14. GENETIC EVIDENCE FOR ENDOLITHIC MICROBIAL LIFE COLONIZING BASALTIC GLASS/SEAWATER INTERFACES1

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ABSTRACT

The majority of the Earth’s shallow crust is composed of basalt that erupted on the seafloor and was subsequently altered by chemical exchange with seawater. One aspect of this alteration is the replacement of glass by secondary minerals, including clays. Petrographic thin sections from ODP Holes 504B and 896A revealed characteristic patterns of pitting (channels) at the interface of fresh basaltic glass and secondary clay. Fluorescent dyes that bind specifically to nucleic acids (Hoechst 33342, PO-PRO-3, and Syto11) were used to examine thin sections for evidence of cellular life. Independent experiments with three dyes indicated the presence of particulate nucleic acids at the interface of altered and unaltered glass, particularly at the distal tips of channels. Organic material was extracted from crushed basalts from Holes 504B and 896A and examined for the presence of microbial DNA by a polymerase chain reaction (PCR) assay. This technique detects specific ribosomal RNA genes that are present in all cellular life-forms. The PCR assays demonstrated the presence of prokaryotic genomic DNA in the rock extracts. Among the geochemical reactions that could provide energy for biological processes are the oxidation of iron, manganese, and sulfur. Electron microprobe analyses of areas of glass that appear to be affected by microbial action have low iron relative to the fresh glass. Also, elements that are consistent with the presence of cellular life, phosphate and potassium, were elevated in the channels (P is 0.02 wt% in glass and up to 1.4 wt% in channels and K is 0.01 wt% in glass and up to 2.0 wt% in channels). These data raise the possibility that chemical transformations of basalt at the seafloor are mediated by microorganisms.

INTRODUCTION

Substantial evidence supports the view that microorganisms can effect widespread geochemical changes in the atmosphere, hydrosphere, and sediments. In the lithosphere, in contrast, the significance of the influence of microbial life on chemical change is unknown. Thus, it is of considerable interest that microbial communities associated with basalt have been reported (Thorseth et al., 1992). The alteration of basalts to palagonite has been viewed as a physical/chemical process (Staudigel and Hart, 1983; Crovisier et al., 1987; Thorseth et al., 1991). Evidence that microorganisms participate in this process could precipitate a fundamental shift in scientific perceptions of this problem.

The surface area of pillow basalts in contact with seawater is vast. Water circulation through the ocean crust results in chemical exchange that modifies seawater chemistry and leaves enormous accumulations of basal alteration products. The presence of microorganisms at the interface between fresh basaltic glass and seawater raises the possibility that microbial communities may influence chemical exchange in this global arena. Of particular interest are the possibilities that microbial life might control both rates of alteration processes and the composition of products. From a biological standpoint, the temperature range, pressure range, substrates, and evolutionary origins of these putative microbial communities are of keen interest.

Previous studies have implicated microorganisms in various glass transformations (Duff et al., 1963; Webley et al., 1963; Silverman, 1979; Krumbien, 1983; Thorseth et al., 1992; Palmer and Hirsch, 1993; Furnes et al., this volume). However, it has not been established whether the mineral transformations provide the energy for microbial growth. Alternately, the minerals could be colonized by biofilms that derive their energy from other sources. The latter is possible because the rock surfaces interface with active ecosystems. The constituents of the microbial communities have not been identified, leaving doubt as to whether mineral transformations are coupled to the activity of specialized microbial communities.

