Bacteria in post-glacial freshwater sediments

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Prokaryote communities in post-glacial profundal freshwater sediments of Windermere, representing 10–12 000 years of deposition, were examined for culturability, viability and community structure. The potential for active geochemical cycles was inferred from the presence of specific groups of bacteria. Direct count procedures revealed $10^{12}$ cells (g dry wt sediment)$^{-1}$ in the surface sediments, which declined to approximately $10^{10}$ cells (g dry wt sediment)$^{-1}$ at 6 m depth of core (representing approximately 10 000 years of deposition). The majority of the cells in the upper sediments were metabolically active when challenged with viability probes and responded to the direct viable count method. Below 250 cm, viability shown by 5-cyano-2,3-dirol tetrazolium chloride (CTC) dye was not significantly different from the direct count; however, counts obtained with 5-carboxyfluorescein diacetate (CFDA) and the direct viable count both declined significantly from the direct count below 250 cm and 1 m, respectively. Culture was achieved from samples throughout the core, although the numbers of culturable bacteria decreased significantly with depth, from $10^7$ c.f.u. (g dry wt sediment)$^{-1}$ to $10^{-2}$ c.f.u. (g dry wt sediment)$^{-1}$ below 3 m depth. Among culturable isolates, Gram-positives and Gram-negatives were found at all levels of the core, and spore-forming heterotrophs dominated. Although sulphate-reducing bacteria were not detected below 20 cm, isolates demonstrating denitrifying activity were detected at all depths. PCR performed on samples taken below 3 m (deposited more than 7000 years ago) using eubacterial and archaeal primers revealed sequences similar to those found in deep sediments of the Pacific Ocean and the presence of methanogenic archaea. These observations indicate that bacteria and archaea are capable of long-term persistence and activity in deep, aged freshwater sediments.

Keywords: subsurface bacteria, freshwater sediments, survival, viability

INTRODUCTION

Sediment bacteria are acknowledged to play a critical role in benthic food webs, nutrient cycling and the decomposition of organic matter. Most research on sediment microbiology has concentrated on the upper few centimetres and the surface sediment/water interface (e.g. Jones & Simon, 1981; Jones, 1982). However, the last decade has seen deeper sediments receiving more attention, mainly due to concerns about aquifer contamination through industrial, agricultural and other human activities (see Reeves et al., 1995; Biovin-Jahns et al., 1996; Fredrickson & Onstott, 1996; Balkwill et al., 1997). Bacteria have now been described extensively in the subsurface environment, and have been shown to be closely involved in mineral formation and diagenic processes, some of which were initially thought to be of purely chemical origin. Bacterial activity is generally highest near the sediment surface, although there is
indirect evidence that microbial activity continues to considerable depths within the sediment (Jones & Simon, 1981; Jones et al., 1982; Parkes et al., 1994).

The environmental conditions that prevail within subsurface sediments are highly variable and are represented by different zones of redox potential. The type of bacterial activity varies with depth and is dependent on the zone encountered (Jones, 1985; Lovley & Chapelle, 1995). Activities associated with carbon dioxide production, methanogenesis, denitrification, sulphate reduction, iron reduction and production of volatile fatty acids have been detected at depths down to 503 m in subsurface marine sediments (Ekendahl & Pedersen, 1994; Phelps et al., 1994; Krumholz et al., 1997; Wellsbury et al., 1997). Furthermore, there are reports of the detection and/or cultivation of bacteria from sediments to depths > 500 m (Kennedy et al., 1994). However, aerobic organisms are most frequently studied due to ease of culture. Intact bacteria have been observed in marine sediment layers more than 4 million years old due to ease of culture. 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sulphate-reducing medium, were incubated under an anaerobic atmosphere generated by Oxoid gas-generating kits. For presumptive sporing anaerobes, Differential Clostridial Medium (DCM; Oxoid) was used. Nutrient agar (Oxoid) was supplemented with KNO₃ (0.1%) to determine the number of nitrate-reducing heterotrophs in sediment samples (Hardy, 1979). Plate counts were corroborated by most-probable-number (MPN) broth counts as follows. Nutrient broth was supplemented with KNO₃ to a concentration of 0.1%, and 4.5 ml aliquots were autoclaved in 13 cm test tubes each containing an inverted Durham tube. Five replicate serial dilutions were performed; 0.5 ml of each diluted sample was used to inoculate the broth. Tubes were examined for growth, gas production and generation of nitrite. Reduction of nitrate was detected using the reagent of Elliot & Porter (1971). Sulphate-reducing bacteria were enumerated by a MPN technique using Postgate medium B (Postgate, 1984) containing 20 mM sodium acetate with a headspace mixture of 80% H₂/20% CO₂ (Jones & Simon, 1984).

