

Habitat partitioning of denitrifying bacterial communities carrying *nirS* or *nirK* genes in the stratified water column of Lake Kinneret, Israel

Pilar Junier^{1,2,*}, Ok-Sun Kim², Karl-Paul Witzel², Johannes F. Imhoff³, Ora Hadas⁴

¹École Polytechnique Fédérale de Lausanne, 1015 Lausanne, Switzerland

²Max Planck Institute for Evolutionary Biology, 24306 Plön, Germany

³Leibniz Institute of Marine Sciences, University of Kiel, 24105 Kiel, Germany

⁴Israel Oceanographic and Limnological Research, Kinneret Limnological Laboratory, 14950 Migdal, Israel

ABSTRACT: The community composition of denitrifying bacteria was studied in the stratified water column of Lake Kinneret. The nitrite reductase genes *nirS* and *nirK* were amplified by PCR from water samples taken at 1, 14, 19 and 22 m depth, which represent the epi-, meta- and hypolimnion of the lake. The PCR products were analyzed with terminal restriction fragment length polymorphism (T-RFLP) and clone libraries. The highest diversity of *nirS* denitrifying communities was observed at 1 m depth. According to the T-RFLP profiles and clone libraries of *nirS* products, 2 groups of denitrifiers were common to and dominant in all depths. Deduced protein sequences from one of these groups displayed low identity (77%) with other *nirS* sequences reported in GenBank. Denitrifying bacterial communities with *nirK* were most diverse at 22 m and showed highest similarity to those at 19 m depth. Sequences unrelated to *nirK* dominated the clone libraries from 1 m depth, suggesting that denitrifying bacteria with copper-containing nitrite reductase were less frequent at this depth. The results suggest that microorganisms with *nirK* and those with *nirS* respond differently to the environmental conditions in the stratified water column of Lake Kinneret.

KEY WORDS: *nirS* · *nirK* · T-RFLP · Lake Kinneret

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INTRODUCTION

Biological denitrification is a respiratory process involving electron transport phosphorylation and defined as the enzymatic reduction of nitrogen oxides with the production of nitric oxide (NO), nitrous oxide (N₂O) and dinitrogen (N₂) gases. The final product and the intermediates are released into the atmosphere so that fixed nitrogen is transformed into less bioavailable forms of nitrogen (Knowles 1982).

Denitrifying bacteria are a diverse group of microorganisms defined by their common respiratory physiology. They are facultative anaerobes that can switch from oxygen to nitrogen oxides as terminal electron acceptors (Knowles 1982, Philippot 2002). Two structurally different nitrite reductases that carry out the same reaction are found among denitrifiers. One,

encoded by the *nirK* gene, contains copper (Cu-Nir), and the other, encoded by the *nirS* gene, contains cytochromes *c* and *d*₁ (*cd*₁-Nir) (Zumft 1997). The gene *nirS* is part of a cluster containing several other genes involved in the production of the active enzyme (Zumft 1997, Philippot 2002). The gene *nirK* does not seem to belong to a gene cluster, with the only exception of nitrifying bacteria in which it has been observed as part of an operon (Beaumont et al. 2005). So far, only 1 of these 2 genes is found in any particular denitrifying strain, but different strains of the same species may contain different *nir* genes (Coyne et al. 1989).

Using the nitrite reductase genes as functional markers, the community composition of denitrifying bacteria has been studied in different habitats, including soil (Avrahami et al. 2002, Prieme et al. 2002, Rösch et al. 2002), groundwater (Yan et al. 2003), estuarine sedi-

*Email: pilar.junier@epfl.ch

ment (Nogales et al. 2002), marine sediments (Braker et al. 2000, 2001), aquifers (Santoro et al. 2006), seawater (Jayakumar et al. 2004, Castro-Gonzalez et al. 2005) and the oxygen minimum zone of the Black Sea (Oakley et al. 2007). In marine environments a shift in the community structure of *nirS* denitrifying bacteria occurs along the physical–chemical gradient in the stratified water column (Castro-Gonzalez et al. 2005, Oakley et al. 2007). However, similar studies have never been carried out in stratified freshwater lakes.

Lake Kinneret is a monomictic subtropical freshwater lake located in the northern part of Israel. The water column shows a distinct seasonal pattern of chemical stratification characterized by changes in the concentration of dissolved oxygen, nitrate and sulfide. During the period of mixing (December to March), the concentrations of oxygen and nitrate are high throughout the water column. With the onset of thermal stratification in April, followed by the degradation and decomposition of the massive bloom of the dinoflagellate *Peridinium gatunense* in May, oxygen is consumed and gradually depleted in the hypolimnion so that anoxic conditions arise. At the onset of anaerobiosis, the activity of denitrifying bacteria in the anoxic hypolimnion leads to the use of nitrate. After depletion of the nitrate, sulfate reduction becomes the dominant microbial process in this layer (Hadas & Pinkas 1995, Eckert et al. 2002).

During stratification, the distribution of denitrifying communities in the water column is unknown, but is probably coupled to nitrification in the chemocline. We applied molecular techniques combining PCR amplification, terminal restriction fragment length polymorphism (T-RFLP) and clone library analysis to examine the community composition of denitrifying bacteria at various depths of the stratified water column of Lake Kinneret.

MATERIALS AND METHODS

Site description. Lake Kinneret is situated in the northern part of Israel at 210 m below sea level. The lake is stratified from May to December. All water samples were collected on the same day at Stn A (maximum depth 42 m) in October 2004, during the stable stratification period. The composition of the denitrifying bacterial communities was analyzed at 4 depths of the water column, defined by their chemical characteristics (Table 1): 1 m represented the oxygenated surface mixed layer of the epilimnion, 14 m the boundary between epi- and meta-

limnion (lower dissolved oxygen concentrations and decline in temperature), and 19 and 22 m the lower metalimnion and hypolimnion (anoxic water), respectively.

