)*	0.66 (p=0.096)

## Table 5. MRPP A-vales of the denitrifier community composition.

765 \* means p<0.05 (statistical difference between whole *nirS* and *nirK* profiles assessed using multi-response
 766 permutation procedure).

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Legends 769

770

771	Figure 1. Phylogenetic trees for <i>nirS</i> (A) and <i>nirK</i> (B) genes, based on the translated
772	amino acid sequence, constructing by neighbor-joining method in MEGA 5. Sequences
773	from this study were shown as the percentage of environmental clones from each station.
774	Significant bootstrap values (>50) are shown at branch nodes.
775	Figure 2. Canonical correspondence analysis of the denitrifier community composition of
776	mat samples. (A) and (C): analysis based on <i>nirS</i> and <i>nirK</i> clone data and points represent
777	the denitrifier community from seasonal samples at indicated station. (B) and (D):
778	analysis based on nirS and nirK GeoChip data and points represent replicated denitrifier
779	community from summer and winter samples at indicated station (S: summer; W:
780	winter). Arrows represent the relationship between environmental parameters with the
781	denitrifier communities.

782

## 784 Figures:

785 Fig. 1