Fluorometric enumeration of bacteria. Acridine orange direct count (AODC) was performed as described by Fry (1990). Two fluorometric methods were used for viable counts. (a) The fluorescent formazan derivative CTC method of Rodriguez et al. (1992; detects respiratory activity), in which samples were diluted 1:2 with R2A broth (Reasoner & Geldreich, 1985) and 5-cyano-2,3-dicyethyl tetrazolium chloride (CTC) was added to a concentration of 3.5 mM. Samples were incubated statically at 20 °C for 4 h. (b) Sterilized 5-carboxyfluorescein diacetate (CFDA; detects esterase activity and denotes viability through enzyme activity and membrane integrity) was added to samples to a final concentration of 0.01 mM. Samples were statically incubated for 10 min at 20 °C followed by 15 min on ice. In both cases, the sample was filtered through a 0.22 μm black membrane and then stored in the dark before counting within 20 min. Negative controls comprised samples that were autoclaved or treated with membrane-filtered formaldehyde (to a final concentration of 4% for 1 h before incubation). Direct viable count (DVC) was done by a modification of the method of Kogure et al. (1987). Filter-sterilized piromidal acid, pipemidic acid and nalidixic acid were added to cultures to final concentrations of 10, 10 and 20 μg ml⁻¹, respectively, then incubated at 20 °C for 16–18 h with shaking. After incubation, samples were immediately stained with CTC and CFDA, and numbers of bacteria counted by microscopy. Negative controls comprised samples treated with membrane-filtered formaldehyde (to a final concentration of 4% for 1 h before incubation).

Microscopic counting strategy. For AODC, DVC, CTC and CFDA counts, stained cells were visualized with a Zeiss Axioplan fluorescence microscope using appropriate optical filters. Triplicate filters were wet mounted in low-fluorescence immersion oil (Leica) and counted within 20 min of staining. Counts were made in accordance with the recommendations of Fry (1990). Bacteria were assumed to be Poisson distributed; 40 randomly chosen microscope fields (or enough to count 400 bacteria) were scanned per filter, and bacteria-like objects counted. Only samples diluted >3000-fold, visualized within fields containing ≤50% particulate obstruction, were counted; no correction of counts for particulate obstruction was performed.

Transmission electron microscopy (TEM). Electron microscopy was carried out on deep sediment samples using a JOEL 100CX Temscan electron microscope. Previously frozen sediment (−70 °C) was allowed to thaw, and 2.5 microlitres of the melt water was removed and spotted directly on to a copper electron microscopy grid. Four grids were prepared for each subsection examined. All grids were osmium tetroxide fixed; two from each section were shadowed with chromium, and subjected to TEM at various magnifications.

Direct extraction of DNA from sediment. This method was derived from those of Selenska & Klingmuller (1991), Rochelle et al. (1992) and Smalla et al. (1993). Sediment samples (5 g wet wt sediment) were vortexed in 10 ml extraction buffer [0.12 M sodium phosphate buffer, pH 8.0; lysozyme, 5 mg ml⁻¹; sodium dodecyl sulphate, 1.5%; acid-washed polyvinylpyrrolidone (PVPP; 6%)], sonicated at 18 μm amplitude for 20 s on ice and then incubated for 1 h at 70 °C with frequent shaking. Following this, the samples were centrifuged at 2800 g, 4 °C for 15 min, and the supernatant was retained at 4 °C. This procedure was repeated twice without further addition of PVPP. The pooled supernatants were then centrifuged at 8000 g, 4 °C for 30 min. Proteins were precipitated from the final supernatant by the addition of 0.1 vol. 5 M NaCl and polyethylene glycol 6000 (to a final concentration of 15%) and overnight incubation at 4 °C followed by centrifugation (5000 g, 4 °C for 40 min). The pellet was resuspended and purified following ethanol precipitation (Sambrook et al., 1989).

PCR amplification. The PCR was carried out using a Perkin Elmer Cetus 480 thermal cycler and performed using primers pA, pD’ and pH’, specific to the eubacterial 16s rRNA gene (Edwards et al., 1989), and 1Af and 1100Ar, specific to the archaeal 16s rRNA gene (Embley et al., 1992). Prior to 'hot start', samples were serially diluted tenfold to achieve the optimal sample concentration for amplification. Template DNA was added to a standard PCR reaction mix, giving a total volume of 45 μl comprising 20 pmol of each primer, 10 nmol of each deoxynucleoside triphosphate (Pharmacia), PCR buffer [50 mM KCl, 10 mM Tris/HC1 (pH 8.3), 1.5 mM MgCl₂, Boehringer Mannheim]. Reactions were overlaid with 50 μl liquid paraffin (BDH) in thin-walled PCR reaction tubes (Anachem). The amplification reaction comprised an initial cycle of denaturation at 95 °C for 4 min followed by addition of 5 μl 1× PCR reaction buffer containing 0.5 U Taq DNA polymerase (Boehringer Mannheim) and 1 U Perfect Match polymerase enhancer (Stratagene) at 80 °C. This was followed by 30 cycles of denaturation at 95 °C for 1 min, annealing for 1 min at 55 °C and extension at 72 °C for 2 min with a final extension step of 72 °C for 10 min.