Microbial identification commonly limits investigations in microbial ecology because many bacterial species cannot be grown in culture. The introduction of molecular methods to microbial ecology has encouraged the study of communities of organisms that are not cultivable (Ward et al., 1992; Giovannoni and Cary, 1993). The 16S ribosomal RNA genes have played a pivotal role in these investigations. Ribosomal RNA databases include sequences from more than 1000 strains of Bacteria (eubacteria) and Archaea (archaeabacteria), making it possible to accurately identify unknown genes by phylogenetic methods. When genes from novel organisms are encountered in these investigations, phylogenetic information commonly indicates biochemical potential. For example, organisms such as methanogens, sulfur-and iron-reducing bacteria, and extreme thermophiles can be identified by characteristic signatures in ribosomal RNA genes. Furthermore, ribosomal RNA gene sequences can be used to construct oligonucleotide probes that hybridize specifically to the nucleic acids of target organisms. Probes of this type are used for the microscopic identification of microorganisms and to measure the abundance of specific target sequences in mixtures of nucleic acids that have been extracted from natural systems (Stahl et al., 1988; DeLong et al., 1989).

1Unlike its initial appearance in The Ocean Drilling Program (ODP) Hole 896A, drilled during Leg 148, Fluorescence imaging studies with three dif-
SAMPLES AND METHODS

Samples

Three samples from Hole 896A and Hole 504B from the Costa Rica Rift were used for this study. The two samples from Hole 896A were collected during Leg 148 and the sample from Hole 504B was collected during Leg 70 of DSDP. Sample 148-896A-7R-1, 52–55 cm (Piece 10), from 247.92 mbsf is a basalt pillow margin at the base of petrologic unit 13 (Shipboard Scientific Party, 1993). A 250 mbsf the temperature is estimated to be about 60°C (Shipboard Scientific Party, 1993). The sample has a 1-cm glass rim that contains about 5% plagioclase phenocrysts and 1% olivine phenocrysts. The glass has been partially replaced with palagonite, the zeolites phillipsite and possibly chabazite, and 1- to 5-µm grains of pyrite. Where fractures penetrate the glass, the glass is typically altered to clay, and along the boundary between the glass and the clay we observed irregular pits and channels that we suspected were produced by microbes (Fig. 1). This shipboard thin section was stained with a dye that indicates the presence of DNA.

Three dyes were used to stain separate thin sections. Hoechst 33342 was used on Sample 148-896A-7R-1, 52–55 cm (Piece 10). Six thin sections of Sample 148-896A-7R-1, 47 cm (Piece 1) were prepared at the Smithsonian Institution using techniques that were expected to preserve biological material in the rocks—no water or ultrasonic cleaners or high temperatures were used. The sample was obtained from the curated collection of ODP cores at Texas A&M University, vacuum impregnated with epoxy and cured overnight at room temperature. The sample was then cut in half with a diamond saw that was lubricated and cooled with Texaco almag oil and rinsed in 1:1 trichloroethane. After being polished on a 600-mesh diamond wheel that was lubricated with alcohol (95% ethanol), the surface was glued to a glass slide with Epo-tek 301 epoxy, and cured overnight at room temperature. The sample was then cut in half with a diamond saw, leaving a 0.5-mm-thick wafer of rock attached to the slide, which was polished with 6-µm diamond paste and then with 1-µm diamond paste. The thin sections were swabbed with alcohol to remove the paste.

Electron Microprobe

Ten-element chemical analyses of the glass, etch pits, and channels were done with the nine-spectrometer ARL microprobe at the Smithsonian Institution Division of Mineral Sciences. The standard used for the calibration were VG-2 (USNM 111240/52) for SiO2, Al2O3, FeO, MgO, and CaO. The standard for P2O5 was apatite (USNM 104021), for MnO was manganite, for Na2O was Kakanui anorholase (USNM 133868), and for K2O and TiO2 was Kakanui hornblende (USNM 143965). Counting time was 10 s on each element and backgrounds for all elements except SiO2 were calculated based on the background values for the elements on pure SiO2. The background for SiO2 was based on the background at the Si peak position on pure Al2O3. Standard glass A-99 (Jarosewich et al., 1980) was analyzed to check the calibration (Fisk et al., this volume).