DNA extraction and PCR amplification of *nirS* and *nirK* genes. Water samples (400 ml) were filtered onto 0.2 µm pore size filters (Supor-200, PALL Life Sciences), and stored at –18°C. DNA was extracted using the UltraClean Soil DNA kit (MoBio) following the manufacturer's guidelines. Concentration and quality of the DNA was checked by electrophoresis in 0.8% agarose gels stained with ethidium bromide. For *nirS* amplification the primers cd3aF (Michotey et al. 2000) and R3cd (Throbäck et al. 2004) were used. For *nirK* the primers F1aCu and 3Rcu (Hallin & Lindgren 1999) were applied. These primer sets were selected according to the results of a recent re-evaluation of primers for amplifying genes involved in denitrification in which these combinations performed the best for cultured strains and environmental samples (Throbäck et al. 2004). PCR reactions were carried out as mentioned elsewhere (Hallin & Lindgren 1999, Throbäck et al. 2004) except that the annealing temperature was increased to 57°C. The expected size of the products was about 410 bp for *nirS* and 470 bp for *nirK*.

T-RFLP. In total 8 samples were analyzed with T-RFLP. PCR products that had been purified with multi-screen plates (Millipore) were diluted 1:100 in autoclaved HPLC water and used as template for reaction with the labeled primers R3Cd-HEX (*nirS*) and F1aCu-FAM (*nirK*). To minimize the effect of PCR drift factors (Polz & Cavanaugh 1998), each amplification was carried out in triplicate. The products were pooled, purified using the Qiaquick PCR Purification Kit (Qiagen) and quantified in the gel using Molecular Analysis software (BioRad). The labeled product (20 to 40 ng) was digested with the restriction enzymes *HaeIII*, *AluI* and *MspI* (New England Biolabs), respectively. Restrictions were carried out in 10 µl with 10 U of enzyme and 1× the indicated buffer. After overnight digestion at 37°C the reactions were incubated for 10 min at 95°C to inactivate the restriction enzymes. DNA was ethanol precipitated (BigDye 3.1v sequenc-

Table 1. Chemical characteristics at the sampling depths. NTU = nephelometer turbidity units, SpCond = electrical conductivity

Depth (m)	Temp (°C)	Dissolved oxygen (% saturation)	Dissolved oxygen (mg l ⁻¹)	SpCond (mS cm ⁻¹)	Turbidity (NTU)	pH	Redox (mV)
1	28.0	92.2	7.4	1.0	1.5	8.8	150.0
14	26.8	23.3	1.9	1.0	0.0	8.7	–66.0
19	26.1	1.6	0.1	1.1	4.2	7.7	–62.0
22	18.2	1.6	0.2	1.1	0.1	7.6	–205.0

ing kit manual, Applied Biosystems), and resuspended in 9 μ l of high-denaturing formamide mixture containing the size standard ROX-500 (Applied Biosystems). Restriction fragments were separated in an ABI 3100 automatic sequencer with the GeneScan36-POP4 default module.

Data analysis. Data were analyzed using GeneScan 3.1 software (Applied Biosystems). Values of peak height were normalized relative to the sample with the lowest fluorescence and the threshold of peak detection set to 50 relative fluorescence units (FU) of the normalized data. After normalization, a table with the relative peak height of each terminal restriction fragment (T-RF) was constructed. Statistical analyses were carried out with Primer 6 software (Primer E) and Statistica 6 (Statsoft), as described elsewhere (Wolsing & Prieme 2004). In the principal component analysis (PCA) the samples were classified based on the presence or absence of T-RFs. Additionally, the environmental variables shown in Table 1 were included as secondary variables in the PCA. These secondary variables were also classified based on the 2 principal components calculated with the presence/absence matrix of T-RFs. Simulation of T-RF profiles of the clones was carried out with the program TRiFLe (P. Junier et al. unpubl.). This program simulates PCR and restriction reactions with those sequences that can be detected with the selected primers, producing a list of T-RFs that can be compared with the experimental T-RFs obtained from the environmental samples.

Cloning. In order to minimize PCR errors, the proof-reading *Pfu* DNA polymerase (Promega) was used to produce the PCR products for cloning with the Zero Blunt PCR Cloning Kit (Invitrogen) according to the manufacturer's guidelines. Products from 3 independent PCRs were combined, concentrated on a Multi-screen plate (Millipore), and agarose gel-purified using the Gel Extraction Kit (Qiagen). Forty-eight clones were picked randomly from each cloning and checked for inserts of the expected size by PCR with the plasmid-specific primers M13f/M13r, and agarose electrophoresis. For screening of these clones, one-shot sequencing with M13f primer was performed using the BigDye terminator v3.1 cycle sequencing kit. Sequences were analyzed with an ABI 3100 Genetic Analyzer (Applied Biosystems) according to the manufacturer's recommendations. The resulting nucleotide sequences were compared with sequences in the GenBank database by using translated BLASTX (Altschul et al. 1997) on the NCBI homepage (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). Clones were grouped according to the BLAST results. From each group 1 clone was selected for sequencing of both strands. These sequences have been deposited in GenBank under the accession numbers EF623493 to EF623527.

Phylogenetic analyses. Phylogenetic analyses were carried out with the software ARB (www.arb-home.de). Databases of *nirS* and *nirK* were created using all sequences available in GenBank. Phylogenetic trees of protein-coding genes were constructed from amino acid sequences with the PHYLIP subroutine in ARB, in which a neighbor-joining algorithm used results of a substitution matrix calculated with the Jones-Taylor-Thornton (JTT) model.

Nitrogen species (NH_4^+ , NO_2^- , NO_3^-) concentration measurements. Lake water samples were filtered through a 0.45 μ m filter (S&S), and then NH_4^+ , NO_2^- and NO_3^- concentrations were measured spectrophotometrically by Standard Methods or with a Flow Injection System (QuikChem 8000 Automated Ion Analyzer, Lachat Instruments).

RESULTS

T-RFLP analysis of *nirS* and *nirK*

Nitrate concentrations were about 25 μ g l^{-1} in the epilimnion (1 and 14 m) and undetectable below 18 m. Ammonium concentrations were low in the epilimnion (~ 25 μ g l^{-1}) and increased in the thermocline/chemocline, reaching values of 500 μ g l^{-1} at 22 m.

The enzymes that produced the highest number of T-RFs in the water samples were *AluI* (13 T-RFs) for *nirS* and *HaeIII* (20 T-RFs) for *nirK* (Figs. 1 & 2).

