Prokaryotic community analysis with CARD-FISH in comparison with FISH in ultra-oligotrophic ground- and drinking water

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Introduction

Well and spring water from oligotrophic groundwater habitats are of great importance for public water supply in many areas throughout the world. Thus, understanding the biogeochemistry and microbiology of these aquatic resources is of general interest as it has an impact on the ability to distribute and store this water (i.e. ‘biostability’) in water supply systems. Only limited information exists on the occurrence and significance of prokaryotic populations, both, in the aquifer itself as well as in the abstracted water (Ghiorse and Wilson 1988; Gounot 1994; Griebler et al. 1999). Studying the microbiology and ecology of these oligotrophic to ultra-oligotrophic systems provides also knowledge on basic ecological questions such as adaptation and survival strategies of micro-organisms under nutrient-poor conditions. However, a basic requirement to study these micro-organisms in such systems is the availability of methods which allow for sensitive, specific and quantitative detection of prokaryotic cells in a reduced metabolic state.

Keywords
Archaea, Bacteria, CARD-FISH, drinking water, FISH, oligotrophic ground- and mineral water.

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2006/1186: received 20 August 2006, revised 6 November 2006 and accepted 3 January 2007
doi:10.1111/j.1365-2672.2007.03319.x

Abstract

Aims: We compared the applicability of catalysed reporter deposition fluorescence in situ hybridization (CARD-FISH) and FISH to enumerate prokaryotic populations in ultra-oligotrophic alpine groundwaters and bottled mineral water

Methods and Results: Fluorescent oligonucleotide probes EUB338 and EUB338mix (EUB338/EUB338-II/EUB338-III) were used to enumerate bacteria and probes EURY806 and CREN537 for Euryarchaea and Crenarchaea, respectively. Improved detection of Planctomycetales by probe EUB338-II was tested using a different permeabilization step (proteinase K instead of lysozyme). Total detection efficiency of cells in spring water of four different alpine karst aquifers was on average 83% for CARD-FISH and only 15% for FISH. Applying CARD-FISH on bottled natural mineral waters resulted in an average total hybridization efficiency of 89%, with 78% (range 77–96%) bacteria and 11% (range 3–22%) identified as Archaea.

Conclusions: CARD-FISH resulted in substantially higher recovery efficiency than FISH. Hence, CARD-FISH appears very suitable for the enumeration of specific prokaryotic groups in ground- and drinking water.

Significance and Impact of the Study: This study represents the first evaluation of CARD-FISH on ultra-oligotrophic ground- and drinking water. Results are relevant for basic research and drinking water distributors. Archaea can comprise a significant fraction of the prokaryotic community in bottled mineral water.
During the last decade, the fluorescence in situ hybridization (FISH) method ascended towards an indispensable tool to enumerate specific prokaryotic populations in marine and freshwater habitats (DeLong et al. 1989; Behrens et al. 2003a, 2003b; Yeates et al. 2003; Chen et al. 2004; Sekar et al. 2004). Although FISH has gained widespread acceptance, there are some technical and conceptual problems that can lead to a great variability in the detection of target cells, especially under oligotrophic conditions (Oda et al. 2000; Pernthaler et al. 2002b; Bouvier and del Giorgio 2003). Frequently only a low fraction of all planktonic cells (<50%) can be visualized by FISH in freshwater systems (Sekar et al. 2003) as well as in coastal surface waters (Pernthaler et al. 2002a). Considerable efforts have been made to overcome some of these problems (Amann et al. 1990; Ouverney and Fuhrman 1997; DeLong et al. 1999; Fuchs et al. 2000; Pernthaler et al. 2002b; Behrens et al. 2003a, 2003b). Probably the most significant improvement was the development of the catalyzed reporter deposition-FISH (CARD-FISH) (Schönhuber et al. 1997; Pernthaler et al. 2002a), facilitating the detection of cells with low ribosomal content. The critical step in the CARD-FISH approach is the diffusion of large molecules, in this case the horse radish peroxidase (HRP)-labelled probe, into whole cells embedded in an agarose matrix. A directed permeabilization of prokaryotic cell walls prior to the hybridization step is therefore of crucial importance to enable the penetration of the probe (Pernthaler et al. 2002a).