Staining and Epifluorescence Microscopy

Three dyes were used to stain separate thin sections. Hoechst 33342 was used on Sample 148-896A-7R-1, 52–55 cm, and PO-PRO-3 and Syto11 were used on Sample 148-896A-12R-1, 0–4 cm. These dyes bind specifically to nucleic acids-Hoechst 33342 and PO-PRO-3 to DNA and Syto11 to DNA and RNA. Three dyes were used because they fluoresce at different wavelengths (Hoechst 33342 is blue, PO-PRO-3 is orange, and Syto11 is green) and therefore elimi-
nate the possibility that fluorescence observed in the thin sections is from minerals within the thin section or materials used in preparing the sections. Dyes were obtained from Molecular Probes Inc., Eugene, OR.

For each dye the staining procedure was slightly different. Sample 148-896A-12R-1, 0-4 cm, was fixed and dehydrated to permeabilize cells to DNA dyes. This was accomplished by immersing sections in a 1:4 mixture of formalin and 50% ethyl alcohol for 10 min, followed by dehydration for 10 min in 80% ethanol, 10 min in 100% ethanol, and drying in air at room temperature. This thin section number 3 of Sample 148-896A-12R-1, 0-4 cm, was stained with the cyanine dye PO-PRO-3 at a concentration of 10 µM in TE buffer (10 mM Tris × HCl pH 8.0, 1 mM EDTA). This section was stained for 30 min, followed by washing for 30 min at room temperature in TE buffer, and mounting in Citifluor (Citifluor LTD, London).

This thin section number 5 of Sample 148-896A-12R-1, 0-4 cm, was stained for 15 min at room temperature with a 50-mM solution of the dye Syto11 in TE buffer, followed by 5 min of washing. The section was then air dried and mounted.

The dyed sections were examined at 400× and 1000× magnifications using a Zeiss Standard 18 epifluorescence microscope equipped with a mercury vapor excitation source. The dyes and filter sets were as follows: Hoescht 33342, excitation band pass 365, dichroic mirror FT395, barrier long pass LP397; PO-PRO-3, excitation band pass 540, dichroic mirror FT 605, barrier long pass 565 nm; Syto11, excitation band pass 480, dichroic mirror FT 505, barrier long pass 535 nm. After photographing the fluorescence produced by the incident ultraviolet light, thin section number 3 of Sample 148-896A-12R-1, 0-4 cm, was treated with 50 µg/mL Dnase I (a DNA-degrading enzyme; Sigma, St. Louis, MO) for 24 hr.

**DNA Extractions**

Two samples (70-504B-48-2, 120-124 cm, and 148-896A-12R-1, 0-4 cm) were treated to remove contaminating DNA from rock surfaces. Single pieces of rock (2.5 g and 12.4 g, respectively) were first irradiated with ultraviolet light (λmax = 254 nm) at 400 J/m² per side, with rotation of the rock 6 times to irradiate each side. This level of irradiation is sufficient to cause approximately one thymidine dimer pair to form per 100 base pairs of DNA (Setlow, 1966), and thus render surface-contaminating DNA inactive to replication by DNA polymerases. Surface irregularities of the rocks might protect contaminating DNA in cells from damage from ultraviolet irradiation. To remove possible cell contaminants from rock surfaces, therefore, the samples were tumbled for 20 min in a DNA-free rock tumbler in the presence of dry, coarse, carborundum grit that had been baked at 220°C for 48 hr to destroy macromolecular contaminants, including DNA. Using forceps that had been heated to eliminate contaminating DNA, the rocks were removed and placed in sterile plastic ware. Rocks were crushed to sand using a mortar and pestle in a laminar flow hood. Organic material was extracted in lysis buffer that contained high salt and EDTA concentrations (100 mM EDTA, 400 mM NaCl, 0.75 M sucrose, and 50 mM Tris × HCl pH 9.0). Proteins were degraded with proteinase K, releasing DNA. The detergent sodium dodecyl sulfate was added to 1% and proteinase K, an enzyme that hydrolyzes proteins, was added to 100 mg/mL, followed by incubation at 37°C for 30 min, then 55°C for 10 min. Lysate was transferred to polypropylene centrifuge tubes, extracted with an equal volume of phenol/chloroform/isoamyl alcohol (25/24/1), and again extracted with an equal volume of CHCl3/IAA (24/1). The purified nucleic acids were precipitated with 2 volumes of 100% ethanol, 0.1 volume 2 M sodium acetate pH 5.2, and resuspended in aqueous solution.