Differences in the community structure between the samples were analyzed based on the relative fluorescence of each T-RF in the T-RFLP pattern. In the community of denitrifying bacteria with *nirS*, a single T-RF (221 bp; Fig. 1) was found most frequently at all depths. Three additional T-RFs (49, 232 and 405 bp) were also common to all samples. These 4 T-RFs represented between 79 and 99% of the total fluorescence in the samples. The community of denitrifiers containing *nirS* at 1 m was the most diverse and contained several unique T-RFs (76, 131, 204, 221, 274 and 374 bp).

The community structure of denitrifying bacteria carrying *nirK* changed more substantially with depth than those containing *nirS* (Fig. 2). Although the *nirK* T-RFs 173 and 231 bp were common to all samples, each depth was dominated by a different T-RF. At 1 m depth, the dominant fragment was the T-RF 264 bp, which also appeared at 14 and 19 m, but not at 22 m. The T-RF 59 bp was observed at depths of 14, 19 and 22 m, but not at 1 m. At 22 m it represented the major proportion of fluorescence. The community of denitrifiers containing *nirK* was more diverse at 22 m depth and showed several distinctive T-RFs (97, 100, 111, 159 and 162 bp).

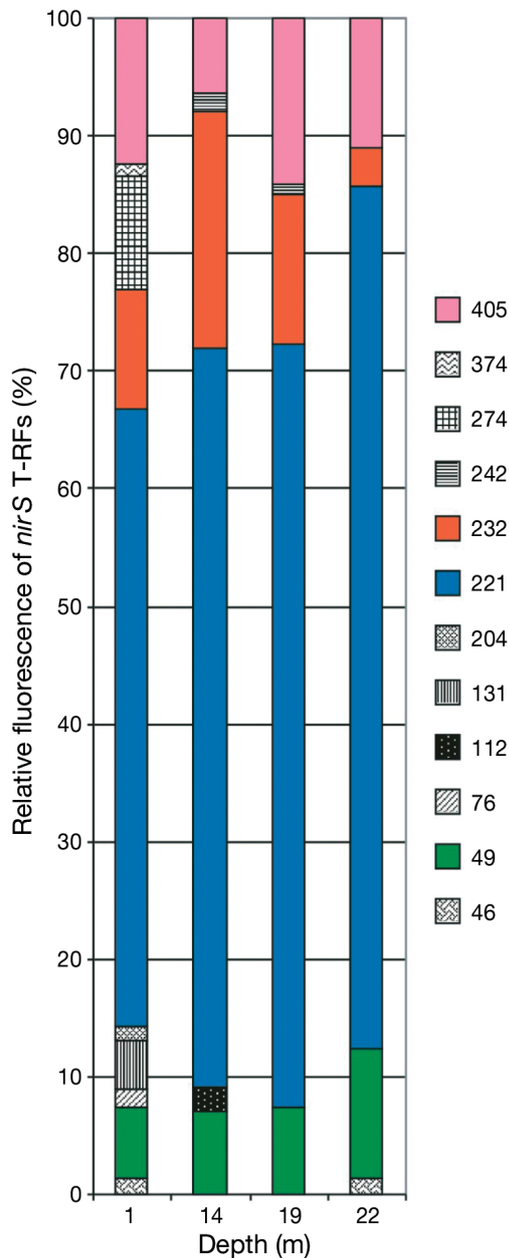


Fig. 1. T-RFLP analysis of *nirS* PCR products digested with *AluI*. T-RF sizes (bp) are shown on the right. T-RFs that were assigned to clones in the libraries are shown in color

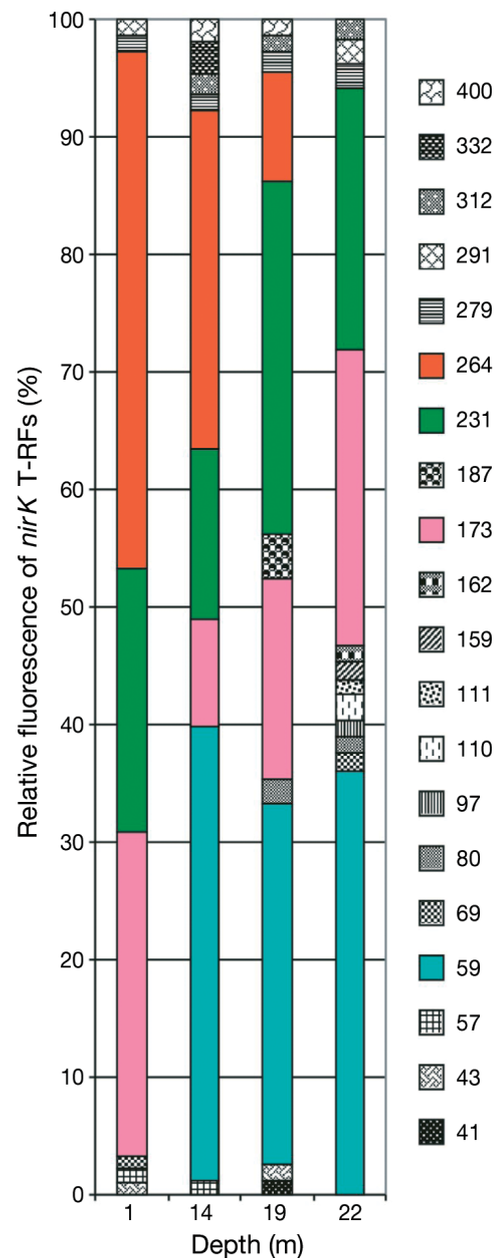


Fig. 2. T-RFLP analysis of *nirK* PCR products digested with *HaeIII*. T-RF sizes (bp) are shown on the right. T-RFs that were assigned to clones in the libraries are shown in color

Statistical analysis of T-RFLP

To determine the underlying community structure, results from restrictions with *HaeIII*, *AluI* and *MspI* were combined and analyzed by multivariate statistical analysis. Three ordination methods were used: PCA, multidimensional scaling (MDS) and cluster analyses of T-RFLP profiles. The results from the different methods were consistent and therefore only results from PCA are shown (Fig. 3).