Another challenge when using FISH-based protocols is the fact that only a small fraction of the occurring prokaryotes are taxonomically characterized. Consequently, all rRNA-targeting probes have to be continuously re-evaluated and modified as new sequence data is reported (Daims et al. 1999). For example, the probe EUB338 was used to enumerate bacteria, however, studies indicate that this probe is insufficient for detecting all bacterial groups (Hugenholtz et al. 2001). Therefore, two supplementary probes, EUB338-II (targeting Planctomycetales) and EUB338-III (targeting Verrucomicrobia), were designed for a more complete detection of this domain (Daims et al. 1999). Unfortunately, there are few studies to date which evaluated the effectiveness of the probe mix (EUB I–III) when performing CARD-FISH in freshwater environments. In addition, there are only very limited data about the applicability of FISH-based protocols to detect members of the domain Archaea in freshwater systems (Bouvier and del Giorgio 2003).

The aim of this study was to evaluate the applicability of CARD-FISH to enumerate specific prokaryotic populations in ultra-oligotrophic ground- and drinking water from various environments. As a first step, the CARD-FISH protocol was adapted (i.e. optimization of permeabilization and amplification time) and evaluated (i.e. cell loss determination, effect of different enzymatic treatments on Planctomycetales permeabilization) for prokaryotic cells from oligotrophic freshwater habitats. The adapted protocol was then used to compare the CARD-FISH procedure with the conventional FISH method in spring water samples from four different alpine karst springs covering a gradient from extremely low to high surface-influenced karst habitats (Farnleitner et al. 2005). A higher mean detection efficiency of CARD-FISH compared with FISH was obtained by applying an EUB338mix (EUB338, EUB338-II, EUB-III) and an improved protocol for archaean cell detection (Teira et al. 2004). Finally CARD-FISH in bottled drinking water samples, originating from various European groundwater habitats, demonstrated its general applicability for water samples from oligotrophic and ultra-oligotrophic environments.

Materials and methods

Study sites and sample preparation

The selected groundwaters from four different alpine springs are located in the Northern Calcareous Alps in Austria (Farnleitner et al. 2005). DKAS 1 and HKAS 3 show a discharge regime with very low variations and an average water storage capacity of >20 years. In contrast, the discharge response after precipitation is very quick at LKAS 2 and SKAS 9, and the average water retention time is in the range of only a few years. Total prokaryotic abundance in the springs is in the range of about 10 7–10 8 cells l−1, reflecting their ultra-oligotrophic conditions (Farnleitner et al. 2005). Because of varying substrate supply and surface runoff conditions, cell counts from aquifers with high water retention times (i.e. DKAS 1 and HKAS 3) are typically lower than in more dynamic aquifers (i.e. LKAS 2 and SKAS 9) (Farnleitner et al. 2005).

Samples were collected from January 2004 to July 2004 at three to four weeks intervals, directly at the spring outlet. Duplicate samples of 60 ml were taken and immediately fixed with paraformaldehyde (final concentration 2%) at 4°C in the dark for 14–18 h. Subsequently, the samples were filtered through polycarbonate filters (0.2-μm pore size; 25-mm diameter; type GTF; Millipore Corp. Bedford, MA, USA), supported by cellulose acetate filters (0.45-μm pore size; Millipore), to ensure an equal distribution of the cells on the filter surface, washed twice with 5 ml of 0.2-μm filtered Milli-Q water, air-dried and stored at −20°C. In addition, the abundance of major prokaryotic groups was examined in 12 different brands of mineral waters, differing in their geological origin and bottled in different European countries. Their brands were made anonymous and they were randomly designa-
ted from 1 to 12. Bottles of mineral water were sampled aseptically. After mixing the water in the bottle, duplicate samples of 90 ml were taken from each bottle immediately after opening it and treated as described.