Cloning of PCR products. PCR products were ligated into the cloning vector pCRII provided in the TA-Cloning Kit (In-vitrogen), following the manufacturer's instructions. Recombinant plasmids were used to transform 50 μl TA One Shot Competent Cells by the manufacturer’s specified protocol. Transformants were plated on selective agar containing X-Gal (Boehringer Mannheim), subcultured and screened for the presence of an appropriate insert by plasmid mini-preparation-restriction analysis using the method of Close & Rodriguez (1982).

Sequencing. Plasmid DNA was extracted from clones using Qiaprep Spin columns (Qiagen). Cloned genes were sequenced using T7 polymerase (Pharmacia). Electrophoresis and reading of the sequences were performed on an automated laser fluorescence ABI 373A DNA sequencer (ALF; Pharmacia). The GenBank accession numbers for the sequences are: Wineub 010, AF045172; Wineub 012, AF045173; Wineub 019, AF045176; Wineub 027, AF045170; Wineub 043, AF045168; Wineub 076, AF045171; Wineub 082, AF045169; Wineub 115, AF045173; Wineub 117, AF045174; Winarc 007, AF045181;
Analysis of DNA sequence data. Sequences were analysed using the GCG suite of programs (Devereux et al., 1984) running on the SEQNET Facility at Daresbury (UK). Sequences were visually aligned with representative sequences from National Science Foundation Ribosomal Database Project (RDP; Larsen et al., 1993) and the GenBank Database (Benson et al., 1993) on SEQNET. Only unambiguously aligned base positions were used in the analysis. Analyses and measures of robustness were corroborated by DNASIST (PHYLIP version 3.5) using the Jukes & Cantor (1969) correction. Phylogenetic trees were constructed by the neighbour-joining method (Saitou & Nei, 1987). Bootstrap analysis was performed using SEQBOOT (PHYLIP 3.4) to evaluate the robustness of the inferred phenograms and CONSENSE (PHYLIP 3.4) (Felsenstein, 1993) was used to generate a consensus phenogram in conjunction with the 'neighbour-joining method'. Topology of phenograms was confirmed by maximum parsimony analysis.

Statistical analyses. The following analyses were performed where appropriate: MPN (Halvorsen & Zeigler, 1933); analysis of variance (probability of 0.01%; Fry, 1993); MINITAB 8.21, Minitab, State College, PA, USA; minimum significant difference (MSD, Tukey–Kramer method; Fry, 1993).

RESULTS

Intercomparability of cores

Visual examination of the varve stratification confirmed that the cores taken on the three occasions were representative of a typical vertical section of the sediment (Collins, 1977; Haworth, 1985). The data are derived from three separate 6 m Mackereth cores taken in the late spring/summer over a period of 15 months. The initial bacteriological survey compared direct, viable and culturable counts at different depths spanning the complete core (40 cm to 6 m). All three cores were comparable with respect to stratigraphy, culturable aerobic heterotrophic counts and AODC ($P<0.05$; data not presented). Data for the top 30 cm were derived from independent Jenkin cores from the same sample point. Measurements of carbon, nitrogen and water content confirmed those made by Pennington (1973), giving further confirmation of core comparability (data not shown).

Microscopic examination of the cores

Electron microscopy. The nature of the sediment prevented TEM procedures that employed resin embedding and sectioning. However, it was possible to visualize cells after freeze/thawing the samples and placing them directly on electron microscopy grids. A range of bacterial morphologies was observed by TEM at depths down to 2.3 m. Rod-shaped bacteria were observed in sediment from subsections 0-1 m, 1.5 m and 2.3 m. The dimensions of these ranged from $1.8 \times 0.3 \mu m$ to $3.8 \times 0.5 \mu m$ (Fig. 1a). In the case of the rod-shaped bacterium from 2.3 m (approximate size $1.5 \times 0.5 \mu m$), intracellular structures were visible (Fig. 1b). Coccolid bacteria undergoing division, ranging in diameter from 0.2 to 0.3 mm, were observed in subsections 0-12 m and 1.5 m (Fig. 1c, d). Bacteria with apparent prosthecate structures were observed at 2.3 m, with protosac lengths between 4 and 12 mm (data not presented). No stalk bands were observed along the length of the protosac in any of the cases. The viability of these organisms could not be determined at this stage.

Direct count. AODC was used to visualize and enumerate bacteria present in the sediment. In general, the bacteria observed were small and rounded when viewed in situ. The majority of cells stained green rather than orange with acridine orange and >90% remained particle-associated. No numerical correction was made to account for bacteria beneath particles; therefore the counts were consistent, but may underestimate cell numbers. Microcolonies were observed, although frequently, in all organic sections examined. Filamentous bacteria were observed only in the top 10 cm of the sediments and similarly no protozoa were observed in subsurface samples below a depth of 10 cm. Total counts were approximately $10^{12}$ cells (g dry wt sediment)$^{-1}$ for the surface sediments (Jenkin core), then declined significantly within the subsurface sediments, but remained consistent from a depth of 0.9 m to 40 m at $3.7-7.3 \times 10^9$ cells (g dry wt sediment)$^{-1}$, at which point a significant ($P<0.01$) drop to $<10^8$ (g dry wt sediment)$^{-1}$ occurred that was coincident with an increasing clay content of the sediments (Fig. 2a, b).