In the PCA of the T-RFLP of *nirS*, the first principal component (PC1), which accounted for 56% of the variance, separates the 1 m sample from the other depths (Fig. 3A). This separation is explained by the presence of several unique T-RFs that appeared only at the 1 m depth and distinguished this community from the others, e.g. the T-RFs 76, 131, 204, 221, 274 and 374 bp from the digestion with *AluI* (Fig. 1). The separation of the *nirS* denitrifying communities at 19 and 22 m from those at 1 and 14 m coincides with differ-

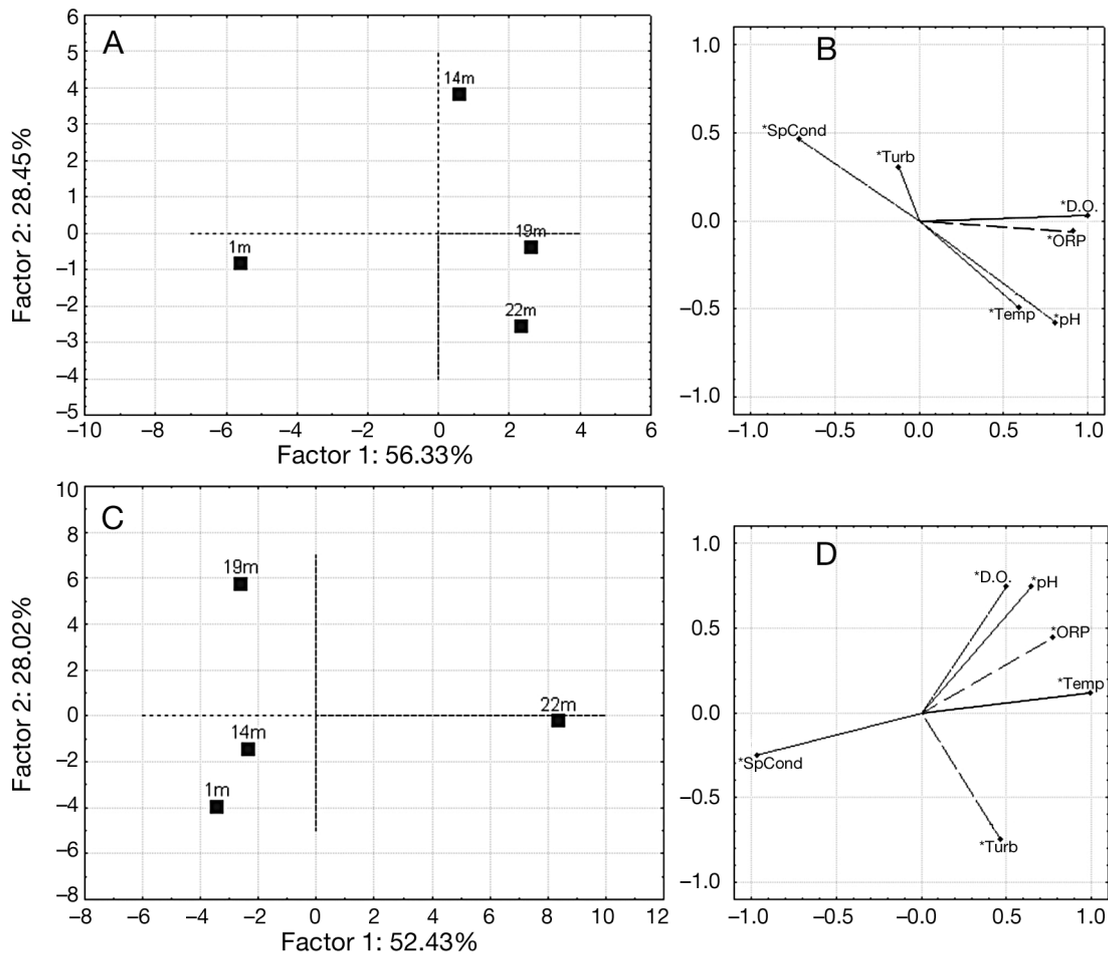


Fig. 3. Principal component analysis (PCA) of T-RFLP patterns of *nirS* and *nirK*. (A) PCA of *nirS*. (B) Separation of environmental variables analyzed as secondary variables in the PCA of *nirS*. (C) PCA of *nirK*. (D) Separation of environmental variables analyzed as secondary variables in the PCA of *nirK*. D.O.: dissolved oxygen; ORP: Redox potential; SpCond: electrical conductivity; Temp.: temperature; Turb.: turbidity

ences in dissolved oxygen, pH, redox and temperature, which are all associated with water stratification (Table 1, Fig. 3B).

For *nirK*, the PC1 (52% of the variance) distinguishes the *nirK* denitrifiers at 22 m from those in the other depths (Fig. 3C). This separation is also due to the presence of various unique T-RFs at 22 m. In the case of *nirK*, temperature was associated with the separation of the sample from the hypolimnion (Fig. 3D).

Cloning and sequence analysis of *nirS* and *nirK*

The samples from 1 and 22 m were chosen for cloning and sequencing because they were differentiated among depths or between genes. Samples from 19 m were also examined because they originated from the boundary between the oxygenated surface water

and the anoxic hypolimnion. Initial screening of the *nirS* and *nirK* clone libraries by one-shot sequencing with the primer M13f and BLAST search showed that the 85 *nirS* sequences could be separated into 9 groups, while the 120 *nirK* clones were separated into 5 groups (Table 2). Matched identity of the inferred protein sequences of these groups with the most similar nitrite reductases in GenBank ranged from 75 to 99% (Table 2), and corresponded in almost all cases (except 1) to uncultured denitrifying bacteria. The only exception was the group IIa of *nirK* that showed 99% matched identity with NirK from *Ochrobactrum* sp. 3CB4.

The library of the PCR products from 1 m prepared with the primers for *nirK* contained more than 90% of clones that did not significantly match with sequences in the GenBank. Similar sequences were also detected at 19 m, but not at 22 m. To characterize these products, specific BLAST tools (tBLAST and PHI-

Table 2. Frequency of clones in the different libraries and closest relative sequences identified by BLASTX in GenBank. nd = not determined, Id = identity (%)