**Fluorescence in situ hybridization**

Filters from spring water samples were cut into halves. One was kept at –20°C for CARD-FISH the other was used for FISH and further cut into three parts for hybridization with different probes. The FISH procedure is based on the standard protocol (Mantz et al. 1993; Glöckner et al. 1999) using 5 ng μl⁻¹ of the respective probe. All probes were purchased from Thermoelectron (Germany). Oligonucleotide probes for FISH were 5’-monolabelled with the indocarbocyanine dye Cy3. One section of the filter was hybridized with the EUB338mix containing probes EUB338 (5’-GTCGCTTCCGTAGGAT-3’; targeting bacteria), EUB338-II (5’-GACGCCACCCGTAGGTTG-3’; targeting Planctomycetales), EUB338-III (5’-GCTGCCACCGTAGGTTG-3’; targeting Verrucomicrobia) (Daims et al. 1999) simultaneously, one section with probe EUB338 only and one section with probe NON338 as a control for nonspecific binding (Amann et al. 1995). After incubation in a prewarmed washing solution (Glöckner et al. 1999), filter sections were dipped in 80% ethanol, dried on Whatman 3M paper (Whatman Ltd., Dassel, Germany) and placed on a glass slide. Subsequently, they were mounted with a drop of DAPI mix (5:5 parts Citifluor (Citifluor, Ltd., Leicester, UK), 1 part Vectashield (Vector Laboratories, Inc., Peterborough, UK), 0:5 parts phosphate-buffered saline (PBS) with DAPI at a final concentration of 1 μg ml⁻¹ for counterstaining the cells. At the first two sampling dates, additional filter sections were hybridized with Cy3-monolabelled *Archaeal* probes (EUB3806, 5’-CACAGGCTTTACACCTAG-3’ targeting Euryarchaeota; CREN537, 5’-TGACCACTTGAGGTGCTG-3’; targeting Crenarchaeota) (Teira et al. 2004). The probes target the same cells as the polynucleotide probes used in previous studies (DeLong et al. 1999; Karner et al. 2001). Positive controls were performed with prefiltered (0:2 μm) spring water samples (two samples from each DKAS 1 and LKAS 2) spiked with *Escherichia coli* [strain K12 (DSM4509 harvested at late exponential phase, purchased from DSMZ)] at two different final concentrations (10⁵ cells ml⁻¹ and 10⁶ cells ml⁻¹). After immediate fixation as described before they were hybridized with the EUB338mix.

**Catalysed reporter deposition-fluorescence in situ hybridization**

The remaining filter halves were dipped in low-gelling-point agarose [0:1% (wt/vol) Biozym, USA; in Milli-Q water] dried upside down on a glass petri dish at 37°C and dehydrated in 96% (vol/vol) ethanol (Pernthaler et al. 2002a). Subsequently, they were cut into four to five sections for hybridization with different HRP-labelled oligonucleotide probes. CARD-FISH was based on the protocol for marine habitats (Teira et al. 2004) using the same permeabilization, hybridization and washing conditions, however, with some modifications as subsequently described. Cell wall permeabilization was performed by incubation of the respective filter sections in either lysozyme (10 mg ml⁻¹; Sigma, Vienna, Austria) for bacteria (probes EUB338, EUB338-III, NON338) or proteinase K (0:2 μl ml⁻¹, 1:84 U mg⁻¹, 10:9 mg ml⁻¹; Fluka, Buchs, Switzerland) solution [0:05 mol l⁻¹ EDTA, 0:1 mol l⁻¹ Tris-HCl (pH 8)] for *Archaea* (EUB3806, CREN537) and EUB338-II at 37°C for 1 h. Filters were washed three times with Milli-Q water and subsequently incubated in 0:01 mol l⁻¹ HCl for 20 min at room temperature to inhibit potentially present intracellular peroxidase and residual proteinase K. After two more washing steps in Milli-Q water, filters were dehydrated with 96% ethanol and dried at room temperature. Filter sections were hybridized at 35°C (Schönhuber et al. 1997) for 10–12 h with the EUB338mix, EUB3806, CREN537 and NON338, respectively. Three hundred microlitres of hybridization buffer [0:9 mol l⁻¹ NaCl, 20 mmol l⁻¹ Tris-HCl (pH 7–7.5), 10% (wt/vol) dextran sulfate, 0.002% (wt/vol) sodium dodecyl sulfate (SDS), 1% blocking reagent (Boehringer Mannheim, Mannheim, Germany) and 55% (vol/vol) formamide (for EUB338, EUB338-II, EUB338-III and NON338) or 20% (vol/vol) formamide (for EUR036 and CREN 537) were transferred into a 0-7-ml reaction vial. The respective HRP probe was added to a final concentration of 0.28 ng μl⁻¹ (0.05 μmol l⁻¹). After hybridization, the filter sections were transferred into 50 ml of prewarmed washing buffer [5 mmol l⁻¹ EDTA (pH 8), 20 mmol l⁻¹ Tris-HCl (pH 7.4–7.6), 0.01% (wt/vol) SDS] containing 13 mmol l⁻¹ NaCl (for EUB338, EUB338-II, EUB338-III and NON338) and 145 mmol l⁻¹ NaCl (for EUR036 and CREN 537) at 37°C for 10 min. Thereafter, all sections were transferred in PBS [145 mmol l⁻¹ NaCl, 1-4 mmol l⁻¹ NaH2PO4, 8 mmol l⁻¹ Na2HPO4 (pH 7.6)] amended with 0.05% Triton X-100 (PBS-T) at room temperature for 15 min. After removal of excess buffer, the filter sections were put into a 1-5-ml reaction vial containing 493 μl of amplification buffer [10% (wt/vol) dextran sulfate, 2 mol l⁻¹ NaCl, 0.1% (wt/vol) blocking reagent and 0.0015% H2O2 (freshly prepared in PBS) and 5 μl of tyramide-Alexa488 (1 mg ml⁻¹; Molecular Probes, Europe) and incubated at 37°C for 45 min. Finally, filter sections were washed at room temperature in PBS-T (15 min) and Milli-Q, dehydrated in 96% ethanol and air-dried. They were put on a
slide, counterstained with DAPI-mix and stored at –20°C until further processing. At three sampling dates, additional sections were hybridized with EUB338 only to determine the fraction of bacteria targeted with the probes EUB-II and EUB-III by comparison with the EUB338mix. Different times for permeabilization, hybridization and amplification were tested to optimize the procedure for freshwater environment. At the first two sampling dates, additional filter sections from spring water samples were stained with DAPI to determine cell losses during the CARD-FISH procedure. In addition, a test series was carried out to examine differences in the hybridization signal of probe EUB338-II when cells were either permeabilized with lysozyme or proteinase K. Additional control experiments were performed without any probe to test for remaining intrinsic peroxidase activity. Filters from mineral water samples were hybridized with the EUB338mix and a mix of the used Archaea probes (EURY806 and CREN537).