**Polymerase Chain Reaction**

Several sets of small subunit ribosomal DNA primers were used to examine rocks for the presence of microbial genomic DNA. The bacterial primer 27F and the universal 16S ribosomal DNA primer 1492R were used to selectively amplify a 1.4-kb fragment of bacterial 16S ribosomal RNA genes (Giovannoni, 1991). The Arch21F and the universal 1492R primer were used to selectively amplify the same DNA region from Archea (Giovannoni et al., 1988). These two primer sets specifically amplify 1.4-kb regions from prokaryotes of the Archea and Bacteria domains, respectively. Universal small subunit ribosomal DNA primers (1492R and 519F; Lane, 1991) were used to amplify ribosomal DNAs from rock extracts using polymerase chain reaction (PCR). Reactions contained 0.1% of the rock extract, 5% acetamide, 200 µM of each primer, 1.5 mM MgCl₂, 200 µM total nucleotide triphosphates, 1X Taq reaction buffer containing 500 mM KCl, 100 mM Tris × HCl pH 9.0, and 1% TritonX-100 (Promega Corporation, Madison, WI), and 2.5 units Taq DNA polymerase (Promega) in a volume of 100 µL. Thirty-five cycles were performed in a thermocycler (MJ Research, Watertown, MA) using the following conditions: denaturation at 94°C for 1 min, annealing at 55°C for 1 min, and elongation at 72°C for 3 min. Unreacted oligonucleotides and deoxynucleotide triphosphates were removed using a Qiagquick PCR purification kit (Qiagen, Chatsworth, CA).

**Oligonucleotide Probe Hybridization**

An oligonucleotide probe specific for *Archaeas* (915R; Stahl and Amman, 1991) was hybridized to PCR amplification products. The primer terminus of the probe (0.5 mg) was labeled with 100 µCi of [32P]-dATP (New England Nuclear Corp., Boston, MA) using T4 polynucleotide kinase (Promega) as previously described (Giovannoni et al., 1988). For dot blots 100 ng of each PCR amplification product was denatured in 0.2 M NaOH and applied to Zetaprobe membranes (Bio-Rad Laboratories, Richmond, CA) using a dot blot manifold (Minifold I SRC 09690; Schleicher & Schuell, Keene, NH). The membranes were dried at 80°C for 15 min and UV cross-linked (120 J/m²) in an ULTRA LUM UV radiation source. Membranes were prehybridized for 15 min at room temperature in 10 mL of hybridization buffer (1 mL EDTA, 0.5 mM NaH₂PO₄, pH 7.2, 7% SDS). The solution was replaced with 6 mL of fresh buffer containing 5 ng of [32P]-labeled DNA probe, and the membranes were incubated at room temperature for 18 hr. Membranes were washed in two changes of wash buffer (0.2X SSPE, 0.1% SDS) at room temperature for 15 min. A final stringency wash was done at 45°C for 30 min. Hybridization of the probe was detected with X-ray film.

**RESULTS**

**Electron Microprobe**

A profile of 56 chemical analyses across a fracture and the surrounding etched glass is shown in Figures 1 and 2. The first nine analyses (Table 1) and the last five analyses in the profile are of fresh glass and appear to be unaffected by the processes that produced the pits and etch trails. This fresh glass is chemically similar to other glass found at this depth in the hole (Fisk et al., this volume). Within the area of etched pits and channels, the composition of the glass is highly variable (Fig. 2), which suggests that the processes that are affecting the glass are operating on a scale of less than 2 µm—the beam diameter of the microprobe analyses.