Gene	Group	Clone	No of clones			BLASTX results		Accession no.	Id	Accession no.	
			1 m	19 m	22 m	First hit	First identified hit				
<i>nirS</i>	I	LK1mS-14	2	0	0	Uncultured bacterium	85	AB162262	Uncultured bacterium ^a	76	DQ159631
	II	LK1mS-12	2	0	0	Uncultured bacterium	99	AB162258	<i>Ralstonia eutropha</i> JMP134	83	CP000091
	III	LK19mS-1	0	2	0	Uncultured organism	88	DQ303100	<i>Ralstonia eutropha</i> JMP134	85	CP000091
	IV	LK1mS-7	2	0	0	Uncultured bacterium	86	AY583443	<i>Azoarcus</i> sp. EbN1	77	CR555306
	V	LK1mS-15	1	0	0	Uncultured bacterium	83	AJ440483	<i>Azoarcus</i> sp. EbN1	82	CR555306
	VI	LK1mS-1	5	0	0	Uncultured bacterium	94	AB164086	<i>Dechloromonas aromatica</i> RCB	88	CP000089
	VII	LK1mS-19	8	4	2	Uncultured bacterium	97	AB162301	<i>Dechloromonas aromatica</i> RCB	88	CP000089
	VIII	LK19mS-12	11	26	20	Uncultured organism	74	AF549036	<i>Azoarcus</i> sp. EbN1	73	CR555306
	IX	LK1mS-17	1	0	0	Uncultured bacterium	77	AY195897	<i>Thauera</i> sp. 28	67	AY829012
	<i>nirK</i>	IIf	LK19mK-37	0	4	5	<i>Ochrobactrum</i> sp. 3CB4	99	AY078250	<i>Ochrobactrum</i> sp. 3CB4	99
IIfj		LK22mK-50	0	19	29	Uncultured bacterium	94	AB162319	<i>Pseudomonas</i> sp. G-179	82	AF083948
IVd		LK19mK-25	4	11	14	Uncultured bacterium	83	AB112897	Uncultured bacterium ^a	80	DQ304184
VI		LK22mK-28	0	0	5	Uncultured bacterium	79	DQ304343	<i>Sinorhizobium meliloti</i> 1021	67	AE007256
nd		LK19mK-4	22	3	4	No significant matches					

^aNo identified organisms among the first 100 hits in BLASTX

BLAST) were used. After this analysis, the deduced protein sequence from the clones was only 40 % similar to the histidyl-tRNA synthetase from *Rhodospirellula baltica*.

In the phylogenetic analysis, the *nirS* sequences from Lake Kinneret were assigned to 9 groups (Fig. 4), having as closest relatives sequences from soil or activated sludge, rather than those from other aquatic environments. Group VIII, which contained the majority of the clones from all depths (Table 2), was distantly related to a clonal sequence from nitrate- and uranium-contaminated groundwater (Yan et al. 2003). The clusters VI and VII, which were also frequent especially in the library from 1 m depth, were related to sequences from nitrate- and uranium-contaminated groundwater and *Dechloromonas* spp. With the exception of group III, which was specific for the library from 19 m, all the other clusters were detected at 1 m depth only (Table 2).

In the phylogenetic analysis of *nirK*, the clones were assigned to 4 groups (Fig. 5). The subgroup IIf dominated the libraries from 19 and 22 m depth, and had as closest relatives *nirK* sequences from uncultured soil bacteria. This subgroup, as well as IIf, belonged to a cluster related to sequences from different species of *Ochrobactrum* and other cultured denitrifiers. The group IVd, which was found in both libraries (Table 2), was related to *nirK* from soil and *Azospirillum* spp. Group IVd formed a subcluster inside cluster IV defined by Prieme et al. (2002). Sequences from group VI were observed only in the library from 22 m, and were distantly related to clonal sequences from soil contaminated with silver.

Sequences that did not have a significant match in GenBank (Table 2) were also included in the phylogenetic analysis. These sequences formed the cluster 'nd' (not determined), which is unrelated to the other clonal sequences from Lake Kinneret.

Assignment of *nirS* and *nirK* clones to T-RFs

To correlate the T-RFs observed in the samples with the sequences in the clone libraries, the T-RFLP data were simulated with the *nirS* and *nirK* clonal sequences (Table 3).

Studies with other genes have shown that differences between theoretical and experimental T-RFs,

Fig. 4 (next 2 pages). Phylogeny of partial *nirS* sequences. Accession numbers of reference sequences are given. Sequences from this study are shown in bold. The number of identical sequences is indicated in parentheses. Bootstrap values >85% are indicated. Numbers inside the boxes represent the number of sequences included in the group