Image acquisition and analysis

The stained filter sections were examined on a Leitz DMRB microscope equipped with a HBO 50-W Hg lamp, a 100× Plan Apochromat oil objective and appropriate filter sets for DAPI, Cy3 and Alexa488 (Chroma Tech. Corp., Rockingham, USA). First, DAPI-stained cells were counted in one randomly selected microscopic field, followed by the determination of the fraction of Cy3-(FISH) or Alexa488-(CARD-FISH) stained cells in the same field. At least 800–900 DAPI-stained cells were counted per filter piece. Images were taken with a KAPPA CF15/4MCC camera system.

Statistical analysis

In order to obtain a standardized indication of variability to compare the FISH- and the CARD-FISH data set, the parametric coefficient of variation (%CV) was used (100 × SD divided by the arithmetic mean). In addition to the comparison of the percentage of total hybridization for FISH and CARD-FISH results, CVs for the respective EUB338mix data sets were calculated. All further statistical comparisons were performed in SPSS version 13.0 using the nonparametric Wilcoxon test with related samples and Spearman rank correlation.

Results

FISH in oligotrophic groundwater

The average total prokaryotic abundance of all four springs stained by DAPI was 2.7 × 10^4 cells ml^-1 ranging from a minimum of 2.2 × 10^4 in February to a maximum of 3.4 × 10^4 in January (Table 1). In all samples, the fraction of hybridized cells was higher after FISH with the EUB338mix (on average 5% higher) than with the EUB338 probe alone (P < 0.01, n = 24; nonparametric Wilcoxon test, related samples). Only very few cells were hybridized with the NON338-probe (mean 0.5%). After subtracting the counts from the negative control the mean percentage of DAPI-stained cells visualized with the

### Table 1: Average detection rates by fluorescence in situ hybridization (FISH) for DKAS 1, LKAS 2, HKAS 3 and SKAS 9 (n = 56; seven sampling dates, four springs and two replicates per sample)

<table>
<thead>
<tr>
<th>Date</th>
<th>Total cell counts (10^4 cells ml^-1)</th>
<th>Fraction per cent of total cells detected with probe*</th>
<th>Per cent of total hybridization§</th>
<th>Range min–max</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EUB338</td>
<td>EUBmix†</td>
<td>ΔEUB‡</td>
<td>NON338</td>
</tr>
<tr>
<td>07.01.04</td>
<td>3.4</td>
<td>21</td>
<td>25</td>
<td>4</td>
</tr>
<tr>
<td>02.02.04</td>
<td>2.9</td>
<td>13</td>
<td>22</td>
<td>9</td>
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<tr>
<td>25.02.04</td>
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<td>n.a.***</td>
<td>11</td>
<td>n.a.</td>
</tr>
<tr>
<td>23.03.04</td>
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<td>12</td>
<td>16</td>
<td>4</td>
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<td>19.04.04</td>
<td>2.4</td>
<td>n.a.***</td>
<td>19</td>
<td>n.a.</td>
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<tr>
<td>17.05.04</td>
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<td>6</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>14.06.04</td>
<td>2.3</td>
<td>n.a.***</td>
<td>9</td>
<td>n.a.</td>
</tr>
<tr>
<td>Avg. ††</td>
<td>2.7</td>
<td>13</td>
<td>16</td>
<td>5</td>
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</tbody>
</table>