Viable count. The number of metabolically active bacteria was assessed by the application of CTC and CFDA as fluorogenic viability dyes (Fig. 2a, b). In addition, the DVC assay method of Kogure et al. (1987) was used to enumerate bacteria capable of demonstrating growth (Fig. 2b). CTC- and CFDA-fluorescent cells were easily distinguishable, with minimal non-specific staining occurring. Control samples, obtained by formaldehyde treatment and autoclaving, generated no visible fluorescence.

There was no significant difference between the number of CTC-reducing cells and the AODC throughout the depth of the core ($P<0.01$; Fig. 2a). By contrast, the CFDA-cleaving populations were high in upper sediments, being equivalent to total and CTC-reducing populations to a depth of 250 cm ($P<0.01$); thereafter, numbers decreased consistently and significantly from the AODC values with depth ($P<0.01$). The relationship between AODC and CTC-reducing cells was confirmed in the analysis of a later core complemented with bacterial enumeration from section of a suture Jenkin core (0–20 cm; Fig. 2b). AODC decreased rapidly with depth from a surface maximum of $1.48 \times 10^{12}$ (g dry wt sediment)$^{-1}$ to approximately $4.45 \times 10^9$ (g dry wt sediment)$^{-1}$ at 74 cm depth; thereafter, populations remained approximately constant in size (Fig. 2b). CTC-reducing populations again were not significantly different from the total populations ($P<0.01$).

The viability of these metabolically active populations
was investigated further by an adaptation of the DVC assay using CTC as the resolving fluorochrome (Fig. 2b). A 16–18 h incubation was performed in the presence of yeast extract and a cocktail of antibiotics inhibitory to cell division (Kogure et al., 1987). The total number of CTC-reducing cells decreased in all samples after incubation. However, cells capable of growth under these incubation conditions were clearly distinguishable as elongated rods. The number of substrate-responsive cells remained proportional to the pre-incubation AODC and CTC-reducing counts, cells demonstrating growth following incubation decreased rapidly in number with depth from the surface sediments until a depth of about 1 m. Thereafter, the population size remained approximately consistent with depth to 400 cm, below which numbers again declined through to the deepest sampling point (Fig. 2a, b).

**Culturable bacteria and fungi**

**Aerobic culture.** In a preliminary study, aerobic heterotrophs from all depths of a 6 m core were recovered by spreading on R2A agar and TSA at 50% and 5% strength (Fig. 3a). The number of culturable heterotrophic bacteria decreased significantly within the first metre ($P < 0.01$), from a surface value of $2.1 \times 10^7$ cells (g dry wt sediment)$^{-1}$ (not shown on graph) and continued to decline to approximately $10^3$ cells (g dry wt sediment)$^{-1}$ at a depth of 3 m. Below this depth, the culturable population declined slowly and was main-

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**Fig. 1.** TEM micrographs of chromium-shadowed bacteria: (a) rod-shaped bacterium recovered from 1.5 m (~ 2800 years deposition); (b) rod-shaped bacterium recovered from 2.3 m (~ 4250 years deposition), showing unidentified intracellular structures; (c) coccoid bacterium from 0.1 m (~ 37 years deposition); (d) sample from 1.5 m (~ 2800 years deposition) showing cells fixed at the time of apparent cell division. Bars, 1 μm.
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log_{10}[Bacterial count (g dry wt sediment)^{-1}]

7 8 9 10 7 8 9 10 11 12 13

Fig. 2. (a) Depth distribution of total bacteria using the AODC technique (■), and of metabolically active bacteria using CTC (▲) and CFDA (●) to 587 cm depth. (b) Depth distribution of total bacteria using the AODC technique (●) and the DVC viability assay. CTC-reducing cells were enumerated to indicate total viability (□). Substrate-responsive CTC-reducing cells (direct count; ▲) became classically elongated. The MSD_{0.01} bars indicate the mean significant difference (P<0.01).

tained at a low level [10^1–10^2 c.f.u. (g dry wt sediment)^{-1}] by 6 m. Regardless of the sedimentary horizon sampled, no significant difference in culture efficiency was observed between the aerobic heterotroph media (P<0.01; Fig. 3a). The exception was sediment agar, which recovered significantly lower numbers (data not presented). Low numbers of fungi and yeast were distributed throughout the sediments with no perceivable pattern. Colonies were selected arbitrarily from each medium at all depths and purified on the medium from which they were originally isolated. As these sediments were anoxic, the anaerobic growth capability of the isolates was tested. Isolates were subcultured on to nutrient agar supplemented with 0.1% KNO₃ and plates were incubated anaerobically under N₂/CO₂/H₂ (85:10:5) for 2 weeks at 20 °C. It was found that 98% (n = 150) of isolates were facultatively anaerobic.

Bacterial isolates were crudely classified as rods, cocci and branched chains by microscopy; >95% of the isolates were rods, with equal numbers of cocci and branched chains. Gram-staining was ambiguous for approximately 45% of the isolates. To a depth of 2 m, Gram-negative organisms were twice as abundant as Gram-positive ones, whereas below this depth Gram-positive and Gram-negative organisms were in equal proportions. It should be noted, however, that the gross numbers recovered below 2 m were reduced.