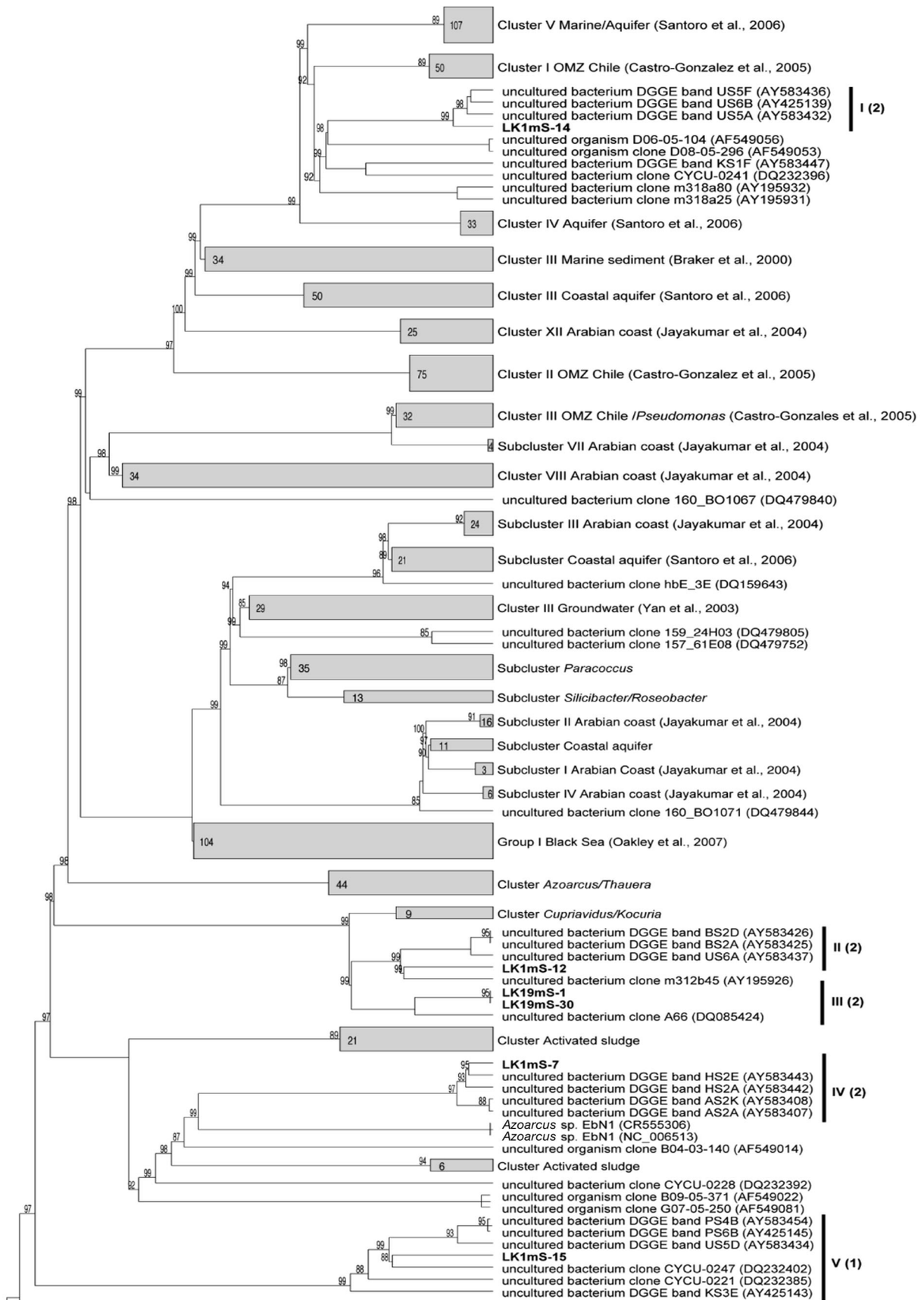


Fig. 4 (continued on next page)

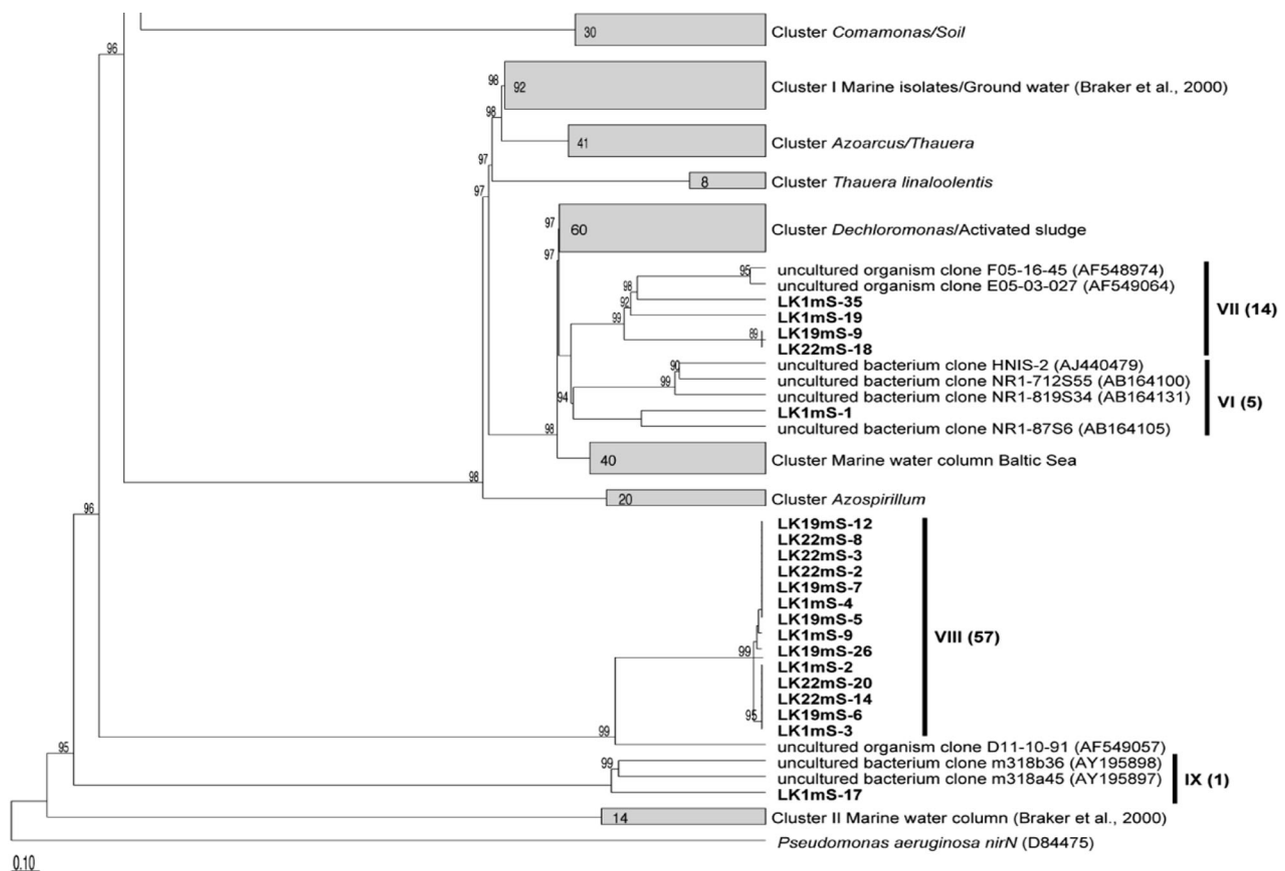


Fig. 4 (continued)

also called T-RF drift (Kitts 2001, Kaplan & Kitts 2003, Lueders & Friedrich 2003), do exist. In most of our experiments, this drift was $\leq 1\%$ of the size of the T-RF (data not shown). Therefore a tolerance limit of 1% of the size of the fragment was chosen for comparison between simulated and experimental T-RFs, meaning that an experimental T-RF of 100 bp was assigned to a theoretical T-RF of 99 to 101 bp (1 bp of tolerance).

Despite the fact that *AluI* produced the highest number of fragments in the T-RFLP analysis of the *nirS* samples, *AluI* had a very low resolution for discriminating groups of clones in the libraries (Table 3). Considering the T-RF drift, the dominant T-RF observed in the samples (T-RF 221 bp, Fig. 1) can be assigned to several groups (I, II, III, V and VIII with a T-RF 224 bp). Other T-RFs that could be assigned were 49 bp (group IV), 232 bp (groups VII and VI) and 405 bp (group IX).