Several differences between fresh and altered glass are evident. Al₂O₃ and K₂O are significantly enriched in that altered glass, and
FeO, MgO, and CaO are significantly lower in the altered glass than in the fresh glass. These differences indicate that the glass is being broken down and that FeO, MgO, and CaO are being preferentially removed from the altered glass. The sink for the CaO could be the calcium carbonate that cements the fragments of this breccia. Both FeO and MgO may be taken up by clays, as the clay in Hole 504B has 15% to 20% FeO and 18% to 24% MgO (Noack et al., 1983; Alt et al., 1985).

There are several significant correlations between elements within the profile. For example, high K2O is associated with high Al2O3 and low MgO, MnO, and FeO are also strongly correlated. Al2O3 has a strong negative correlation with MgO, FeO, and CaO, which may result from the immobility of Al relative to Mg, Fe, and Ca during the breakdown of the glass. Na2O, TiO2, and P2O5 are not strongly correlated with other elements in the chemical profile (Fig. 2). For Na and P, this may be due to their high solubility and mobility in the fracture, but Ti is considered insoluble and immobile so reason for the absence of a correlation with Al is not clear.

### Staining and Epifluorescence Microscopy

The dyes used in these experiments form complexes with greatly enhanced fluorescence quantum yields after binding specifically to DNA and/or RNA. The fluorescence quantum yield of Syto11 increases 10-fold after binding to DNA or RNA. The cyanine dye PO-PRO-3 exhibits a 20- to 200-fold increase in fluorescence upon binding to DNA. Hoescht 33342 binds specifically to adenine/thymidine-rich regions in the minor groove of DNA, with a 30-fold increase in fluorescence (Kapuscinski, 1990). The intensities observed (Fig. 3) are not caused by background fluorescence of the uncombined dye, which was checked by examining areas of the thin section that were free of cracks but where dye was present. Rock samples were studied by fluorescence microscopy before staining, and it was determined that the samples exhibited weak fluorescence in the blue and green regions of the spectrum without staining; however, no autofluorescence of rocks was observed in the orange and yellow parts of the spectrum.

The thin sections of Sample 148-896A-7R-1, 52-55 cm (Piece 10) were stained using PO-PRO-3 and Syto11, two dyes with high affinities for nucleic acids and enhanced fluorescence yields in the presence of nucleic acids. The dyes indicated that DNA was present, but of the two, PO-PRO-3, which exhibits an orange fluorescence upon binding to DNA, was the most informative.

Hoescht 33342 was used to stain one thin section from Hole 896A (Pl. 1, Fig. 1). Hoescht 33342 binds specifically to the minor groove of DNA molecules, with a resulting enhancement of fluorescent quantum yield in the blue region of the spectrum. Hoescht dyes are similar to DAPI, which was used by Furnes and coworkers (this volume) to indicate the presence of DNA in Sample 148-896A-11R-1, 73-75 cm, but are somewhat more specific to DNA and usually less bright. One drawback of Hoescht 33342 was the presence of blue autofluorescence in unstained rock. This background autofluorescence was generally more uniform in appearance and less bright than the fluorescence associated with regions of alteration after staining with Hoescht 33342. However, significant blue autofluorescence of very uniform appearance was commonly associated with cracks in the sections, leading us to seek additional DNA stains with emission maxima in parts of the spectrum that are not subject to interference from autofluorescence.

