Diversity and distribution of subseafloor Thermococcales populations in diffuse hydrothermal vents at an active deep-sea volcano in the northeast Pacific Ocean

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[1] The presence, diversity, and distribution of a key group of subseafloor archaea, the Thermococcales, was examined in multiple diffuse flow hydrothermal vents at Axial Seamount, an active deep-sea volcano located in the northeast Pacific Ocean. A polymerase chain reaction (PCR) approach was used to determine if this group of subseafloor indicator organisms showed any phylogenetic distribution that may indicate distinct subseafloor communities at vents with different physical and chemical characteristics. Targeted primers for the Thermococcales 16S rRNA (small subunit ribosomal RNA) gene and intergenic transcribed spacer (ITS) region were designed and applied to organisms filtered directly from a variety of diffuse flow vents. Thermococcales were amplified from 9 of 11 samples examined, and it was determined that the ITS region is a better phylogenetic marker than the 16S rRNA in defining consistent groups of closely related sequences. Results show a relationship between environmental clone distribution and source vent chemistry. The most highly diluted vents with elevated iron and alkalinity contained a distinct group of Thermococcales as defined by the ITS region, suggesting separate subseafloor Thermococcales populations at diffuse vents within the Axial caldera.


1. Introduction

[2] The active circulation of hydrothermal fluids and seawater within the upper 500 m of porous oceanic crust provides a rich environment for microbial growth in the subseafloor at mid-ocean ridges. However, because of the difficulties in sampling the subseafloor environment directly, the microbial and chemical properties of venting fluids can serve as indicators of the processes and habitats beneath the seafloor. Since diffuse flow vents were first discovered, the potential relationship between their chemistry and microbial populations was recognized [Edmond et al., 1979b; Jannasch and Mottl, 1985; Lilley et al., 1984; Von Damm and Lilley, 2004]. Possible microbiological signatures in vent fluids include the consumption of oxygen, sulfide, nitrate, sulfate, hydrogen, and oxygen, as well as the production of ammonia and methane [Butterfield et al., 2004; Edmond et al., 1979a; Jannasch and Mottl, 1985; Lilley et al., 1984; Von Damm and Lilley, 2004]. In addition, thermophilic and hyperthermophilic methanogens and heterotrophs cultured from low-temperature diffuse fluids are viewed as indicator organisms of hot subseafloor habitats because they usually have a minimum growth temperature that is higher than the temperature of the fluids sampled [Holden et al., 1998; Holland et al., 2004; Summit and Baross, 1998, 2001].

[3] Owing to their ability to grow well in artificial media, the Thermococcales (Pyrococcus, Thermococcus, and Palaeococcus spp.) is one of the most thoroughly studied groups of cultured organisms from high-temperature environments, including marine hydrothermal vents, solfatarea, oil reservoirs, and freshwater thermal pools [Zillig and Reysenbach, 2001]. Most have a fermentative metabolism, require multiple amino acids for growth, and are anaerobic

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thermophiles or hyperthermophiles [Zillig and Reysenbach, 2001]. However, the Thermococcales also exhibit tremendous metabolic versatility, such as the ability to grow lithotrophically on CO coupled with H₂ production [Sokolova et al., 2004], form biofilms [Reysenbach et al., 2000; Rink and Kelly, 2000], and grow over wide temperature and organic carbon concentration ranges [Holden et al., 2001; Summit and Baross, 2001]. Syntrophic relationships with subseafloor autotrophic primary producers, such as methanogens and the Aquificales, may also help support Thermococcales populations [Bonch-Osmolovskaya and Stetter, 1991; Rink and Kelly, 2000]. Additionally, members of the Thermococcales have a high tolerance to oxygen exposure and show long-term survival at cold temperatures, characteristics thought to be important for their dispersal and wide distribution, as well as their dominance in culture collections from deep-sea hydrothermal vents [Jannasch et al., 1992; Stetter, 1989].