The assignment of the simulated T-RF was clearer for *nirK* than for *nirS*. Based on the results of the most polymorphic enzyme (*HaeIII*), the T-RF 59 bp can be assigned to group IIj, the T-RF 173 bp to group IVd, and the T-RF 231 bp to group IIf. The T-RF 264 bp, which dominates in the profiles and the clone library from 1 m, can be linked to the group of non-*nirK* sequences.

DISCUSSION

In the present study, denitrifying bacterial communities were analyzed in water samples from 4 depths of the stratified water column of Lake Kinneret. While *nirS* and *nirK* sequences from freshwater environments are scarce in the public databases, both types of nitrite reductase genes were detected in all depths. The simultaneous amplification of *nirS* and *nirK* has been difficult in environments such as soil and sediments (Braker et al. 2000, Avrahami et al. 2002, Prieme et al. 2002, Wolsing & Prieme 2004). In water samples, however, this seem to be possible (Oakley et al. 2007).

The simultaneous detection of both genes in the present study is an opportunity to compare the structure of denitrifying bacterial communities carrying *nirS* or *nirK* and their habitat preferences in this stratified system. Virtually nothing is known about the environmental preferences of denitrifying bacteria (Prieme et al. 2002). According to our results from T-RFLP and clone libraries, the diversity of denitrifying communities carrying *nirS* or *nirK* varies at different depths of the water column of Lake Kinneret. Oxygen, nitrate (Liu et al. 2003, Castro-Gonzalez et al. 2005), dissolved organic carbon, inor-

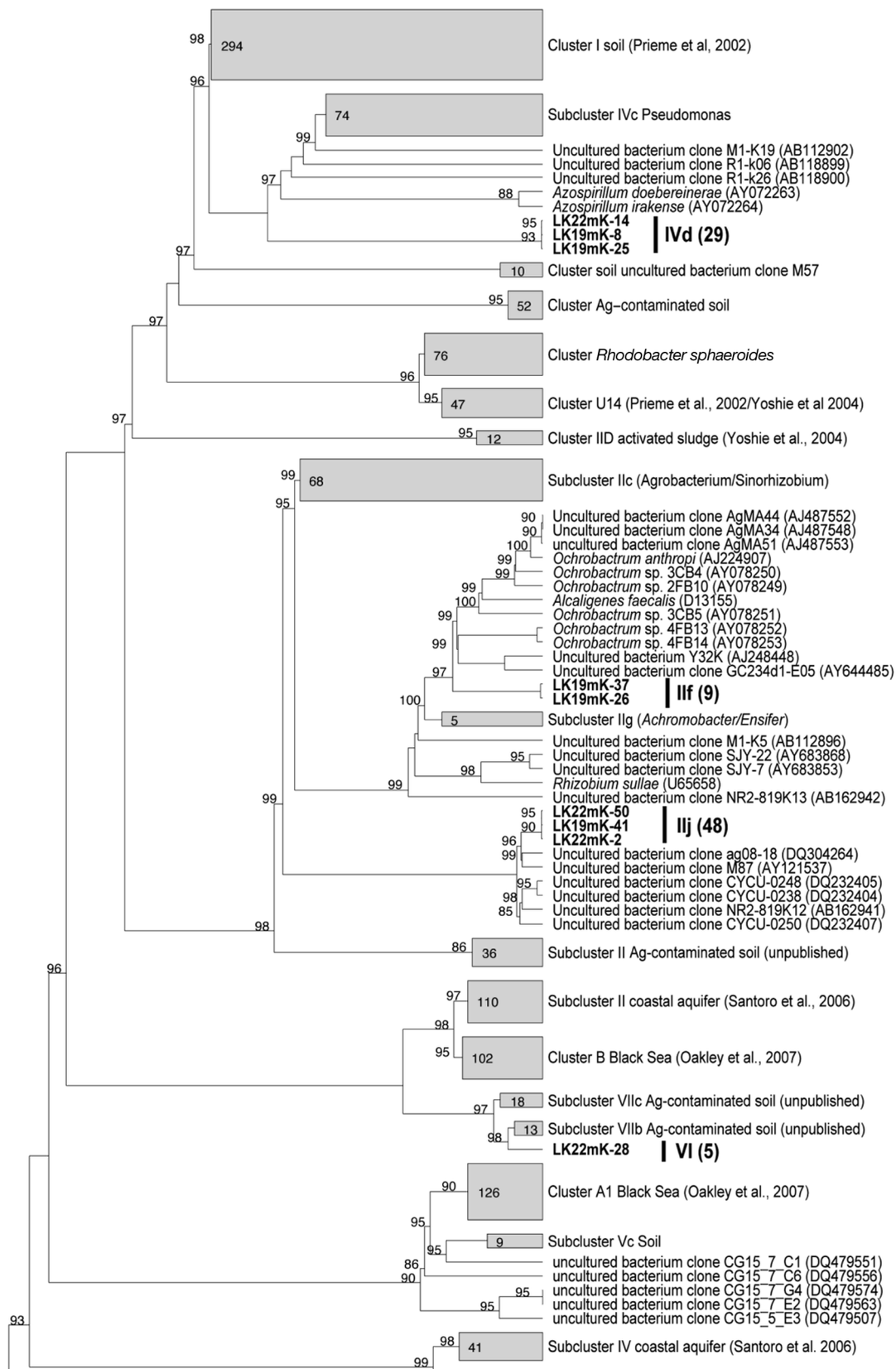
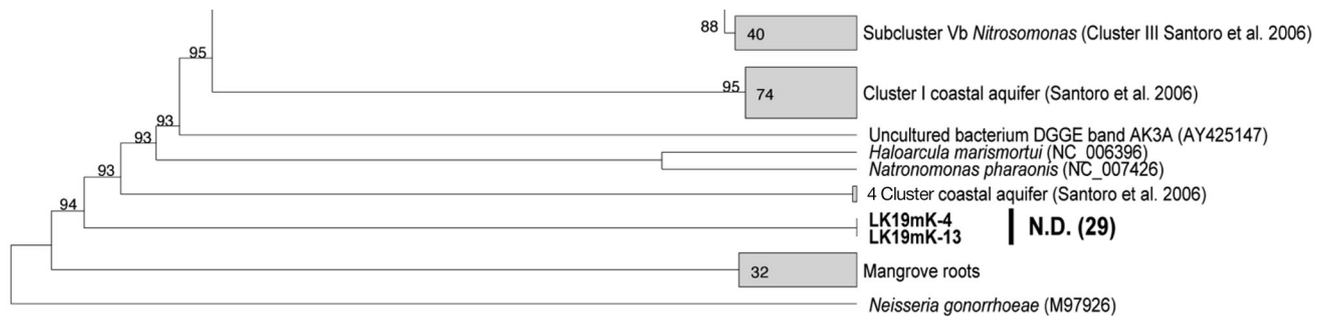