The presence of sporing bacteria was assessed. Surface sediments contained greater numbers of total culturable cells than sporing ones (P<0.01). However, in sub-surface samples there was no difference between the heat-resistant and heat-sensitive populations (Fig. 3b). The single exception was the deepest sample (5.9 m), where recovery was significantly higher after heat

Fig. 3. (a) Depth distribution of aerobic heterotrophic bacteria isolated on R2A agar (▲), 50% TSA (■) and 5% TSA (●). The MSD_{0.01} bar indicates the mean significant difference (P<0.01). (b) Distribution of bacteria recalcitrant to pasteurization at 80 °C. To select actinomycetes, samples were treated for 10 min before plating on M3 medium (▲). For aerobic heterotrophic bacteria, samples were plated on 50% TSA before (●) and after (■) 30 min pasteurization. The MSD_{0.01} bar indicates the mean significant difference (P<0.01) (MSD value calculated upon data from TSA plates only). (c) Distribution of anaerobic heterotrophic bacteria isolated on Differential Clostridial Medium (DCM; ■) and nitrate agar (●). Sulphate-reducing (●) and nitrate-reducing (▲) populations were also enumerated by MPN broth culture. The MSD_{0.01} bar indicates the mean significant difference (P<0.01) (MSD value calculated upon data from the DCM and nitrate agars only).
Bacteria in post-glacial freshwater sediments

Listeria monocytogenes

Mycobacterium malmoense

Arthrobacter globiformis

Listeria monocytogenes

Bacillus mycoides

Bacillus thuringiensis

Wineub 010 (5-2 m)

Bacillus subtilis

Bacillus pumilus

Wineub 027 (5-2 m)

'Bacillus macroides'

Wineub 019 (5-2 m)

Methanococcus jannaschii (outgroup)

High-G+C

Gram-positive

bacteria

Low-G+C

Gram-positive

bacteria

y-Proteobacteria

Legionella feeleii

Pseudomonas aeruginosa

R. solanacearum

Wineub 117 (4-9 m)

Wineub 076 (4-4 m)

R. picketti

Wineub 082 (4-9 m)

Wineub 043 (4-4 m)

6-Proteobacteria

Desulfovibrio desulfuricans

Wineub 012 (4-4 m)

JAP751 (unknown eubacterium)

Wineub 115 (5-0 m)

Methanococcus jannaschii

(outgroup)

Fig. 4. Phylogenetic relationships of eubacterial 16S rRNA gene sequences isolated from Windermere subsurface sediments (prefix Wineub). The depth from which the clones were obtained is shown in parentheses. Data for reference strains were obtained from GenBank or RDP (Larsen et al., 1993). R., Ralstonia (formerly Burkholderia). Base positions 107-180 and 220-316 were considered in the analysis (numbering based on E. coli; Brosius et al., 1978). Methanococcus janaschii was used as the outgroup. Bootstrap values were derived from 100 analyses; only values greater than 40% are shown at nodes. The scale bar represents base changes per nucleotide position.

treatment \((P<0.01)\). This strongly suggests that all culturable bacteria below 30 cm are spore-formers. Actinomycete populations consisted of only Microactinomycetaceae sp., identified on the basis of its highly characteristic colony morphology (Cross, 1981). Maximum concentrations were recovered from the surface sediments \((1.65 \times 10^6 \text{ (g dry wt sediment)}^{-1})\), with the number of isolates decreasing rapidly with depth; no actinomycetes were recovered below a depth of 74 cm (Fig. 3b).

**Anaerobic culture studies.** A variety of culture media were utilized to assess the sedimentary bacterial populations capable of anaerobic growth. Spread-plate and MPN broth methods detected denitrifying populations. Neither technique was found to be significantly more sensitive \((P<0.01); \text{ Fig. 3c}). Denitrifying heterotrophs were cultured from all depths, with maximal populations recovered from surface sediments \((3.46 \times 10^6 \text{ c.f.u. (g dry wt sediment)}^{-1})\). Culturable anaerobic heterotrophs were recovered at a frequency indistinguishable from that of the aerobic heterotrophs throughout the length of the core, dropping by approximately 5 orders of magnitude over the entire depth \((P<0.001)\) and being finally maintained at \(10^5 \text{ c.f.u. (g dry wt sediment)}^{-1}\) below 3 m.

Sporing anaerobic isolates were recovered from all samples. The pattern of culturability was comparable to that of the aerobic and anaerobic heterotrophs down to 4 m. Below this point the sporing anaerobes declined further, to \(<10^4 \text{ c.f.u. (g dry wt sediment)}^{-1}\). By contrast, sulphate-reducing bacteria were cultured only in the three uppermost samples: \(2.5 \times 10^5 \text{ c.f.u. (g dry wt sediment)}^{-1}\) in the surface sediments, approximately \(10^3 \text{ c.f.u. (g dry wt sediment)}^{-1}\) at a depth of 10 cm before declining to \(8.4 \text{ c.f.u. (g dry wt sediment)}^{-1}\) at 18 cm; Fig. 3(c).