[4] Using cultured isolates, the distribution, and genetic and phenotypic diversity of Thermococcales populations have been studied from different biotopes at deep-sea hydrothermal vents [Holden et al., 2001; Lepage et al., 2004; Marteinsson et al., 1995; Summit and Baross, 2001; Wery et al., 2002]. Furthermore, using PCR-based assessments with the 16S rRNA gene, a highly conserved gene found in all living prokaryotes, Thermococcus spp. have been detected on enrichment surfaces in deep-sea vent fluids (in situ growth chambers) [Nakagawa et al., 2005; Nercessian et al., 2003; Reysenbach et al., 2000; Takai et al., 2004a] and black smoker chimneys and plumes from Manus Basin, the Juan de Fuca Ridge, Central Indian Ridge, and the Okinawa Trough [Hoek et al., 2003; Nakagawa et al., 2005; Schrenk et al., 2003; Takai et al., 2001; Takai et al., 2004b]. However, the few microbial diversity studies of deep-sea hydrothermal vent fluids without any enrichment on surfaces have not detected the Thermococcales [Huber et al., 2002; Takai and Horikoshi, 1999]. Another genetic marker, the 16S–23S rDNA intergenic transcribed spacer (ITS) region, has also proved useful in examining microbial diversity and distribution in marine environments. Owing to its high sequence and length variability, the ITS region is often used in phylogenetic studies to identify closely related species and strains [Garcia-Martinez et al., 1999; Gurtler and Stanisch, 1996]. Currently, the ITS region is being used to examine ecotypes and environmental niche adaptation in marine prokaryotic population studies [Garcia-Martinez et al., 2002; Garcia-Martinez and Rodriguez-Valera, 2000; Holden et al., 2001; Rocap et al., 2002; Summit and Baross, 2001].

[5] The purpose of this study was to apply PCR-based environmental sequence analysis to one of the key group of vent hyperthermophiles, the Thermococcales, to assess their presence, diversity, and distribution in low-temperature hydrothermal vent fluids at an active deep-sea volcano, Axial Seamount. We used both the 16S rRNA gene and ITS region to determine if the Thermococcales showed any phylogenetic distribution that may indicate distinct subseafloor communities at vents with different chemical characteristics. The results show that Thermococcales are widespread in diffuse vents at Axial Seamount, and that environmental sequences could be clustered to specific vents, suggesting that distinct subseafloor communities may exist at individual diffuse vents.

2. Study Site and Methods

[6] Located 480 km off the coast of Oregon at a depth of 1520 m on the Juan de Fuca Ridge, Axial Seamount (45°58’ N; 130°00’ W) is an active submarine volcano (Figure 1). The caldera of the volcano (3 × 8 km) lies between two rift zones to the north and south and is bordered on three sides by a boundary fault, and active long-term venting is located along these rift zones and near the boundary fault [Johnson and Embley, 1990]. Three distinct hydrothermal fields exist at Axial in the caldera; at the CASM site to the north [Canadian American Seamount Expedition, 1985], at the ASHES site to the southwest [Embley et al., 1990], and within the 1998 lava flow along the southeast portion of the caldera [Baker et al., 1999] (Figure 1). Other vents exist in the southern portion of the caldera, including a region just west of the 1998 lava flow, termed pre-1987 lava flow.

[7] The systematic sampling and analyses of a variety of diffuse flow vents at Axial since the eruption show that these vents are chemically and geologically diverse and support a microbial subseafloor community, including hyperthermophilic methanogens and heterotrophs and potential nitrogen fixers [Butterfield et al., 2004; Huber et al., 2002, 2003; Mehta et al., 2003]. Because of the nearshore location, the presence of well-characterized hydrothermal systems, and the abundance of diffuse venting, Axial Seamount is a readily accessible and excellent site for studying the subseafloor in young oceanic crust and examining linkages between the geochemical environment and subseafloor microbial communities.

2.1. Sample Collection

[8] Using the ROV ROPOS, basalt hosted diffuse fluids were collected in 1998, 1999, 2000, and 2001 from vents throughout the caldera, as indicated in Tables 1 and 2 and Figure 1. Fluids were sampled using the Hydrothermal Fluid and Particle Sampler (HFPS), which pumps vent fluids through a titanium intake nozzle and measures the temperature of the fluid throughout the sampling period. This sampler allows for the collection of both whole fluids for chemical analyses, cell counts, and culturing, as well as in situ filtered fluids for DNA-based and chemical analyses. Filtered and unfiltered fluids were sampled at a vent after a stable temperature was reached on the intake probe, and fluids were pumped at a known rate until the desired volume was collected. Temperature and volume of fluid collected was monitored throughout the 10–15 minute sampling time required to obtain approximately one liter of fluid. For filtered samples, fluid was pumped through either a 47 mm Millipore (3 μm pore size) cellulose ester filter, followed by a Sterivex-GP (0.22 μm pore size) filter, or only the >0.22 μm fraction was isolated with a Sterivex-GP filter unit. Different size filter units were used owing to both space constraints on the sampler and multiple experimental needs for each sample. A background (no detectable hydrothermal plume) seawater sample from 2540 m depth (47°42’N, 127°38’W) was collected using a 10 liter Niskin Bottle mounted on a CTD (conductivity, temperature, depth) and filtered through a sterile 47 mm 0.22 μm pore...
Figure 1. Map of Axial Seamount showing (a) caldera and three main vents fields and (b) individual vents from the southeast portion of the caldera. Figures were modified from Bill Chadwick.
### 2.4. Primer Design and Testing

