Composition and temporal dynamics of planktonic archaeal assemblages from anaerobic sulfurous environments studied by 16S rDNA denaturing gradient gel electrophoresis and sequencing

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ABSTRACT: The planktonic archaeal assemblages of several anaerobic, sulfide-rich, aquatic environments were analyzed in space and time by PCR-denaturing gradient gel electrophoresis and sequencing of 16S rRNA gene fragments. The systems were sampled in different years between 1992 and 1998. PCR products were obtained directly from the original DNA without previous nested amplification and yielded successful fingerprints with mostly sharp bands in the gel. Nineteen samples from the anaerobic hypolimnia of 8 lakes and 1 coastal lagoon in NE Spain, Mallorca and Switzerland were compared and a temporal survey was carried out in one of the lakes (Lake Vilar). Between 4 and 14 well-defined bands appeared. All the sequenced bands belonged to Archaea. Although most of the water bodies shared the same climatic conditions and presence of sulfide, the limnological parameters were different among them and different fingerprints were observed in different lakes. Euryarchaeota, i.e., methanogen- and thermoplasma-related sequences, appeared in all the samples but crenarchaeota were recovered only from Lake Vilar. A temporal shift in the predominant members of the archaeal assemblage from crenarchaeota to members of the cluster of thermoplasmal and relatives took place in Lake Vilar between February and June. Sequences related to thermoplasmal and crenarchaeota were distantly related to cultured strains (81% similarity in 16S rDNA) and clustered with branches represented only by environmental clones, whereas sequences related to methanogens grouped with a sequence from an endosymbiont of 1 anaerobic ciliate. A new branch of freshwater euryarchaeota appeared within the cluster of thermoplasmal and relatives. Our study indicates the presence of dynamic archaeal populations in the water column of nonthermophilic, sulfide-rich environments, further extending the diversity and distribution of Archaea in nature. The temporal shift in community composition in Lake Vilar suggests that Archaea grow under in situ conditions. If this is the case, Archaea would be active players in the anaerobic biogeochemical cycles of these environments.

KEY WORDS: Archaea · DGGE · Fingerprinting · 16S rDNA · Sulfurous lakes · Diversity

INTRODUCTION

Archaea illustrate better than any other group of organisms how dramatically our perception of the liv-
tats with high temperature, extreme values of pH, high salinity or anaerobic environments allowing methano genesis. Thus, the genetic and metabolic diversity of Archaea and their ecological distribution seemed more limited than those of Bacteria. Uncultured Archaea, however, extend from marine planktonic environments to soils and sediments (see a review in DeLong et al. 2000), showing up also in close association with animals (Preston et al. 1996, Van der Maarel et al. 1998) and bacteria (Boetius et al. 2000). Therefore, the expected metabolic capabilities of Archaea have increased greatly. Recently, a combination of in situ hybridization and microautoradiography has shown that marine Archaea are active and able to take up amino acids (Ouverney & Fuhrman 2000).

During the last few years, more and more sequences have been deposited in databases and new groups of uncultured Archaea have been described (DeLong 1992, Fuhrman et al. 1992, Barns et al. 1996, Jurgens et al. 1997, 2000, Buckley et al. 1998b). These groups are the marine and freshwater sediments cluster (Crump & Baross 1999), the terrestrial cluster (Buckley et al. 1998) and some cosmopolitan marine clusters (Murray et al. 1998, 1999, Pernthaler et al. 1998, Crump & Baross 2000, Cytryn et al. 2000, Karner et al. 2001). Such studies provide insight into changes in community structure and population dynamics and yield information useful to investigate their functional roles. In Archaea most of these studies have concentrated on marine and terrestrial samples, in a hypersaline lake or in sediments. Only a handful of studies have been carried out in the water column of freshwater habitats: 2 meromictic lakes (Ovreås et al. 1996, Casamitjana & Roget 1999), a high mountain lake (Crump et al. 2000b) and some boreal forest lakes (Jurgens et al. 2000). Whether Archaea are a common component of freshwater plankton is still uncertain.