*Per cent detection of DAPI-stained cells. Between 800 and 900 DAPI-stained cells were counted per filter piece. Average values were based on eight analyses.
†Hybridization with a mixture of probes EUB338, EUB338 II and EUB338 III.
‡Difference between fraction (%) EUB338 and EUBmix.
§Percentage of total hybridization (%EUBmix) normalized to total DAPI counts. Numbers have been corrected by subtracting NON338 counts.
***Not analysed.
††Total average.
n.a., not available.
Euryarchaea age of prokaryotic cells identified as with the two specific archaeal probes. The mean percent-an archaeal contribution of 6–12% of DAPI-stainable cells was 75 ± 7%. Permeabilization with proteinase K resulted in a fraction of hybridized cells with the EUB338mix was increased by 9% when the EUB338mix was applied (Table 2). The mean percentage of DAPI-stainable cells during the CARD-FISH procedure could be detected (data not shown). The mean number of cells hybridized with the EUB338 probe alone was 66 ± 8% (±SE; n = 16; nonparametric Wilcoxon test, related samples). Therefore, no loss of DAPI-stainable cells occurred during the CARD-FISH investigation period (Fig. 2). Nevertheless, the recovery efficiency using CARD-FISH decreased in all four springs during the winter months and steadily increased from end of February until June [Spearman correlation (independent data sets); r = 0.61, P < 0.01, n = 40]. Generally, higher recovery efficiency was obtained for DKAS 1 and HKAS 3, the two springs representing the hydrogeological more stable spring type than for LKAS 2 and SKAS 9 in winter to 99% for HKAS 3 in spring and was significantly higher than the maximum value reached with the FISH protocol (34%) throughout the entire investigation period (Fig. 2).

**CARD-FISH in oligotrophic groundwater**

The described CARD-FISH protocol resulted in high fractions of hybridized prokaryotic cells and intense fluorescent labelling of the cells in water samples from all four springs (Table 2). The average numbers of DAPI-stainable cells (2.7 × 10^8 cells ml⁻¹) were not significantly different from the results of the corresponding DAPI counterstains during CARD-FISH analysis (P = 0.66, n = 16; nonparametric Wilcoxon test, related samples). Therefore, no loss of DAPI-stainable cells during the CARD-FISH procedure could be detected (data not shown). The mean percentage of cells hybridized with the EUB338 probe alone was 66 ± 8% (±SE; n = 24), increasing on average by 9% when the EUB338mix was applied (Table 2). The fraction of hybridized cells with the EUB338mix was 75 ± 7%. Permeabilization with proteinase K resulted in an archaeal contribution of 6–12% of DAPI-stainable cells with the two specific archaeal probes. The mean percentage of prokaryotic cells identified as Euryarchaea was 6 ± 2%, the fraction identified as Crenarchaea was 3 ± 1%. After subtracting counts from the negative control (~0-6%) the total DAPI-stained cells detected with all five oligonucleotide probes averaged 83% (±6%). After lysozyme treatment, 1% of the total cells could be visualized with probe EUB338-II in DKAS 1 and 3% in LKAS 2. Proteinase K treatment augmented these fractions to 2% in DKAS 1 and 6% in LKAS 2 (P < 0.01, n = 16, nonparametric Wilcoxon test, related samples). As expected, controls without probe, to test for remaining intrinsic peroxidase activity, did not give any fluorescent signal.