[11] A forward PCR primer for the 16S rRNA gene was designed to amplify selected members of the order Thermococcales, including *Pyrococcus*, *Thermococcus*, and *Palaeococcus* spp. A segment of the gene with near consensus was targeted on the basis of an oligonucleotide probe designed for fluorescence in situ hybridization analyses [Schrenk et al., 2003]. Approximately 20 16S rRNA sequences for the Thermococcales were compiled from the GenBank database via the National Center for Biotechnology Information (NCBI) website and aligned with 60 lab isolates of the Thermococcales from a variety of hydrothermal vent environments in the BioEdit sequence alignment editor [Hall, 1999]. The previous probe, Tc589F, was modified with the insertion of a Y (serving as C or T), which expanded its matches to include *Palaeococcus* spp. The modified primer was submitted to the Probe Match function of the Ribosomal Database Project II [Cole et al., 2003] to check for self-complementarities and matches in the database. The new primer only matched with members of the Thermococcales and was designated Tc589FII. The sequence for the primer is 5′-GCC YGT AAG TCC CTG

### Table 2. Source of DNA Samples and Summary of Results From Microscopic, Culturing, and Molecular Analyses

<table>
<thead>
<tr>
<th>Sample</th>
<th>Vent</th>
<th>Size Fraction</th>
<th>Volume (L)</th>
<th>Fluid cells/mL</th>
<th>±95% Conf. cells/mL</th>
<th>Cultures(^a)</th>
<th>MPN results(^c)</th>
<th>Percent(^d)</th>
<th>PCR(^e)</th>
<th>Number of Clones Sequenced</th>
</tr>
</thead>
<tbody>
<tr>
<td>FS011</td>
<td>Marker 33</td>
<td>&gt;3.0 1998 Lava Flow</td>
<td>1.5</td>
<td>2.24E+05</td>
<td>1.61E+04</td>
<td>≥48,000</td>
<td>&gt;0.021</td>
<td>+</td>
<td>72</td>
<td></td>
</tr>
<tr>
<td>FS016</td>
<td>Marker 33</td>
<td>&gt;3.0 1998 Lava Flow</td>
<td>1</td>
<td>1.12E+05</td>
<td>3.82E+03</td>
<td>3000–96,000</td>
<td>0.086</td>
<td>+</td>
<td>64</td>
<td></td>
</tr>
<tr>
<td>FS047</td>
<td>Marker 33</td>
<td>&gt;3.0 1998 Lava Flow</td>
<td>1</td>
<td>1.23E+05</td>
<td>1.59E+04</td>
<td>&gt;48,000</td>
<td>0.004</td>
<td>+</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>FS009</td>
<td>Easy</td>
<td>&gt;3.0 1998 Lava Flow</td>
<td>1</td>
<td>1.19E+05</td>
<td>2.56E+04</td>
<td>no data</td>
<td>no data</td>
<td>+</td>
<td>72</td>
<td></td>
</tr>
<tr>
<td>FS024</td>
<td>Magnesia</td>
<td>&gt;3.0 1998 Lava Flow</td>
<td>1</td>
<td>5.02E+05</td>
<td>3.04E+04</td>
<td>80–2400</td>
<td>≤0.0005</td>
<td>+</td>
<td>63</td>
<td></td>
</tr>
<tr>
<td>FS062</td>
<td>Old Worms</td>
<td>0.2 E of 1998 Lava Flow</td>
<td>1</td>
<td>2.38E+05</td>
<td>2.03E+04</td>
<td>no data</td>
<td>no data</td>
<td>+</td>
<td>55</td>
<td></td>
</tr>
<tr>
<td>FS068</td>
<td>Marshmallow R624 2001</td>
<td>&gt;0.2 ASHES</td>
<td>3</td>
<td>3.00E+04</td>
<td>8.02E+03</td>
<td>80–2400</td>
<td>≤0.0008</td>
<td>+</td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>FS075</td>
<td>Fe-City</td>
<td>&gt;0.2 Pre-1987 Lava Flow</td>
<td>1</td>
<td>1.59E+05</td>
<td>1.83E+04</td>
<td>60–880</td>
<td>≤0.0006</td>
<td>+</td>
<td>55</td>
<td></td>
</tr>
<tr>
<td>FS076</td>
<td>Bag City</td>
<td>&gt;0.2 Pre-1987 Lava Flow</td>
<td>1</td>
<td>3.26E+05</td>
<td>2.45E+04</td>
<td>no data</td>
<td>no data</td>
<td>+</td>
<td>51</td>
<td></td>
</tr>
<tr>
<td>FS045</td>
<td>Cloud</td>
<td>&gt;0.2 Pre-1987 Lava Flow</td>
<td>1</td>
<td>1.19E+05</td>
<td>1.74E+04</td>
<td>720–26,000</td>
<td>≤0.0002</td>
<td>-</td>
<td>no data</td>
<td></td>
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<tr>
<td>FS067</td>
<td>Gollum</td>
<td>&gt;0.2 ASHES</td>
<td>3</td>
<td>2.01E+05</td>
<td>2.20E+04</td>
<td>60–880</td>
<td>≤0.0004</td>
<td>-</td>
<td>no data</td>
<td></td>
</tr>
<tr>
<td>CTD005</td>
<td>Seawater</td>
<td>&gt;0.2 47°24′N, 127°38′W</td>
<td>3</td>
<td>3.97E+04</td>
<td>5.97E+03</td>
<td>ND</td>
<td>ND</td>
<td>-</td>
<td>no data</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)Volume of fluid filtered.