The orange fluorescence of PO-PRO-3 was confined primarily to regions of rock alteration. Basaltic glass exhibited almost no PO-PRO-3 staining (Pl. 1, Fig. 2). Staining of channels was vivid and decreased inward from the glass/alteration interface toward the fracture. The intensity of staining with PO-PRO-3 was consistent with the presence of DNA, but less than the fluorescence observed in control experiments with intact bacterial cells, suggesting that the DNA present in the rock preparations was partially degraded. Cell-like
structures of approximately 1 mm diameter were occasionally observed, but in general cellular morphologies could not be discerned. Fractal-like structures elsewhere in the rock did not stain, nor did particulates associated with the rock or cracks in the rock. DNase treatment resulted in a decrease in fluorescence, but fluorescence was not eliminated. In control experiments, E. coli cells treated with DNase 1 lost most of their PO-P-3 fluorescence. The limited effect of DNase 1 on rock sections stained with PO-P-3 may be attributed to the poor penetration of DNase 1 into the channels and pits where the fluorescence was strongest. The diffusion of DNase 1 through clay alteration products may be limited by its molecular weight (31,000 daltons), which is much greater than that of the dyes used in these experiments (<1000 daltons).

With Sytol 1, as with the other dyes, significant staining of channels and alteration products was observed. However, with Sytol 1, other structures within the rock, including particulate matter, were also stained brightly (Pl. 1, Figs. 3, 4). A combination of interference from autofluorescent particles and endogenous green fluorescence of the rock rendered the signal from Sytol 1 somewhat equivocal, but nonetheless consistent with the possibility that DNA is present in channel regions and alteration products. In summary, all three nucleic acid dyes revealed evidence of DNA in basaltic glass alteration products and supported the highest concentrations of DNA at interfaces between basaltic glass and alteration products, particularly in channels.

DNA Extractions

DNA was extracted from Samples 70-504B-48-2, 120–124 cm, and 148-896A-12R-1, 0–4 cm, using procedures that were developed for the isolation of DNA from natural samples. A major concern was the possibility that rock samples were contaminated with DNA from drilling muds, seawater, or handling after retrieval. To minimize the possibility of carrying contaminating DNA forward into DNA extraction protocols, samples were decontaminated as described above. Control experiments indicated that extractions performed by the same procedure in the absence of rock were not contaminated with detectable DNA as assayed by polymerase chain-reaction amplifications of ribosomal RNA genes. However, extracts of rock samples did contain ribosomal RNA genes that could be amplified.

Polymerase Chain Reaction

Several different sets of oligonucleotide primers for small subunit ribosomal RNA genes were used to examine rocks for the presence of microbial genomic DNA. The oligonucleotide primer sets used were of three types, either universal for all cells, specific for the Bacteria, or specific for the Archaea. Small subunit ribosomal RNA genes are about 1540 bases in length. The universal primers we employed span an internal region of the gene which is about 1000 bases long. Both of the specific sets of primers we used span nearly the entire gene, approximately 1400 bases. Positive results indicating the presence of ribosomal DNA genes were obtained only with the universal primers. Neither set of specific primers yielded products from rock extracts in amplification reactions.

The universal ribosomal RNA primers 1492 reverse and 536 forward were successfully used to amplify a 1-kb fragment from both rock samples (Fig. 3). The amplified DNA is probably microbial in origin because eucaryotic ribosomal RNA genes are significantly larger for the same domain and are thus easily detectable by agarose gel electrophoresis.

Two factors may explain the amplification of these shorter DNA regions with universal primers. Smaller DNA target regions often amplify more efficiently. This phenomenon is enhanced considerably when low molecular weight DNA is used as a template. Thus, both the fluorescence data and the PCR data can be interpreted as evidence for partially degraded DNA in rock samples.

DISCUSSION

Electron Microprobe

The chemical differences of the fresh glass and the associated etch pits and tracks (Figs. 1, 2) are clearly not the same as chemical changes caused by secondary alteration of basalt glass. For example, previously reported palagonite in Sample 70-504B-4R-5, 87–89 cm, has less SiO2 and Na2O, more MgO and FeO, and about the same Al2O3 as the glass from which it formed (Noack et al., 1983). The area affected by secondary processes in Sample 148-896A-12R-1, 4–7 cm, however, shows no significant change in SiO2, a slight increase in Na2O, significant decreases in FeO, MgO, and CaO, and a significant increase in Al2O3 relative to the surrounding glass.