Fig. 5 (this page and next page). Phylogeny of partial *nirK* sequences. For explanation see Fig. 4 legend



0.10

Fig. 5 (continued)

Table 3. Theoretical terminal restriction fragments (T-RFs) calculated for each group of clones from Table 2. nd = not determined

Gene	Group	Clone	Theoretical T-RFs		
			<i>MspI</i>	<i>AluI</i>	<i>HaeIII</i>
<i>nirS</i>	I	LK1mS-14	89	224	42
	II	LK1mS-12	146	224	43
	III	LK19mS-1	133	224	70
	IV	LK1mS-7	130	50	42
	V	LK1mS-15	133	224	70
	VI	LK1mS-1	32	236	144
	VII	LK1mS-19	31	236	151
	VIII	LK19mS-12	89	224	88
	IX	LK1mS-17	38	410	70
<i>nirK</i>	IIf	LK19mK-37	137	473	235
	IIj	LK22mK-50	137	473	64
	IVd	LK19mK-25	136	473	176
	VI	LK22mK-28	297	473	23
	nd	LK19mK-4	82	24	269

ganic nitrogen and salinity gradients (Taroncher-Oldenburg et al. 2003) have been identified as influencing the composition of denitrifying bacterial communities along physical–chemical gradients. In our study, differentiation of the communities in the meta- and hypolimnion (19 and 22 m in depth) was related to dissolved oxygen concentration, pH, and temperature, parameters that are directly linked to stratification. However, the differentiation of the communities in the epilimnion, in which a higher diversity of denitrifiers carrying *nirS* was found, was not directly related to any of the environmental variables measured (Table 1).

It is well established that trace metals, particularly iron (NirS) and copper (NirK) (Zumft 1997), are necessary for the activity of nitrite reductases. In Lake Kinneret most of the bioavailable iron occurs in particles that are restricted to the epilimnion during stratification (Shaked et al. 2004). It can be hypothesized that differences in bioavailability of iron and other trace metals might favor denitrifying bacterial communities with *nirS* in the epilimnion of the lake.

Considering the physiological diversity of the denitrifying bacteria, it is possible that the factors affecting the distribution of the different groups observed at specific depths are not at all correlated with their denitrifying capabilities. Other variables such as the source of carbon or the general tolerance to oxygen may better explain the diversity of denitrifying bacteria carrying *nirS* in the epilimnion, but this needs to be verified in further studies. The bacterial distribution may be influenced by the nature of the microniches with low oxygen concentrations, which may exist in larger aggregates or detritus particles, in the otherwise oxic epilimnion.

According to the T-RFLP analysis and clone libraries, denitrifying bacterial communities carrying *nirS* were dominated by organisms that were common in all the depths, but that seem to be able to exploit different environmental niches. In contrast, only one cluster of *nirK* sequences was common to all the depths; this cluster did not correspond to the dominant group. In Lake Kinneret, bacteria carrying *nirK* appear to be adapted to the specific environmental conditions in each layer of the stratified water column. A similar tendency was observed in a salinity gradient in a beach aquifer at Huntington Beach (California) where different clades of *nirK* sequences were associated to specific salinities, whereas *nirS* did not show any specific environmental adaptation (Santoro et al. 2006). Also, a recent study in the suboxic zone of the Black Sea (Oakley et al. 2007) showed much greater variation of *nirK* sequences at the bottom of the suboxic zone, while *nirS* sequences were more homogeneously distributed.

The *nirS* and *nirK* sequences from Lake Kinneret differed from those of marine habitats and coastal aquifers (Jayakumar et al. 2004, Castro-Gonzalez et al. 2005, Francis et al. 2005, Santoro et al. 2006). This suggests a differentiation of marine and purely freshwater denitrifying bacteria carrying *nirS*, but has to be substantiated by additional studies. Group VIII, which contained most of the clones of this study and dominated in the libraries from all depths, seems to be a

hitherto undescribed group of denitrifiers specific for Lake Kinneret.

The topology of the *nirK* phylogeny obtained in our study resembled those from previous studies (Casciotti & Ward 2001, Avrahami et al. 2002, Prieme et al. 2002). Sequences from Lake Kinneret formed 3 new subclusters inside the clusters II and IV defined by Prieme et al. (2002). The new cluster VI appears to be specific for Lake Kinneret. The closest relatives of most *nirK* clonal sequences from Lake Kinneret were uncultured clones from agricultural soil (Avrahami et al. 2002) or marsh soil (Prieme et al. 2002). Although this situation might suggest a terrestrial origin of the denitrifying bacteria carrying *nirK* in the lake, it might also be due to the low number of *nirK* sequences from aquatic environments that are available for comparison.

Targeting the *nirK* and *nirS* genes would detect denitrifiers irrespective of their activity (Braker et al. 1998). Thus, the data obtained in the present study represent a genetic characterization that constitutes a framework for a more detailed study of the function and activity of denitrifying communities in Lake Kinneret. It also constitutes one of the first analyses of this kind in the water column of freshwater lakes, where community composition and diversity of denitrifying bacteria have previously been unknown.

Acknowledgements. This research was supported by German-Israel Foundation (GIF) grant no. I-711-83.8/2001 and Binational Science Foundation (BSF) grant no. 2002-206. Samples were taken during the German Israeli Minerva School in October 2004. We thank the Yigal Allon Kinneret Limnological Laboratory personnel for their assistance during the sampling. We thank the Max Planck Society and the GIF for financial support of P. Junier and O. S. Kim during this study.

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*Editorial responsibility: Samantha Joye,
Athens, Georgia, USA*

*Submitted: May 8, 2007; Accepted: February 12, 2008
Proofs received from author(s): April 28, 2008*