**Comparison between FISH and CARD-FISH**

The CARD-FISH protocol was not only superior in terms of the intensity of fluorescence labelling but also with regard to mean detection efficiency (15% for FISH vs 83% for CARD-FISH) (Fig. 1). Total recovery efficiency of prokaryotic cells using CARD-FISH ranged from 68% for SKAS 9 in winter to 99% for HKAS 3 in spring and was significantly higher than the maximum value reached with the FISH protocol (34%) throughout the entire investigation period (Fig. 2). Nevertheless, the recovery efficiency using CARD-FISH decreased in all four springs during the winter months and steadily increased from end of February until June [Spearman correlation (independent data sets); r = 0.61, P < 0.01, n = 40]. Generally, higher recovery efficiency was obtained for DKAS 1 and HKAS 3, the two springs representing the hydrogeological more stable spring type than for LKAS 2 and SKAS 9 (Fig. 3). None of these trends could be discerned with FISH results, where the highest recovery was found in

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**Table 2**: Average detection rates by catalysed reporter deposition-fluorescence in situ hybridization (CARD-FISH) for DKAS 1, LKAS 2, HKAS 3 and SKAS 9 (n = 56; seven sampling dates, four springs and two replicates per sample)

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*Per cent detection of DAPI-stained cells. Between 800 and 900 DAPI-stained cells were counted per filter piece. Average values were based on eight analyses.
†Hybridization with a mixture of probes EUB338, EUB338 II and EUB338 III.
‡Difference between fraction (%) EUB338 and EUBmix.
§Percentage of total hybridization of probes (EUBmix, EURY806, CREN537) normalized to total DAPI counts. Numbers have been corrected by subtracting NON338 counts.
**Not analysed
††Total average.
n.a., not available.
DKAS 1 during the winter period and the lowest value (7%) was also in DKAS 1 in May. Furthermore, variability in the recovery efficiency for FISH, indicated by the parametric coefficient of variation, was rather high (CV $47\pm5\%$, $n=28$) compared with CARD-FISH (CV $11\pm5\%$, $n=56$). Variability within the data sets obtained with the EUB338mix showed similar results (FISH, CV $47\pm5\%$, $n=56$; CARD-FISH, CV $14\pm3\%$, $n=56$). One drawback when using CARD-FISH, however, is that cell shapes can hardly be obtained. Image analysis after FISH allows for determination of cell morphologies or sizes (Alfreider et al. 1996). In contrast to that, even the morphological differentiation between rods and cocci is sometimes ambiguous after CARD-FISH hybridization, especially in environments where the majority of cells are rather small, because the cells get bloated during the amplification step.

CARD-FISH analysis of mineral water

The CARD-FISH method was applied on samples from seven carbonated and five noncarbonated mineral waters originating from geologically different aquifers (Table 3). Total counts of DAPI-stained cells ranged between $4\times10^3$ and $1.1\times10^5$ cells ml$^{-1}$. The mean percentage of prokaryotic populations hybridized with all five oligonucleotides probes was $89\pm5\%$, of which 78% were bacteria (stained with the EUB338mix) and 11% (range

Figure 1 Photomicrographs of DAPI-stained hybridized cells from DKAS 1. Pictures (a) and (b) depict corresponding DAPI-stained vs fluorescence in situ hybridization (FISH)-labelled (Cy3 probes) cells. Pictures (c) and (d) depict corresponding DAPI-stained vs catalysed reporter deposition (CARD)-FISH-labelled (Alexa488-labelled tyramide) cells. Both samples were hybridized with the EUB338mix.
1–22%) were Archaea (stained by the ARCHmix) (Table 4). Negative controls ranged between 0% and 0–4% (data not shown) and were not subtracted from the total hybridization results.

**DISCUSSION**

**CARD-FISH vs FISH in oligotrophic freshwater environments**

The presented data demonstrate that CARD-FISH is an effective tool to overcome significant problems occurring when using FISH in oligotrophic and ultra-oligotrophic groundwater environments (e.g. no detection of cells with low ribosomal content). Although FISH was successfully applied in various habitats (Franks et al. 1998; Glöckner et al. 1999; Nielsen et al. 2002), the method is limited in certain environments (e.g. Kenzaka et al. 1998; Pernthaler et al. 2002a,b; Bouvier and del Giorgio 2003). Hybridization assays carried out in ice, snow and drinking waters gave an estimated mean recovery efficiency for bacteria lower than 40% (Bouvier and del Giorgio 2003).

**Table 3** Chemical characterization of different mineral waters*

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<td>5</td>
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<td>3.3</td>
<td>28.3</td>
<td>17.5</td>
</tr>
<tr>
<td>12</td>
<td>n.a.</td>
<td>11.4</td>
<td>9.9</td>
</tr>
</tbody>
</table>

*Parameters were copied from bottles.
†Total dissolved solids.
‡Not available.