The secondary minerals that are prevalent at Hole 896A are smectite clays, iron hydroxides, aragonite (and some calcite), zeolite (philolipsite) and pyrite (Shipboard Scientific Party, 1993). None of these minerals have chemistries that could account for the chemical differences between the fresh and altered glass. Furthermore, these miner-
The clay produced in the 150°C experiment is high in Mg and low in Ca. Alkalis are not observed in the amorphous area around the fracture (Fig. 1). The chemistry around the fracture indicates that none of these minerals is present, even as submicroscopic grains.

The experimental alteration of oceanic basaltic glass by seawater at 70°C and 1-atm pressure results in the loss of Si and Ca from the glass and the uptake of Mg by the glass (Seyfried and Bischoff, 1979). At 150°C and 500 bars the effects are similar but accelerated. The clay produced in the 150°C experiment is high in Mg and low in Si and Ca relative to the original glass and is similar to clays that are the natural alteration products of basalt glass (Melson and Thompson, 1973) although not all clays are as high in Al and low in Mg as these. The clays produced in the experiments, however, are distinctly different from the areas of alteration analyzed by the microprobe in Sample 148-896A-12R-1, 0-4 cm. This suggests that the form of alteration observed in the ODP samples in this study is different from the chemical alteration of basaltic glass to form clay that has been extensively described in the literature.

The high K_2O of the altered glass associated with “microbial” pits and tracks is interesting because the alteration of K_2O-poor basalt glass results in clays that are also K_2O-poor. K_2O-rich minerals form in Hole 504B but they are associated with veins and the rare replacement of plagioclase feldspar with K-rich feldspar. Where K-rich clays exist, they always have less than 10% Al_2O_3 but at stated above, in the altered sections of the samples that we studied there is a strong correlation of K_2O with Al_2O_3 and Al_2O_3 reaches 22 wt%.

The elevated P_2O_5 in some areas of the altered glass is interesting because it has been recognized for some time that the submarine weathering of basalt results in an increase in the P_2O_5 content of the basalt (Miyashiro et al., 1969; Hart, 1970). The P_2O_5 increase during weathering is interesting in light of the fact that seawater is extremely low in phosphate (about 3 × 10^{-6} mol/L at 2000 m depth). The sink for phosphorus in the basalt appears to be calcium carbonate in veins that may have as much as 0.8 wt% P_2O_5 (Laverne, 1983). The areas within Sample 148-896A-12R-1, 0-4 cm, that are high in phosphorus are not elevated in calcium, which indicates that carbonate is not forming in the areas of altered glass. Elevated phosphorus in this area occurs in the microprobe analyses with the lowest total oxides, which suggests additional elements were present that were not analyzed. These could be hydrogen and oxygen in water, carbon, nitrogen, sulfur, or chlorine. In future analyses we will quantify the amount of oxygen, sulfur, and chlorine.

An additional feature of the microprobe analyses is the strong correlation of FeO and MnO (Fig. 4). During the process of alteration of the glass in the area shown in Figure 1 both FeO and MnO are removed from the glass (Fig. 4, FeO-MnO). This is interesting because both Fe and Mn occur primarily in their reduced state (Fe^{2+} and Mn^{2+}) in basalt glass and in their oxidized state (Fe^{3+} and Mn^{4+}) in seawater, and the oxidation of Fe and Mn is a possible energy source for the diagenesis of organic matter in marine sediments (Froelich et al., 1979).

### CONCLUSIONS

Rock cores are among many scientific samples that cannot be manipulated to the complete satisfaction of investigators. In the case of the basalt cores that were the principal subject of this investigation, a major concern is that contamination occurred during the recovery process. It may never be possible to rule out this possibility, absolutely, with a single experiment. Therefore, our approach was to develop several lines of experimental evidence that can provide evidence for or against the presence of microbial life at interfaces between basaltic glass and alteration products.