In the present work, we have carried out a comparative survey on the composition of the archaeoplankton from several anaerobic sulfuriferous water bodies and a temporal study centered on one of them. We used denaturing gradient gel electrophoresis (DGGE) fingerprinting of 16S rRNA genes, PCR-amplified directly from the original DNA (Muyzer et al. 1998). This technique allows extensive comparison on a spatio-temporal scale. In addition, bands were excised from the gel and sequenced, and the identity of the populations was investigated. DGGE has been successfully applied to Bacteria but, to our knowledge, direct DGGE analysis of Archaea has been shown in only 1 recent study (Cytryn et al. 2000) besides our own work (Casamayor et al. 2000b). Sequence analysis of the 600 bp 16S rDNA fragments indicated the presence of a new lineage within euryarchaeota. For the first time, we describe temporal changes in the composition of freshwater archaeal assemblages involving a shift between distantly related phylogenetic groups.

MATERIALS AND METHODS

Environments, sampling and analysis. The systems sampled are located in karstic regions of northeastern Spain (Banyoles and Cuenca), in the coastal area of Mallorca (El Cibollar, Balearic Islands), and in the Swiss Alps (Lake Cadagno) and were sampled in different years between 1992 and 1998. Basins III, IV and VI from Lake Banyoles, Lake Vilar and Lake Cisó are in the Banyoles karstic area (42° 8’ N, 2° 45’ E) and the microbial communities inhabiting these systems have been extensively studied by both conventional (for reviews see Guerrero et al. 1987, Pedros-Aliò & Guerrero 1993) and molecular methods (Casamayor et al. 2000a, b). Lake Banyoles, covering a surface area of 1.1 km2, is a polje comprising 6 main basins. The 3 basins studied are located in the northern area of the lake. Basins III and IV are meromictic with 25 and 18 m maximal depth, respectively, and incoming water seeps through bottom springs (Casamitjana & Roget 1993). The chemocline oscillates between 16 and 21 m in Basin III and between 12 and 17 m in Basin IV depending on the season. Basin VI is 17 m deep and holomictic, with a thermocline situated around 12 to 13 m, and does not have bottom seepage. Blooms of the photosynthetic bacteria *Chorobium phaeobacteroides* and *Thiocystis minor* (formerly *Chromatium minus*) have been described in the 3 basins (Garcia-Gil et al. 1996). Lake Vilar is a meromictic lake formed by 2 basins with a maximum depth of 9 m and a surface area of 11 000 m2, and is connected to Lake Banyoles through a small and shallow channel. High sulfide concentrations are found throughout the year, although sulfide is restricted to the deeper, high-conductivity waters. A stable chemocline exists at 4.5 m, where dense populations of photosynthetic sulfur bacteria develop. Lake Cisó is a small holomictic lake (650 m3), 1 km away from the former, with a maximum depth of 6.5 m. The thermocline is at 1.5 m, where dense populations of photosynthetic sulfur bacteria develop. The lake becomes anoxic during winter holomixis (complete mixing) and high sulfide concentrations (up to 500 µM) are present in the entire water column (Pedros-Aliò & Guerrero 1993). Lake El Tobar and Lake La Cruz belong to the Cuenca karstic system (39° 59’ N, 1° 52’ W). Biological and physico-chemical properties of these lakes have been well described (Miracle et al. 1992 and references therein, Casamayor...
et al. 2000a). Lake El Tobar is meromictic, 19.5 m in depth, and permanently anoxic saline waters (consisting mostly of a sodium chloride brine) starting at about 12 m depth, where blooms of photosynthetic bacteria develop (García-Gil et al. 1999). Lake La Cruz is a meromictic freshwater lake, 24 m deep, with bottom waters enriched in calcium, magnesium and iron bicarbonates, and low amounts of sulfide; a thermocline exists at 8 to 12 m and gas-vacuolate species of photosynthetic bacteria *Pelodictyon* and *Amoebobacter* develop there.