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study demonstrates that FISH detection rates in ultra-oligotrophic systems can be as low as 8–23% of DAPI-stainable cells. The CARD-FISH method is hardly affected by a low number of ribosomes, although hybridization results might still be influenced by modification (Kalpaxis et al. 1998) or degradation (Davis et al. 1986) of target sites, providing a possible explanation for the decrease of detection efficiency observed with CARD-FISH during February and March (Fig. 3). It can be speculated that these modifications, besides a low ribosomal content, could be responsible for the high variability of total hybridization results with FISH (Fig. 3), also found in other studies where slow growing communities prevail (Bouvier and del Giorgio 2003). Another reason for the observed decrease in the recovery efficiency using CARD-FISH during the winter months could be alterations in bacterial or archaeal cell walls induced by stress conditions (Gounot and Russell 1999) or increased carbon limitation (Nystrom et al. 1992), rendering the cell wall more resistant to the permeabilization step and prevent the probe from entering the cell (Wagner et al. 1998). As this study identified organisms on a low taxonomic resolution level, it remains unknown whether the lower recovery efficiency during the winter months is caused by a less efficient hybridization of the entire community or individual groups (Farnleitner et al. 2005). Although CARD-FISH results showed a fraction of at least 6% belonging to the domain Archaea (Table 2) in these karst water aquifers, identification with FISH was neither possible for Euryarchaeota nor for Crenarchaeota (Table 1). Crenarchaeota comprised 3% of the total archaeal community. This group appears to be ubiquitously present in aquatic habitats and soil and has been reported in a wide variety of temperate and cold environments (Sjöling and Cowan 2003). The majority of the archaeal community was identified as Euryarchaeota (6%). The recovery efficiency using CARD-FISH likely represents the upper level of active prokaryotes as DAPI not only stains viable cells, but also detects dead or ghost cells lacking ribosomes and hence, are not detectable by CARD-FISH (Zweifel and Hagström 1995).

Using the EUB338mix

The importance of an optimized permeabilization step becomes evident when using the EUB338mix. Probe EUB338-II was designed for the order Planctomycetales (Schlesner 1994; Ward et al. 1995). Planctomycetales exhibit some unique characters, most importantly their cell wall is similar to that of Archaea (Kandler and König 1998; Neef et al. 1998). This validates our results indicating better permeabilization of Planctomycetales when cell

### Table 4 Average detection rates by catalysed reporter deposition-fluorescence in situ hybridization (CARD-FISH) for different mineral waters (n = 24, 12 bottles, in duplicate)

<table>
<thead>
<tr>
<th>Type</th>
<th>Country</th>
<th>Total cell counts (10⁴ cells ml⁻¹)</th>
<th>Fraction per cent of total cells detected with probes*</th>
<th>Per cent of total hybridization §</th>
<th>Range min–max</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Spain</td>
<td>2.0</td>
<td>96</td>
<td>3</td>
<td>99</td>
</tr>
<tr>
<td>2</td>
<td>Iceland</td>
<td>0.4</td>
<td>80</td>
<td>8</td>
<td>89</td>
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<tr>
<td>3</td>
<td>Western Austria</td>
<td>8.3</td>
<td>85</td>
<td>7</td>
<td>93</td>
</tr>
<tr>
<td>4</td>
<td>Germany</td>
<td>6.0</td>
<td>77</td>
<td>15</td>
<td>92</td>
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<tr>
<td>5</td>
<td>Switzerland</td>
<td>2.9</td>
<td>82</td>
<td>6</td>
<td>87</td>
</tr>
<tr>
<td>6</td>
<td>Belgium</td>
<td>10.5</td>
<td>80</td>
<td>11</td>
<td>91</td>
</tr>
<tr>
<td>7</td>
<td>France</td>
<td>3.4</td>
<td>80</td>
<td>13</td>
<td>92</td>
</tr>
<tr>
<td>8</td>
<td>Central Austria</td>
<td>0.8</td>
<td>73</td>
<td>20</td>
<td>93</td>
</tr>
<tr>
<td>9</td>
<td>Eastern Austria</td>
<td>0.4</td>
<td>68</td>
<td>18</td>
<td>86</td>
</tr>
<tr>
<td>10</td>
<td>Western Austria</td>
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<tr>
<td>12</td>
<td>Spain</td>
<td>1.8</td>
<td>79</td>
<td>1</td>
<td>80</td>
</tr>
<tr>
<td>Avg.**</td>
<td></td>
<td>3.7</td>
<td>78</td>
<td>11</td>
<td>89</td>
</tr>
</tbody>
</table>

Samples 1–7 represent carbonated mineral water and samples 8–12 represent non-carbonated mineral water.