**Extraction of DNA from core samples.**

Total DNA was extracted directly from all 14 subsamples. AODC indicated that the lysis efficiency of the prokaryotic population was approximately 98%. DNA recovered was of a suitable purity, when diluted to \(10^{-2}\) and \(10^{-3}\), to act as template for amplification by PCR. PCR amplification was carried out using both eubacterial and archaeal 16S rRNA primers.

**Eubacterial 16S rDNA amplification.**

A region of approximately 1.5 kb of the eubacterial 16S rRNA gene using the primer pair pA/pH' was successfully amplified from all samples (Edwards et al., 1989). These amplions were confirmed as eubacterial *rrn* genes by hybridization with digoxigenin-labelled eub338 probe (data not presented). In addition, as amplification and cloning procedures are more efficient with smaller products, amplification of a 528 bp region of the *rrn* gene was performed on selected samples (0.9 m to 5.2 m) using the primer pair pA/pD' (Edwards...
et al., 1989). Fourteen eubacterial 16S rDNA gene sequences, generated by both pA/pH' and pA/pD', were analysed from the deeper samples. Five of the sequences were found to be homologous to each other: Wineub 049/076 (pA/pD') and Wineub 117 (pA/pH') and Wineub 063/065 (pA/pD') were isolated from depths of 4.4, 5.0 and 5.2 m, respectively, using the primers indicated in parentheses.

Of the 14 cloned sequences, Wineub 010, 012, 019, 027, 076, 082 and 117 were found to be >90% homologous to previously described 16S rDNA sequences available in GenBank. Two major sequence groups were apparent (Fig. 4). The clones Wineub 010, 019 and 027 clustered within the low-G+C Gram-positive bacteria (100% confidence value from 100 analyses) and all were very closely related to Bacillus sp. Clones Wineub 076, 082 and 117 grouped within the β-Proteobacteria, with the closest relative for Wineub 076 and 117 being Ralstonia (formerly Burkholderia) picketti (bootstrap value 75%; Fig. 4). Wineub 082 and 043, by contrast, represent a distantly branched novel lineage within the Proteobacteria. The same comparison positioned Wineub 012 and 115 outside the Proteobacteria. However, further phylogenetic assessment of Wineub 012 showed it to have the greatest homology with JAP751 from the JAP504 cluster (bootstrap value 96%; Fig. 4), originally described by Rochelle et al. (1994).

Archaeal 16S rDNA amplification

The primers 1Af and 1100Ar (Embley et al., 1992) were used to amplify a 1.1 kb region of 16S rDNA specific to archaea. Products were obtained from all samples except 5.0 m and 6.0 m; this result was attributed to sample failure rather than indicating the absence of an archaeal community at these points.

Nine archaeal 16S rDNA clones were isolated from 3.4 and 5.9 m depth. They were of two general types, each showing highest identity to 16S rDNA sequences isolated from uncultured archaeal populations of the subsurface blanket bog peat of the Pennine high moors (UK) (Hales et al., 1996). A 350 bp region of the 1100 bp sequence of the nine archaeal clones was found to represent four different sequence types. Two, represented by Winarc 045 and Winarc 031, clustered with R10, the predominant methanogen sequence type found in Cum- brian blanket bog peat (Hales et al., 1996; bootstrap value 84%; Fig. 5). The two other sequence types, represented by Winarc 041 and 150, grouped with R17 (bootstrap value 100%), a minority sequence type found
in Cumbrian blanket bog peat (Hales et al., 1996). On the basis of distance matrix analysis, the R10-like sequences grouped within the Methanogenium group, with close association to Methanospillum hungatei (bootstrap value 45%). R17-like sequences were placed closer to the Methanosarcina group, the confidence of this placement being similarly low (bootstrap value 59%).

Sequences of the R10 group gave high homologies to the 16S rRNA genes from the endosymbiotic bacteria of the anaerobic protozoa Trimyema compressa and Metopus contortus, which are closely related to, but distinct from, the free-living methanogens (Fig. 5). No protozoa were observed throughout the extensive microscopic examination of the prokaryotic populations of these sediments. Indeed, there have been no reports of their isolation from these, or other, lacustrine subsurface sediments of Cumbria. However, the possibility that protozoal cysts present in the sediments may, after lysis, serve as a source of archaeal template could not be discounted due to possible presence of archaeal endosymbionts (Embley et al., 1992). For this reason, PCR amplification of protozoan small-subunit 18S rRNA (∼1.7 kb) was attempted using the primers of Embley et al. (1992); however, no products were obtained.