El Cibollar is a small coastal lagoon on the northeast of the S‘Albufera of Mallorca, 750 m from Alcudia Bay (39° 50' N, 3° 10' E), for which some limnological studies have been carried out (Moyà et al. 1987, Borrego et al. 1997). The lagoon has a surface area of almost 4 ha, and average depth is 3.3 m and maximal depth 8.7 m. The oxygen-sulfide interface is located at 3 to 4 m with maximum H$_2$S concentrations up to 2000 µM. Finally, Lake Cadagno (46° 33' N, 8° 43' E) is a permanently stratified, sulfide-rich meromictic lake (H$_2$S concentrations around 1000 µM), located at 1923 m above sea level in the Piora valley (Southern Switzerland). It is about 850 m long and 420 m wide, with a maximum depth of 21 m and a permanent chemocline found between 9 and 14 m. The lower layer is rich in dissolved salts brought by sulfurous sublacustrine springs. Several biogeochemical and molecular studies have been carried out in this lake (Wagener et al. 1990, Putschew et al. 1996, Schanz et al. 1998, Bosshard et al. 2000, Fritz & Bachofen 2000, Tonolla et al. 2000), reporting blooms of the purple sulfur bacteria *Chromatium okenii* and *Amoebobacter purpureus* at the chemocline.

For most of the samples, depth profiles of water temperature, conductivity and oxygen concentration were measured *in situ* using a portable multisensor probe (Hydrolab DS-3; Hydrolab Instruments, TX, USA). Sampling depths were selected according to physicochemical parameters. Water samples were taken using a PVC cone connected through tubing to a battery-driven pump as described elsewhere (Miracle et al. 1992). Sulfide was measured with the methylene blue colorimetric method in 10 ml subsamples fixed quickly by addition of 100 µl of 10 N NaOH and Zn-acetate to a final concentration of 0.1 M (Golterman et al. 1978). Cells were counted in samples fixed with 4% formaldehyde (v/v) and stained with DAPI (Porter & Feig 1980) using epifluorescence microscopy. For DNA analysis, plastic containers were filled with 1 to 5 l of water and were kept in the dark on ice until processed in the laboratory a few hours later. Sampling depths for molecular studies are indicated in Table 1. Samples were concentrated using a refrigerated centrifuge at 8000 × g (Sorvall Instruments, DuPont, DE, USA) and cell pellets were kept frozen at –70°C until further use. Microscopic observation of the supernatants showed that only between 1 and 3% of the cells were not recovered by this method.

Table 1. Temperature, conductivity, sulfide concentration and DAPI counts of prokaryotes (values × 10$^6$ cells ml$^{-1}$) for the depths sampled for molecular analysis. The depth of the oxic-anoxic interface in each lake is also indicated. ND: not determined.

*Sample from the oxic-anoxic interface (presence of 1.4 mg l$^{-1}$ oxygen)

<table>
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<th>Sample</th>
<th>Date (d.m.o.yr)</th>
<th>Interface (m)</th>
<th>Depth (m)</th>
<th>Temperature (°C)</th>
<th>Conductivity (µS cm$^{-2}$)</th>
<th>pH</th>
<th>H$_2$S (µM)</th>
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DNA extraction and analysis. DNA was extracted with hot SDS-phenol and purified with phenol-chloroform-isooamylalcohol (25:24:1 v/v) followed by ethanol precipitation as reported elsewhere (Casamayor et al. 2000b). Fragments of the 16S rDNA suitable for subsequent DGGE analysis were obtained with the primer combination ARC344F (ACG GGG YGC AGC AGG CGC GA) (Stahl & Amann 1991) and ARC915R (GTG CTC CCC CGC CAA TTC CT) (Raskin et al. 1994). A 40 nucleotide GC-rich sequence was attached to the 5' end of the forward primer according to Muyzer et al. (1998). Annealing temperature was 61°C. PCR conditions and DGGE were as described before (Casamayor et al. 2000b), running for 3.5 h at a constant voltage of 200 V and at 60°C in a 20 to 80% vertical denaturant gradient (100% denaturant agent is 7 M urea and 40% deionized formamide). After electrophoresis, the gels were stained with ethidium bromide and photographed with UV transillumination (314 nm) with a Polaroid camera. The photographs were scanned, and the digitized images were processed with the NIH Image software (National Institutes of Health, Bethesda, MD, USA) to measure relative band intensities. A DGGE band was defined as an ethidium bromide signal higher than 0.2% of the total intensity of all bands in each lane. The variability in intensities measured among replicate analyses was less than 4%. Prominent bands were excised from the gels and sequenced as reported elsewhere (Casamayor et al. 2000b).