\(^b\)Positive enrichment of thermophiles or hyperthermophiles from indicated vent.

\(^c\)Percent MPN cultured microbes represent of total microbial population.

\(^d\)Positive PCR product with Tc589FII and 64Ra primer set.
GCG-3’. A similar but longer primer designed to be used in the reverse direction was recently published for identifying hyperthermophilic archaea of the family Thermococcales [Slobodkina et al., 2004].

[12] Primer Tc589FII was paired with primer 64Ra (5’-GCG NRG GCT TAT CGC AGC TT-3’) targeting the 23S rRNA [Summit and Baross, 2001]. This combination allows for amplification of approximately 1000 bases from the 16S rRNA gene and the complete ITS region. The primer set was then tested on 30 different members of the Thermococcales, including 25 isolates from Axial Seamount and other Juan de Fuca Ridge hydrothermal environments, as well as T. littoralis, T. peptonophilus, T. celer, and T. barophilus. Successful amplification revealed a band of approximately 1250 base pairs. The primers were also applied to DNA from bacterial species, including Escherichia coli, Halomonas spp., Thermosiphon spp., and Desulfuorobacterium spp., as well as two Methanococcus species. No amplification product was observed. A variety of PCR conditions were tested with different annealing temperatures and magnesium chloride concentrations to find the optimal set of PCR conditions for amplification of isolate DNA. Each PCR reaction (25 µl) contained 5% acetamide, 1.5 mM MgCl₂, 0.8 mM deoxyribonucleoside triphosphates, 50 ng (each) primer, 1X PCR Buffer (Promega), and 1 U of Taq DNA Polymerase (Promega). The profile was an initial denaturation step of 2 min at 94 °C, followed by 30–36 cycles of 94°C for 30 s, 59°C for 45 s, and 72°C for 2 min. The final extension step was 72°C for 10 min.

2.5. Environmental PCR and Clone Library Construction

[13] The primers were applied to the environmental DNA samples listed in Tables 1 and 2. If amplification was observed, replicate positive amplification products were reconditioned to minimize heteroduplexes [Thompson et al., 2002]. To recondition, 1 µl of positive PCR product was placed into a fresh cocktail as template and put through 3 more cycles of amplification, including both the hot start and final extension. The separately reconditioned PCR products were then consolidated and cleaned using the Qiaquick Gel extraction columns (Qiagen) according to the manufacturer’s instructions, and visualized on 1% (w/v) agarose gels stained with SYBR green (Molecular Probes). PCR bands of the correct size were gel extracted using the Qiaquick Gel extraction columns (Qiagen) according to the manufacturer’s instructions. Final PCR products were cloned according to Huber et al. [2003]. In some cases, if no product was observed with the Tc589FII and 64Ra primer set, a nested PCR approach was applied. PCR was performed on DNA with the universal archaea specific primer 21F (5’-TTC CGG TTG ATC CGY CGG GA-3’) [DeLong, 1992] targeting the 16S rRNA and the 23S rRNA primer 64Ra. Each PCR reaction (25 µl) contained 5% acetamide, 1.5 mM MgCl₂, 0.8 mM deoxyribonucleoside triphosphates, 50 ng (each) primer, 1X PCR Buffer (Promega), and 1 U of Taq DNA Polymerase (Promega). An initial denaturation step of 2 min at 94 °C was followed by 15, 20, 25, and 30 cycles of 94°C for 30 s, 62°C for 45 s, and 72°C for 3 min. The final extension step was 72°C for 10 min. The bands were visualized and quantified on 1% (w/v) agarose gels stained with SYBR green (Molecular Probes). The lowest possible cycle number for which a band was visible was then chosen as the template (1 µL) for the second round of PCR with Tc589FII and 64Ra using the profile as previously described. Again, the lowest number of cycles (15–25 cycles) for successful amplification was used for cloning as visualized on 1% (w/v) agarose gels. PCR products were reconditioned, pooled, gel extracted, and subsequently cloned according to Huber et al. [2003].