*Percent detection of DAPI-stained cells. Between 800 and 900 DAPI-stained cells were counted per filter piece. Average values were based on two analyses.

†Hybridization with a mixture of probes EUB338, EUB338 II and EUB338 III.

‡Hybridization with a mixture of probes EURY806 and CREN 537.

§Percentage of total hybridization of probes (EUBmix, ARCHmix) normalized to total DAPI counts.

**Total average.

n.a., not available.
walls were permeabilized with proteinase K than with lysozyme. Some studies describe Planctomycetes attached to particulate matter or cell clusters and suggest a possible role in the degradation of more recalcitrant compounds (Neef et al. 1998). This could be advantageous in alpine karst ecosystems and one reason for the high detection rate in LKAS 2 (6%) than in DKAS 1 (2%) is that LKAS 2 is a more dynamic spring type and likely contains more particulate matter (Farnleitner et al. 2005).

CARD-FISH in natural mineral water

Although consumption of natural mineral water is constantly increasing, there are only very few studies on prokaryotic community composition in this environment. Most approaches are based on isolation and subsequent identification (Defives et al. 1999; Dewettinck et al. 2001; Leclerc and Moreau 2002) and/or subsequent molecular typing (Nichols et al. 2003; Villari et al. 2003) and therefore suffer from well-recognized cultivation biases. Evaluation of new methods for microbiological assessments of the microbial community in mineral water are not only interesting because of the unique habitat characteristics, but are an industry-driven need (Ramalho et al. 2001). The CARD-FISH approach we used, applying the EUB338mix and ARCHmix, gain reliable and reproducible results. Even with total cell numbers as low as 3.6×10^3 cells ml⁻¹, total hybridization rates were still higher than 80% (Table 4). Our data strongly indicates the presence of an archaeal community in all ground-mineral waters analysed, comprising between 1% and 20% of the total prokaryotic abundance. Recently, changes in bacterial community composition in mineral water after bottling were reported using FISH (Loy et al. 2005). Although only 63% of all cells were present after the initial growth phase (about 1 week after bottling), unreliability of FISH in oligotrophic systems became especially evident during the first days after bottling. The number of FISH-detectable cells was only on a maximum level of 3% when total cell counts remained constant at 2×10^3 cells ml⁻¹. This highlights the (in)sensitivity of the FISH method when prokaryotic activity is low (Jones et al. 1999; Leclerc and Moreau 2002) such as in bottled mineral water. The low detection rate is in accordance with earlier findings that report hybridization of 5% of total counts with a bacteria-specific probe (Leclerc and Moreau 2002). As bottles in our study were not obtained directly from the producer, the microbial community was analysed after an unknown time span after bottling and thus, after the initial growth phase, when total cell counts were already at a stable level (Defives et al. 1999; Dewettinck et al. 2001). Although the applicability of CARD-FISH for a time course analysis during the first weeks after bottling was not tested, the rather small variations in prokaryotic abundance among the different brands of mineral water indicate that this method is as an adequate tool for future investigations. It would be important to further examine not only community shifts after the bottling process but also the autochthonous prokaryotic community of the source habitat and vulnerability indicators with more specific probes using the CARD-FISH approach. A possible combination of CARD-FISH with microautoradiography (MAR) (Teira et al. 2004) would not only provide reliable data on the abundance of particular populations but also on the physiological state of the community. In conclusion, the applicability of FISH and CARD-FISH in ultra-oligotrophic freshwater systems was compared. We found severe limitations for the FISH approach in environments where the prokaryotic community is likely to be in a starvation-survival state. CARD-FISH resulted in high and reproducible recovery efficiencies that were at the upper end of what might be reached with this method. Therefore, CARD-FISH was found to be a useful and straightforward tool to obtain enfolding insights into the prokaryotic community composition including the archaeal fraction in ultra-oligotrophic aquatic environments like ground- and drinking water.

Acknowledgements

This study was funded by the FWF project P18247-B06 granted to A.H. Farnleitner and supported by the Vienna Waterworks. Special thanks also to Hermann Kain for crucial technical assistance.

References


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