Sequences were submitted to basic local alignment search (Altschul et al. 1990) to get a first indication of what sequences were retrieved. Then, sequences were aligned by using the ARB program package (Technical University of Munich, Munich, Germany; http://www.arb-home.de). Partial sequences were inserted into the optimized and validated tree available in ARB (derived from complete sequence data), by using the maximum-parsimony criterion and a special ARB parsimony tool that did not affect the initial tree topology. Nucleotide sequence accession numbers at EMBL are as follows: AJ239988 to AJ239990 (CIARC-1 to CIARC-3), AJ240004 to AJ240006 (VIARC-1 to VIARC-3), AJ306414 (VIARC-0), AJ306415 (VIARC-4), AJ306416 (VIARC-5), AJ306409 (CIBARC-1), AJ306410 (TOARC-2), AJ306411 (CRARC-3), AJ306412 (CRARC-4) and AJ306413 (BANARC-5 to BANARC-7).

RESULTS

All systems studied had 2 water layers separated according to gradients of temperature or conductivity. The top water layer (epilimnion) was oxygenated and with light. The bottom water layer (hypolimnion) was anoxic, sulfide was present, and it was in the dark. Very little archaeal PCR product (or no product at all) was obtained from the upper aerobic layers. Even at the oxic-anoxic interface most of the DGGE bands found in the hypolimnia were absent. Thus, we focused all the analyses in the anoxic hypolimnia. Different sulfide concentrations were measured for the hypolimnion of each system, ranging from 6 µM in Basin III of Lake Banyoles to 1050 µM in Lake Cadagno (Table 1). Temperature in the study sites ranged between 7 and 25°C. Conductivity ranged between 1600 and 48 000 µS cm⁻² and pH was between 7 and 8. Most systems were sampled in summer (July to September), whereas Lake Cisó and Lake Vilar were sampled several times and in different seasons. The concentration of prokaryotes in the bottom waters was around 10⁷ cells ml⁻¹ (Table 1). This count includes all DAPI-stained particles with a prokaryote-like morphology. Morphologically conspicuous cells such as green and purple photosynthetic sulfur bacteria (Chlorobium sp. and Thiocystis minor, respectively) were counted separately (see results in Casamayor et al. 2000a).

Lake Vilar was sampled twice each month, from February to June 1996. No marked differences in conductivity, temperature or pH were found during the survey in the hypolimnion of the lake (Table 1). Sulfide concentrations, however, increased from winter to summer in the hypolimnion (from 800 to 1200 µM at around 4.5 m). Lake Cisó was sampled at the end of summer 1992 and in winter and early spring 1996. Small differences were found in the physicochemical parameters measured in this lake, and sulfide ranged between 150 and 400 µM (Table 1). Lake Cisó becomes
anoxic during winter holomixis because of its special limnological conditions, and sulfide concentrations up to 400 µM were measured from the bottom of the lake to the surface.

Anaerobic archaeal assemblages inhabiting these systems were compared by DGGE fingerprinting. The number of well-defined sharp bands ranged between 4 and 14 among lakes (Fig. 1), and in most cases just a few bands (1 to 3) appeared as the most intense. The fingerprints showed differences in the mobility of the DGGE bands among hypolimnia of lakes (Fig. 1). Very similar fingerprints were observed in winter and in spring for Lake Vilar (Fig. 2) and Lake Cisó (see Fig. 3 in Casamayor et al. 2000b). In Lake Vilar, however, a shift was apparent in summer from the rest of the year (lane for 20 June in Fig. 2). Very little archaeal PCR product was obtained from the upper aerobic layers of Lake Vilar. As was the case with other lakes, most of the archaeal DGGE bands appearing in the hypolimnetic samples were absent from the oxic-anoxic interface. As an example of the DGGE band pattern from such an interface, the 3.75 m sample is presented in Table 1 and in the last lane in Fig. 2.