2.6. Sequencing

[14] Clones were randomly chosen for complete sequencing of the insert with PCR primers and 958R (5’-YCC GCC GTT GAM TCC AAT T-3’) [DeLong, 1992], 956Fa (5’-TTA ATT GGA KTC AAC GGC GG-3’), 1492Ra (5’-GGY TAC CTT GTT AGC ACT T-3’), and 1492Fa (5’-AAG TCG TAA CAA GGT ARC CGT-3’) [Summit and Baross, 2001]. The DYEnamic ET Dye Terminator Kit (Amersham Pharmacia Biotech Inc.) with subsequent analysis on a MegaBACE 1000 (Molecular Dynamics) was used for sequencing.

2.7. Phylogenetic and Cluster Analysis

[15] Sequencer was used to assemble sequences, and the contiguous sequences were manually aligned and manipulated in the BioEdit v4.7.8 Program Sequences [Hall, 1999]. Sequences that stood out in the alignment as possible chimeras or members of a different order were noted. Each clone was then divided into the 16S rRNA and ITS region, and treated separately for the remainder of the analyses. To find closely related sequences for phylogenetic analysis, all sequences were submitted to the Advanced BLAST search program (available through the National Center for Biotechnology Information), and closest matches added to the alignment. All 16S rRNA gene sequences were submitted to the Bellerophon server for chimera detection [Hugenholtz and Huber, 2003]. Maximum-likelihood phylogenetic trees were constructed using the Hasegawa-Kishino-Yano evolutionary model [Hasegawa et al., 1985] with 50,000 puzzling steps and with mutation rates estimated from the data sets using TREE-PUZZLE version 5.0 [Schmidt et al., 2002]. Branches were collapsed for quartet-puzzling support values of <50% and graphed using TreeView version 1.6 [Page, 1996].

[16] The ITS region and 16S rRNA gene were treated separately in phylogenetic analyses owing to their different levels of sequence variability. For the ITS, sequence identity matrices were made for each sample, and clones with exactly 100% sequence match were identified. A representative clone for each 100% identity group was then chosen and individual trees for each vent constructed. Repetitive groups with the same culture outgroups were identified from all 9 sample trees, and a final consensus tree was constructed to depict the major groups found, with a representative clone from each sample and each group included (Figure 2). For the 16S rRNA gene, the same 100% identity procedure was carried out; however, because all clones were 94–100% identical, it was determined that the 100% sequence identity cutoff did not allow for enough resolution in the individual sample trees for comparison between samples. Therefore a ≥99% similarity index was applied to group together clones within each sample, and a final tree constructed with a representative of each ≥99%
similarity group from each sample included [Acinas et al., 2004a]. From this tree, groups were identified and their distribution among samples noted (Figure 3).

A cluster analysis was performed on the microbial groups (A–F for ITS, Figure 2, and C–I for 16S, Figure 3) found at each vent using Jaccard’s coefficient. A matrix was constructed where designated groups were scored for presence or absence in each vent. Jaccard’s coefficient was calculated, and a cluster diagram of the samples was constructed using the unweighted pair group method using
3. Results

Vent fluids were collected from 11 different hydrothermal vents located throughout the Axial caldera (Figure 1). Some chemical and microbiological characteristics of the vent sites are given in Tables 1 and 2. The samples analyzed for this study provide a cross section of the type of diffuse vents seen at Axial. These included: Marshmallow vent (FS068), an anhydrite-dominated high-temperature vent in the ASHES field with a hot fluid source just below the seafloor; Gollum vent (FS067), a stable low-temperature vent in the ASHES field; Bag City, Old Worms, and Fe-City (FS076, FS062, and FS075), stable diffuse vents outside of Figure 3.