Two experimental approaches were used to examine basaltic glass samples for the presence of life. In the first set of experiments, fluorescent stains, which exhibit enhanced fluorescence when bound to nucleic acids, were used to localize nucleic acids, and particularly DNA, in thin sections of basaltic glass. In the second set of experiments, basaltic glass samples were surface-treated to remove contaminating DNA, crushed, and organic material extracted to chemically isolate any nucleic acids that might be present. Both sets of experiments indicated the presence of DNA in basaltic glass samples. Because the amounts of DNA extracted from rock samples were too small to be chemically measured by direct means, a genetic assay, gene amplification by the polymerase chain reaction, was used to detect ribosomal RNA genes in extracts.

Using this method, ribosomal RNA genes were recovered from rock extracts. The nucleotide sequencing of these ribosomal RNA genes is presently in progress. The results of that analysis will reveal the evolutionary relationships of the cloned genes. It is likely that the identification of these gene clones will influence our perceptions of the origins of these genes. If these genes represent novel microorganisms, or show phylogenetic affinities with known metal-oxidizing groups of bacteria associated with geochemical processes, then the results will be interpreted as strong support for the rock origin of the DNA. If, on the other hand, the cloned genes reveal affinities to organisms commonly associated with humans or seawater, then contamination will seem a likely source.

Finally, electron microprobe analysis of areas of enhanced fluorescence showed that phosphorus and potassium (two elements consistent with the presence of life) were present in high abundance (analyses of C and N by microprobe were not attempted). High potassium may be caused by the presence of chemical alteration of the glass in the presence of seawater, but phosphorus is not commonly found in secondary minerals. The affected glass was depleted in iron, calcium, magnesium, and manganese relative to the fresh glass. Iron and manganese in fresh basalt glass is mostly in the reduced form (Christie et al., 1986), so the low Fe and Mn in the altered glass may result from the oxidation and release of iron from the glass by microbes. The low Ca is similar to that observed in the chemical alteration of basalt glass, so the combined effect of chemical and biological processing of basalt glass is probably responsible for the chemistry of the altered areas analyzed by microprobe.

The results reported herein provide significant support for the hypothesis that bacteria are somehow involved in the alteration of basaltic glasses to clays.

**Note added in proof:** Nucleotide sequences indicate that the cloned ribosomal RNA genes are bacterial. At least six different classes of genes have been identified in 61 clones originating from the two rock samples. One of these was identified as a gene from the bacterial genus Thermus, and is almost certainly a contaminant that...
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originated in reagents used for PCR. The other genes, which have not been completely studied, belong to a variety of bacterial groups, but are predominately genes from proteobacteria. The largest class of genes in the library has a mole% guanine + cytosine content close to 50%, which is characteristic of organisms adapted to mesophilic environments. These sequence data suggest that only a traces of DNA were extracted from these rocks, and do not provide proof of the phylogenetic affiliations of the basalt glass microbial community. However, carbon was detected by electron microprobe at the tips of channels, but not elsewhere in the rocks, confirming that the regions of most intense nucleic acid staining have an elemental composition consistent with organic material.

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REFERENCES


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Plate 1. 1. Thin section from Sample 148-896A-7R-1, 52-55 cm (Piece 10) (284 mbsf) viewed under a combination of plain polarized light and epillumination for Hoechst dye. The long dimension of the photograph is 0.6 mm. The top of the photograph is clear glass with brown microlites. The bottom of the photograph is clay surrounding a crack. A dark brown irregular interface separates the clay and the glass. Hoechst 3342 becomes fluorescent (blue, emission maximum 460 nm) upon binding to adenosine/thymidine-rich regions in the minor-groove of DNA. 2. Thin section from Sample 70-5048B-48R-2 viewed by epifluorescence microscopy. The section was stained with PO-PRO-3, a cyanin dye with high affinity for DNA. 3. Thin section from Sample 70-5048B-48R, 120-124 cm, viewed by transmitted light. 4. The thin section shown in Figure 3 stained with the green fluorescent dye Syto11, which binds to DNA and RNA. The long axis of each image above is 250 µm.