The most prominent bands in the profiles (in terms of intensity and frequency of appearance) were excised, re-amplified, purified and sequenced (27 bands). From these, only 17 bands yielded good-quality sequences. Some of them were obtained from the same position in different lanes and gave the same sequence (e.g., VIARC-2 in Fig. 2). Thirteen sequences were included in a phylogenetic tree (Fig. 3) together with bands previously obtained from Lake Cisó in 1996 (CIARC-1, -2 and -3; Casamayor et al. 2000b). The sequences were named, in addition to the number shown in Figs 1 & 2, with a code for the lake and a code for the primer used (ARC). The components of the archaeal assemblages were distributed among 3 groups: methanogens, thermoplasmals and crenarchaeota (Fig. 3). Sequences from thermoplasmals and relatives cluster and crenarchaeota were very distantly related to cultured organisms (e.g., 81% to Desulfurococcus mobilis or Thermoplasma acidophilum) whereas methanogens where more similar to cultured strains (e.g., 90% to Methanospirillum hungatei). The sequences belong-

ing to methanogens formed a closely related group and were all related to endosymbionts of anaerobic ciliates (97% similarity). Thermoplasmales-related sequences obtained from different lakes grouped in a single branch that was not related to previously described sequences. Lake Vilar was the only water mass where a planktonic crenarchaeota-like population was detected.

The relative band intensities were used as an estimate of abundance to compare the composition of the archaeal assemblages among samples that were amplified in the same PCR run. For this comparison, the sequences were pooled into 3 phylogenetic groups: methanogens, thermoplasmales-related and crenarchaeota. An average of 22% of the total band intensity in each lane could not be sequenced and assigned to known organisms and, therefore, these bands were assigned to an ‘unidentified’ class (black bars in Figs 4 & 5). In general, this fraction consisted of minor components of the community (i.e., weak bands). It must be kept in mind that these bands may correspond to any of the 3 identified groups or to other groups of Archaea. On the basis of these results, we found environments where thermoplasmales-related organisms reached >50% of the total intensity (Lake Vilar in summer and the 3 basins of Lake Banyoles), environments where methanogens reached >50% of the total intensity (Lake Vilar in spring and winter), and environments dominated by 1 crenarchaeota (Lake Cisó) and environments dominated by 1 thermoplasmales-related organism (Lake Vilar in winter). In Lake La Cruz 40% was assigned to thermoplasmales-related organisms and 35% to methanogens. In El Cibollar coastal lagoon, methanogens were 40% but the composition of the remaining 60% was not possible to allocate. Lake El Tobar had a very high unidentified fraction too and thermoplasmales-related organisms were found at 20% of the total intensity.

When a temporal survey was carried out in Lake Vilar a shift from crenarchaeota to thermoplasmales-related sequences was observed from winter to summer (Fig. 5). Crenarchaeota accounted for up to 80% of the total band intensity in February and progressively decreased to 10% in June. In parallel, thermoplasmales-related sequences increased from <10% in February up to 60% in June. The methanogen contribution was largest in spring (around 45% of the total band intensity) but similar percentages were measured in winter and summer (between 5 and 10%). Only 1 sequence of crenarchaeota was recovered from Lake Vilar (VIARC-2). On the other hand, 2 closely related sequences within the thermoplasmales cluster (VIARC-1 and VIARC-5, 99% similarity in 16S rDNA between them) and 2 of methanogens (VIARC-3 and VIARC-4, 98% similarity) were recovered from Lake Vilar. These sequences showed temporal changes in their respective band intensities (Fig. 6).
DISCUSSION

Up to now, freshwater planktonic culturable Archaea were restricted to methanogens and sulfur-metabolizing thermophiles. Archaea inhabiting sulfurous environments had been restricted mostly to crenarchaeotes with thermophilic and acidophilic metabolisms (Brock 1997, Huber et al. 1998, Vásquez et al. 1999). Methanogens have also been detected coexisting with sulfate reducers in sulfurous environments although at very low rates of activity (Senior et al. 1982) and were expected not to be very abundant in such systems because sulfate-reducing bacteria are better competitors for available acetate and H₂ (Abram & Nedwell 1978). Recently, a planktonic crenarchaeote-related 16S rDNA sequence was obtained from a Norwegian meromictic lake (Lake Sælenvan- net; 150 µM sulfide, salinity 2%) and methanogens were also present (Øvreås et al. 1997). Methanogens and thermoplasmal-related sequences have also been detected in the sulfide-reducing water layers of the saline Solar Lake (1 mM sulfide; salinity 19%) (Cytryn et al. 2000). Obviously, members of these groups are able to thrive in a wide range of environments and are not limited to the extremophiles known from cultures.