Figure 3. Maximum-likelihood quartet puzzling phylogenetic tree of cloned 16S rRNA genes for the Thermococcales. Tree shows one representative from each sample for ≥99% sequence identity groups. Clones from this study are denoted in bold type face and refer to specific samples and clone number, with the number of clones they represent from that library in brackets. Accession numbers for all cultures and clones are indicated in parentheses. The complete amplified 16S rRNA (931–943 bp) was used in the alignment. Quartet puzzling support values are shown at each internal branch. The scale bar represents the expected number of changes per nucleotide position.
the 1998 eruption zone; Marker 33 and Cloud (FS045), stable vents within the boundary of the 1998 lava flow; and Magnesia (FS024) and Easy vent (FS009), short-lived low-temperature "snow-blower" vents that died out within two years of the 1998 eruption. Details of the geological setting, chemical composition, and chemical indications of microbial activity in and near the eruption zone have been presented previously [Butterfield et al., 2004]. Prior results indicate the following microbial processes in the subseafloor: oxidation of sulfide and hydrogen; reduction of nitrate, sulfur, and iron; and methanogenesis [Butterfield et al., 2004]. In addition, in some diffuse vent fluids more methane is present than can be produced through methanogenesis by reaction of primary hydrogen (<1 mmol/kg) in representative high-temperature source fluids with the abundant carbon dioxide (>100 mmol/kg) in those same fluids, suggesting fermentation of organic matter, the primary metabolism of the Thermococcales (data not shown) [Butterfield et al., 2004]. The three lowest-temperature vents examined in this study, Easy, Magnesia, and Fe-City, show distinct chemical characteristics. These vents have the highest average Mg content (Table 1 and Figure 5a), indicating that they contain the least amount of high-temperature hydrothermal end-member. This is supported by all of the indicators of mixing. Easy, Magnesia and Fe-City have the highest seawater components (Mg, SO₄, U) and the lowest hydrothermal components (Si, Li, Cs) (Table 1). The higher degree of subseafloor mixing with seawater at these vents is one factor that can help explain their differences compared to other vents, but in addition, there are significant chemical differences that cannot be explained simply by the degree of mixing. Although Easy, Magnesia, and Fe-City vents are the coolest and most highly diluted, they have among the highest iron concentrations of any of the diffuse vents (Table 1) and by far the highest iron content when normalized to heat content (Figure 5). In addition, these three vents are distinguished by alkalinites higher than ambient seawater, while all of the other vent fluids in this study have alkalinites less than seawater (Figure 5b). The high dissolved iron and increased alkalinity is consistent with mobilization of iron at relatively low temperature, probably owing to reduction of the ubiquitous ferric iron precipitates that exist around the margins of diffuse hydrothermal areas on Axial Volcano [Butterfield et al., 2004; Kennedy et al., 2003].

Figure 4. Cluster diagram showing distribution of Thermococcales groups at individual vents based on ITS region.

[19] Positive enrichment cultures were obtained for heterotrophic thermophiles or hyperthermophiles from all hydrothermal fluids sampled at Axial Seamount (Table 2). Most-probable-number estimates of thermophilic and hyperthermophilic heterotrophs indicate that these cultured organisms make up less than 1% of the total DAPI-stained population (Table 2), which is consistent with MPN estimates of Thermococcales from other vent environments [Holden et al., 1998; Holland et al., 2004].

[20] Clone libraries from in situ filtered diffuse fluids were constructed using primers targeting the Thermococcales. As summarized in Table 2, 9 of the 12 environmental samples yielded a positive PCR product of approximately 1250 base pairs. In two vents, Cloud and Gollum, Thermococcales were successfully cultured although no positive PCR product was obtained. This is likely due to the low recovery of DNA from these samples or the low abundance of Thermococcales at these sites. While no PCR product was obtained from Cloud regardless of primer set used, universal archaean primers did result in positive amplification from Gollum even though the Thermococcales-specific primers were unable to produce a product. Out of 526 clones sequenced, 5 were Methanococcus spp., and one was a member of the Marine Group I Crenarchaeota. Those 6 clones were eliminated from further analysis. All of the remaining 520 clones were members of the Thermococcales and appeared to be nonchimeric. All cloned ITS regions (241 bp to 261 bp) contained a tRNA-alanine as observed in other Euryarchaeota [Achenbach-Richter and Woese, 1988].
ITS clone identities ranged from 75% to 100% sequence similarity, whereas 16S rRNA clones (931 bp to 943 bp) ranged from 94% to 100% sequence similarity.

The majority of clone sequences in this study grouped most closely with cultured members of the Thermococcales, including both *Pyrococcus* and *Thermococcus* spp. We included outgroups from vent locations throughout the world in both the 16S rRNA gene and ITS region analyses, and no global biogeographical trend was seen, with clones from this study grouping with cultured organisms from Axial Seamount, sulfide chimneys at the Mid-Atlantic Ridge, shallow marine springs in Vulcano, Italy, and mineral precipitates from the Okinawa Trough. This supports other studies which have shown the Thermococcales to be widespread and ubiquitous in hydrothermal environments [Lepage et al., 2004].