DISCUSSION

A study on sediments in Lake Constance showed that viable counts and potential activities exhibited by bacteria were a function of depth (Rothfuss et al., 1997). It was shown for post-glacial sediments (down to a depth of 7 cm and equivalent to 13000 years of deposition) that non-sporing heterotrophs were not present below 25 cm and that viable bacteria belonging to specific metabolic groups were present only down to 30 cm. Below this depth, viable heterotrophic bacteria existed only as heat-resistant spores, falling below the detection limit at 4–6 m. We present evidence, obtained through electron microscopy, culture, and cellular and molecular analyses, that post-glacial sediments of Windermere, dating from approximately 12000 years ago to the present day, contain viable bacterial populations comprising mainly sporing bacteria. Both the direct counts and total culturable heterotrophic bacteria recovered demonstrated an exponential decrease in numbers from the surface to a depth of approximately 0.8 m (1200 years). The culturable count then showed a slower decline to 3.0 m (4000 years); thereafter, numbers remained at a constant level. The paleolimnological history of Windermere has been studied extensively (Pennington, 1943, 1973, 1981, 1991) and those studies have shown that the composition of the sediment down to 0.8 m could have been influenced by extensive deforestation occurring in the catchment through early settlements (Vikings) to the extensive habitation of the present day (Pennington, 1991). The main effect of deforestation was to increase the input of sediment, with a subsequent reduction in lake volume. This is consistent with a rise in both organic carbon and phosphate in the sediments. The discontinuities in direct and culturable counts around 0.8 m depth may, therefore, be related to anthropological influences initiated around this period of time (approximately 1500 years deposition). The second but less defined discontinuity in culturable count at 3.0 m occurred in sediments of high organic content influenced by a period of climatic cooling and increased water run-off from the catchment.

In this study, pasteurization of samples prior to plating indicated that the majority of culturable aerobic bacteria were capable of sporulation. Whether the cells isolated were resuscitated from a state of dormancy cannot be determined, but the ability of Bacillus sp. spores to survive for exceptionally long periods is well documented (Kennedy et al., 1994). Isolates recovered aerobically must be capable of anaerobic growth if they are to exhibit in situ activity. Of a representative subset of the deepest aerobic isolates held in the culture collection, almost all were found to be facultatively anaerobic when cultured upon nutrient agar supplemented with 0.1% KNO3. Those isolates that failed to grow cannot be classified as obligate aerobes without further study as this medium cannot be considered universal for anaerobic organisms. Heterotrophs were isolated anaerobically with frequencies equivalent to those cultivated aerobically. It is possible that these independently obtained counts may represent the same subpopulations.

In this study, actinomycetes were only recovered to a depth of 74 cm (1000 years deposition), and Micromonospora sp. was the only actinomycete isolated. A further incubation period at 50 °C failed to promote the growth of any thermoactinomycetes. This confirms previous studies which found Micromonospora sp. populations in sediments down to 1 m depth in Thirlmere and Windermere (both in UK; Cross, 1981). However, in addition to Micromonospora, nocardioform actinomycetes and Streptomyces spp. were isolated from both water bodies, with the latter being predominant with increasing depth (Cross & Atwell, 1974; Johnston, 1972).

Postgate’s (1984) medium B for sulphate-reducing bacteria (SRB) was modified to include both acetate and lactate as carbon sources as a means of optimizing the recovery of SRB. The deepest sample to yield culturable SRB was from 77 cm. At this depth, sulphate is almost certainly depleted (Jørgensen, 1983). Therefore, it is unlikely that these isolates were active in situ with respect to sulphate reduction (Brandl et al., 1993). In Windermere, this can be inferred from the decreasing levels of acid-volatile sulphide below 20 cm (<1 mg g−1; Rowlatt, 1980). At the sample points analysed, no sequences were isolated by PCR that were closely related to any known SRB. Declining populations of SRB are frequently indicative of conditions more likely to favour active populations of methanogens. In fresh water, methanogenesis is responsible for the bulk of terminal metabolism under anoxic conditions (Capone & Kiene, 1988). The nature of the post-sampling handling used in these studies made it impractical to quantify methanogens by culture. However, examination of prokaryotic communities in the deepest core samples by targeting
16S rDNA revealed several sequences that were most likely to have been derived from methanogenic species (see below).

In the sediments of Windermere viable cells were detected by microscopy to depths of 5.8 m, representing deposition events over 10000 years ago. The numbers of CTC-reducing cells were consistently lower than the total bacterial numbers, but, in common with published work using acridine orange/INT (2-[4-iodophenyl]-3-[4-nitrophenyl]-5-phenyltetrazolium), values were usually within a single order of magnitude (Porter et al., 1995). The fact that the proportion of non-respiring bacteria in these sediments did not increase with depth implies that dead bacteria were not accumulating with depth.

To complement the use of the tetrazolium salts, CFDA was applied to the same samples to give an independent assessment of bacterial viability. While in the upper sediments CFDA and CTC counts were indistinguishable, with increasing depth the number of CFDA-cleaving cells decreased in comparison to the CTC-reducing populations, but CFDA-cleaving cells were still detectable in the deepest sample. The reason for this anomaly between techniques is unclear and, as is frequently the case with the analysis of heterogeneous environmental populations, data must be interpreted with caution. The protocol for the staining with CFDA differs from that of the CTC and DVC techniques in that it did not require incubation in the presence of a nutrient source. This being the case, counts of CFDA-cleaving bacteria may provide a more faithful representation of in situ metabolic activity. However, the sensitivity of these different assays can have a significant effect on interpretation. Porter et al. (1995) demonstrated that the efficiency of a range of fluorescent probes was variable for the assessment of physiological status of starved Escherichia coli. In the case of Windermere sediments, the heterogeneity of environmental samples and the variability of viability probes made their use representative of populations rather than as a total viability count.