In the present work we have compared archaeal assemblages from several anaerobic, sulfide-rich hypolimnia. In situ conditions in these habitats match neither with the physiological demands of cultured archaeal strains nor with the environments from where clones deposited in databases were obtained. Therefore, these systems are potential habitats for unknown groups of Archaea. For this comparison we have used a fingerprinting technique (DGGE) and sequence analysis of the resulting bands. The small number of published studies using DGGE separation of 16S rDNA PCR products of Archaea reflects that this separation has been difficult to perform. Most of these publications show DGGE profiles characterized by highly complex and mostly diffuse bands. In several cases, a previous nested PCR step was introduced (e.g., Øvreås et al. 1997) that might increase the potential biases introduced by PCR. We have described a primer set and PCR conditions that yielded successful fingerprints with mostly sharp bands in the gel (Casamayor et al. 2000b). The PCR products were obtained directly from the original DNA and allowed us to carry out a semi-quantitative approximation. In addition, the length of the PCR product (600 bp) allowed more reliable identification of the sequences than using shorter 16S rDNA fragments for comparison in databases. A recent work targeting the same 16S rDNA region (Cytryn et al. 2000) reported nonspecific amplification using a similar primer set when annealing temperature decreased to 52°C. On the contrary, our results showed that all the sequences we recovered from the fingerprints belonged to Archaea. Some of the bands excised from our gel, however, did not yield usable sequence data. This difficulty, intrinsic to DGGE, may be related to the presence of different sequences in the same DGGE band or to the fact that weak bands may not have enough template DNA for their reamplification. A more detailed discussion on potentials and limitations of DGGE has been previously published (Casamayor et al. 2000b).

Archaeal DGGE fingerprints are considerably heterogeneous among systems and differences within each lake were smaller than those between lakes. A temporal survey indicated, however, strong changes in time for the archaeal assemblage in Lake Vilar. Therefore, these assemblages appear to be different among lakes, but also summer and winter populations can be different within a lake. This and the identity of the main populations were confirmed after sequencing of the DGGE bands. As expected, the archaeal sequences retrieved did not share close similarity with any cultured organisms. The sequences recovered belonged to 3 different groups. First, the methanogens that we recovered were closely related to a methanogenic endosymbiont of the anaerobic ciliate *Plagiopyla*. This is a well-known symbiosis (Wagener et al. 1990, Fenchel & Finlay 1991) and in our study sites anaerobic ciliates such as *Plagiopyla* and *Metopus* were also seen by epifluorescence microscopy carrying endosymbiotic...
methanogens (Massana & Pedrós-Alió 1994). Second, we found that the previously described crenarchaeotal sequence VIARC-2 (Casamayor et al. 2000b) was present in Lake Vilar throughout the temporal study. And, third, the phylogenetic affiliation showed a new lineage of organisms within the thermoplasmales and relatives group that contained only our sequences from karstic lakes (Lake Vilar, Lake Banyoles, Lake El Tobar and Lake La Cruz). This lineage is included in a bigger cluster containing sequences from freshwater environments (clones from sediments and plankton).

The same fingerprinting and identical sequences (BANARC-5, -6 and -7) were obtained from the 3 basins of Lake Banyoles (B-III, B-IV and B-VI). This is not surprising because all 3 basins are interconnected within this multibasin lake (Casamitjana & Roget 1993). Interestingly, sequences from Lake Banyoles and Lake Vilar, which were only 10 m apart, were not the same. All these sequences were distantly related to sequences CIARC-3 and CIARC-2 obtained from Lake Cisó (1 km away from the former). These results are in line with previous results, suggesting that anaerobic hypolimnia of stratified lakes may act as islands in an aerobic world for anaerobic microorganisms (Casamayor et al. 2000b). We cannot exclude, however, that different limnological conditions for each lake may select different microbial assemblages.