**Figure 5.** Trends of iron, alkalinity, and magnesium in diffuse vents of this study. Delta properties are calculated as measured property minus the ambient seawater background property. Concentration of iron in background seawater is small enough to be insignificant. Use of delta properties eliminates most of the effects of mixing hydrothermal fluid with seawater. (a) Fe/ΔT versus Mg, with three vents highlighted. An increase in iron occurs in the lowest-temperature vents. (b) ΔAlkalinity/ΔT versus Fe/ΔT shows a roughly linear correlation of the ΔAlkalinity and iron in the lowest-temperature vents, consistent with reduction of solid phase ferric iron to soluble Fe$^{2+}$ and consumption of acid. The remaining vents have relatively low iron and negative ΔAlkalinity.
As discussed in the methods section, the 16S and ITS region were treated separately in all analyses owing to their different levels of sequence similarity. Individual phylogenetic trees were constructed for each vent sample for each gene with the same outgroups included. Consistent groups, those containing the same outgroups with strong support values, were noted and their distribution among samples recorded. We found that the ITS and 16S rRNA did not yield identical phylogenetic trees or groupings (Figures 2 and 3). While some groups (C, D, E, and F) contained the same groups on both trees, the tightly clustered groups A and B on the ITS trees were not found on the 16S rRNA trees. Instead, these groups broke into smaller subgroups (G, H, and I). For both the ITS region and the 16S rRNA, there were sequences that could not be assigned to a defined group.

To determine if there was any relationship between the presence and absence of a certain groups for the ITS and 16S rRNA gene in each vent, we performed a cluster analysis (Figure 4). Microbial communities from individual vents were clustered by application of the unweighted pair-group method with arithmetic mean method (UPGMA) to the Jaccard’s coefficient matrix. Despite the inconsistencies between the ITS and 16S rRNA phylogenies, UPGMA cluster analysis with both genes showed three vents consistently clustering together, Magnesia, Fe-City, and Easy (Figure 4).

4. Discussion

Members of the Thermococcales isolated from low-temperature diffuse fluids with optimum growth temperature well above the temperature of sampled vents serve as indicator organisms of a warm anaerobic subseafloor habitat. While they are the most commonly isolated organisms from deep-sea hydrothermal vents due to relative ease of cultivation and selection methods employed, phylogenetic surveys of hydrothermal fluids often miss the Thermococcales, possibly owing to their low abundance in the subseafloor, their attachment to particles in the subseafloor, or the large amounts of seawater that may dilute their numbers at the point of sampling [Huber et al., 2002]. The targeted environmental PCR analysis with order-specific primers used in this study attempted to minimize the problem of seawater organisms and allowed for the direct examination of the incidence, diversity, and distribution of one member of the warm anoxic subseafloor population in a variety of diffuse flow vents.

It has been suggested that distinct microbial populations may exist at individual vents owing to differences in chemical properties of the vent [Lilley et al., 1984], and that for microbial populations, each vent, even within a vent field like Axial, is potentially an independent habitat [Karl, 1995]. Vents only meters apart can have markedly different chemistries [Butterfield et al., 2004; Nakagawa et al., 2005; Von Damm, 2000; Von Damm and Lilley, 2004], reflecting different fluid sources, pathways, mixing ratios, segregation of fluids, precipitation/dissolution reactions and microbial processes in the subsurface. For example, while diffuse flow is often viewed as a mixture of two end-members, seawater and hydrothermal fluid, there is now evidence for a third “intermediate” fluid type consisting of warm but unreacted seawater [Butterfield et al., 2004; Cooper et al., 2000; Ravizza et al., 2001]. Additionally, the geological context of the vent can also control how fluids move through and out of the crust, such as through linear cracks in the seafloor (Marker 33), through the interior of permeable lava pillars (Marker 113), out of drain-back caverns through partially broken lobate lava roof skylights (Roof), or through what may be a lava tube opening (Cloud) [Butterfield et al., 2004]. These factors, and many others discussed in references included here, influence the sampled chemistry and microbial diversity of a single diffuse flow hydrothermal vent.