The number of substrate-responsive bacteria (DVC) constituted approximately 90% of the total bacterial count in the surface sediments. With increasing depth, the number of substrate-responsive cells decreased, yet positive cells remained detectable in the deepest samples. In all samples, populations were observed that were metabolically active on the basis of CTC reduction yet did not demonstrate cell growth and elongation. Roszak & Colwell (1987) compared the DVC and microautoradiographic methods and observed the same effect. It is possible that these non-responsive, yet viable cells may be unable to grow due to sublethal injury (Gurijala & Alexander, 1988). More likely, as with the other enumeration techniques applied in this work (except AODC), the DVC is subject to bias; the carbon source used and the antibiotics applied will favour the enumeration of some species more than others (Fry, 1990). It is interesting that the numbers of substrate-responsive bacteria more closely mirror the counts of CFDA, and thereby the bacteria active in situ, than do the counts obtained by the direct CTC viability assay. Further investigation to combine the DVC with CFDA or fluorescent in situ hybridization (FISH) would be desirable to confirm any possible correlation between the activity of esterases and the ability to demonstrate growth.

The eubacterial 16S rDNA gene sequences isolated in this study, although limited in number, were diverse in type, being distributed throughout both the Gram-positive and Gram-negative groups. However, similar sequences were observed in different levels within the core (e.g. Wineub 010 and 027). The isolation of rrs genes of high homology to those of Bacillus species (which are frequently recalcitrant to lysis) suggests that the DNA extraction procedure provided a template for PCR that is broadly, but not unequivocally, representative of the bacterial community. The recovery of Gram-positive sequences reported here is commensurate with the findings of several other researchers who have demonstrated the prevalence of culturable Gram-positive organisms in sediments to depths of 500 m (Gerhron et al., 1984; Reeves et al., 1995).

Rochelle et al. (1992) inferred the dominance of subsurface bacterial populations by species of the Proteobacteria (80%) from pristine marine samples. Similarly, proteobacterial sequences were prevalent in Windermere sediment, with identical sequences being isolated from separate sample points down to depths of 5.2 m (10000 years; e.g. Wineub 076, 117; Fig. 4). The isolation of sequences highly homologous to those of Ralstonia (formerly Pseudomonas and then Burkholderia) spp. is particularly interesting since these correlate with the reported isolation of species of the Pseudomonadaceae, identified phenotypically and genotypically, from these sediments (data not presented).

Independent isolation of sequences of novel delineation is presented in this work, corroborating novel sequences previously described by Rochelle et al. (1994) and Hales et al. (1996). The isolation of a sequence, Wineub 012 (from a depth of 4.4 m, 8000–9000 years), of high sequence similarity, and equivalent delineation, to the clones of the JAP504 cluster as reported by Rochelle et al. (1994) from deep marine sediments, is very significant. Independent recovery from freshwater sediments implies that these sequences represent organisms from a single group with wide-ranging phenotypic properties and a wide geographical distribution.

Several archaeal sequences were recovered from a depth of 3.4 m (7000 years deposition) and 5.8 m (11000 years deposition), with one sequence common to both depths (Winarc 038, 057 and 123). Sequences were recovered that were similar to the novel clones isolated from Pennine peat by Hales et al. (1996). This was perhaps not unexpected, as this sediment environment, being permanently anoxic, was predicted to contain methanogenic species. 16S rRNA sequence identities do not guarantee species identity as determined by DNA–DNA hybridization (Fox et al., 1992), nor do they imply...
functional activity. Nevertheless, many physiologically specialized microbial groups (the methanogens among them) are phylogenetically coherent or restricted to a limited number of assemblages (Stahl et al., 1988; Stahl & Amman, 1991). The point of delineation of the archaeal clones amongst other reference methanogenic sequences implies that their source organisms possess methanogenic properties. The description of methanogens presented in this study remains qualitative but suggests that, for this system, methanogenesis is the final important process in anaerobic mineralization, given the absence of sulphate-reducing bacteria. Future studies should focus on a quantitative approach that would allow a direct comparison with the better-studied deep-sea sediments, where different processes predominate; for example, the Japan Sea, where sulphate-reduction processes were detected at depths > 500 m (Parkes et al., 1994).

The recovery of gene sequence information from DNA templates does not necessarily indicate in situ cellular activity. It is possible that the bacteria have lysed and naked DNA is forming the target template for PCR, or that their cells have retained structural integrity in a state of death or dormancy. However, coupled with observations on viability, culturability and direct microscopy with non-specific dyes, the gene sequence information reported here implies that the deep sediments of Windermere do contain active bacteria derived from those deposited during sedimentation events during the last Ice Age in Britain.

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