Recent reports show that the group of thermoplasmales and relatives is more widely spread than previously thought and can be found in both marine and freshwater environments (DeLong 1998, Van der Maarel et al. 2000, Bowman et al. 2000, Jurgens et al. 2000, Massana et al. 2000). We found sequences related to the thermoplasmales and relatives in 7 out of 9 study sites, but we cannot discount that they were also present in the high percentage of undetermined sequences from Lake Cadagno and the El Cibollar coastal lagoon. Therefore, they appear to be a widespread lineage. The sequences we retrieved were not related to clones from marine environments and in some cases were close to sequences from lake sediments. Unfortunately, these organisms have not been recovered in pure culture and phenotypic properties cannot be inferred either, due to their distant genotypic relation with cultured archaea. Therefore, their ecological significance, biochemistry, physiology and impact on biogeochemical cycles remain unknown. Some of the archaeal sequences are closely related to endosymbionts and we cannot rule out that some others also correspond to endosymbiotic microorganisms. It is also possible that new anaerobic metabolisms or microbial interactions are involved, as was shown recently for the anaerobic oxidation of ammonium and methane (Jetten et al. 1998, Strous et al. 1999, Boetius et al. 2000) and the association with metazoan species (Preston et al. 1996, Van der Maarel et al. 1998).

Using the relative band intensity as an estimate of abundance, we found a striking dominance of thermoplasmales-related sequences within the archaeal assemblage of sulfurous lakes, especially in the Banyoles karstic area, and a temporal change in the abundance of populations. These changes were higher than the methodological error of 4% intensity (e.g., 70% range of variability for crenarchaeota and 50% for thermoplasmales; see Fig. 5). We were cautious in the number of PCR cycles run to avoid the ‘plateau’ phase and in using the same amount of template in each reaction. The samples that we compared were run in the same PCR run and analyzed in the same DGGE gel. If there was any PCR bias, it should have been the same in all lanes. Therefore, comparison among these samples is valid. In addition, throughout the seasonal study, gradual changes were seen and were not a random distribution of the intensity of the band for each population, adding further credibility to the changes observed. Moreover, we and others have published acceptable correlation between quantitative information obtained by microscopy and by DGGE (Casamayor et al. 2000b and references therein). Thus, although not free of limitations, signal intensity of DGGE bands obtained after ethidium-bromide staining is a useful tool to calculate relative percentages of the different groups. Obviously, absolute quantitative data on archaeal abundance require the use of other techniques.

Our study found qualitative and quantitative temporal changes in the archaeal assemblage of a lake, and we were able to identify the populations involved by DGGE and sequencing. Temporal changes in archaeal assemblages were recently described affecting morphology, abundance and identity of archaeal assemblages in several systems. Thus, successive peaks of 2 archaeal morphotypes were observed during autumn in a high mountain lake by fluorescence in situ hybridization (FISH) (Pernthaler et al. 1998). A maximum of 5% of all DAPI-stained cells (equivalent to around 1.7 × 10^5 archaeal cells ml^-1) hybridized with the general archaeal probe ARCH915 in this lake but archaeal populations were hardly present thereafter. In Antarctic coastal waters, a strong seasonality of archaeal rRNA concentrations has been reported, with the highest values occurring during the austral winter and early spring (Massana et al. 1998, Murray et al. 1998). Marine group I (crenarchaeota-affiliated) was shown to be more abundant in Antarctic coastal waters in late winter and early spring, whereas the marine group II (thermoplasmales-affiliated) was a minor component but peaked in abundance in summer and early autumn (Murray et al. 1998). Intriguingly, this temporal partitioning of the system between crenarchaeota and thermoplasmales-related micro-
organisms matches what we have found in a karstic lake. The environmental factors responsible for these changes in time, however, remain unknown.

Our results indicate that the genetic diversity of Archaea is relatively large in non-thermophilic, sulfuriferous environments. We were able to identify shifts in sequences at the microdiversity scale (more than 97% similarity in 16S rDNA) after sequencing DGGE bands. Several investigators have indicated that such small differences in 16S rRNA genes may belong to closely related but ecologically different populations (Fuhrman & Campbell 1998, Moore et al. 1998). The abundance of these Archaea remain to be determined but these data suggest that they are novel types of organisms that may be widely distributed and contribute to the anaerobic biogeochemical cycles of karstic lakes and coastal lagoons. The vertical distribution (accumulating in sulfide-rich layers) and the temporal dynamics indicate that these archaeal assemblages are autochthonous. The challenge now is to determine the physiological capabilities and environmental roles of these uncultured but ecologically relevant Archaea.

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