For the environmental clone libraries, with both the 16S rRNA gene and in particular with the ITS region, we found sequences grouping together in robust groups of closely related sequences. The significance of this microdiversity remains unknown. Some of the variation may be due to errors in PCR, cloning, and sequencing [Acinas et al., 2004a; Wintzingerode et al., 1997], though this is likely not as significant for the ITS region, which exhibits considerable sequence variation. In addition, some of the microdiversity may be due to multiple copy rRNA gene sequences present in one organism, although most archaea appear to only have a single RNA (rrn) operon [Acinas et al., 2004b]; Palys et al. [1997] have suggested that these clusters of similar sequences should correspond to ecotypes [Cohan, 2001]. Recent studies seem to support this interpretation and show that some clusters appear to represent closely related yet ecologically distinct groups of organisms [Acinas et al., 2004a; Garcia-Martinez and Rodriguez-Valera, 2000; Moore et al., 1998; Rocap et al., 2002]. In particular, the use of the ITS region as a high-resolution marker has revealed that closely related organisms (>97% 16S rRNA sequence similarity) can represent separate physiological populations [Holden et al., 2001; Jaspers and Overmann, 2004; Rocap et al., 2002, 2003; Summit and Baross, 2001]. Although our cluster analyses with both the 16S rRNA and ITS region showed similar results, we found tighter, more consistent groups using the ITS region, supporting other studies which suggest it is a better genetic marker for indicating habitat diversity [Ferris et al., 2003; Rocap et al., 2002].

We looked for relationships to explain the clustering pattern in Figure 4 by examining the characteristics of the sample sites, as listed in Tables 1 and 2. Our analysis with the ITS region shows that three vents, Magnesia (FS024), Easy (FS009), and Fe-City (FS024) cluster together and have a different Thermococcales composition than the other vents. As discussed, these three vents have some common characteristics in their chemistry that are distinct from all other diffuse vents in this study. They are among the most highly diluted (lowest temperature, highest Mg) and show signs of iron reduction (high Fe concentrations, high alkalinity). The ITS region group A was not detected in these three vents but was found in the other five vents. Instead, these three vents contained ITS region groups D, E, and F, which were not found in the other five vents. ITS region group B appeared in all but 3 vents, while ITS region group C only appeared in Marker 33 1998 and 1999. These two vents contained the highest number of observed unique Thermococcales clones of all vents sampled and are identical in terms of Thermococcales groups identified (data not
shown). What distinguishes the Magnesia, Easy, and Fe-City chemically is the indication that they have increased iron content due to reduction of ferric iron in the subsea-floor. It is possible that the genetic distinctness of Thermococcus spp. in these three vents results from entrainment of organisms present in the iron-rich deposits, and the presence of ITS region groups D, E, and F uniquely in the three high-iron vents can be interpreted as the addition of a subpopulation present in these deposits at Axial Volcano. The common presence of most other ITS region groups in all of the vent sites may represent subpopulations entrained from the basalt-hosted upflow zone.

[28] A relationship between environmental characteristics of the habitat and genetic diversity of the microbial population has been observed in terrestrial hot springs [Blank et al., 2002; Ferris et al., 2003; Skirnisdottir et al., 2000], and geographical trends in distribution of hot spring microbes have also been noted, suggesting that physical isolation may be important for microbial diversification [Fenchel, 2003; Papke et al., 2003; Whitaker et al., 2003]. It is clear from our analyses that Magnesia, Easy, and Fe-City vents are distinct from other vents at Axial chemically as well as genetically on the basis of both the 16S RNA and ITS region of environmentally cloned Thermococcus spp. Other studies at deep-sea hydrothermal vents support the hypothesis that microbial populations are linked to the chemical environment. For example, using 16S rRNA and culturing, one previous study found variability in the composition of deep-sea vent microbial communities from multiple chimneys and fluids in a single hydrothermal field in the Mid-Okinawa Trough, and this variability was linked to the geochemistry of the sampled vent fluids [Nakagawa et al., 2005]. Another study using cultured Thermococcus spp. found the cultures formed groups that correlated with their sample location and source material (fluid, rock, or animal), indicating that they were adapted to the specific environment from which they were isolated [Holden et al., 2001]. Additionally, Summit and Barross [2001] found that Thermococcus spp. in sulfide chimney and basaltic subseafloor habitats from one segment of the Juan de Fuca Ridge formed phylogenetically distinct groups and displayed different phenotypic characteristics. Isolates from sulfides produced more active proteases, tolerated higher zinc ion concentrations, and lived at wider temperature and salinity ranges than those isolated from the subsea-floor. Because there is no evidence of restricted exchange between the two habitats, the authors suggest that this phylogenetic grouping reflects significant differences between sulfide and subseafloor habitats that result in the selection and maintenance of separate groups of Thermococcales [Summit and Baross, 2001].

[29] Our work indicates that there may be a link between the genetic diversity of the Thermococcales and source vent chemistry, suggesting distinct subseafloor Thermococcales populations at individual vents within Axial Seamount. Significantly, vents that have distinct subpopulations in the Thermococcales also have distinct physical and chemical properties, with differences in temperature, degree of mixing, and presence or absence of evidence for iron reduction. While it appears that different vents may select for genetically distinct subpopulations of Thermococcales, the details of this selection process and how it is manifested in physiological diversity are unknown. The key factor in the chemistry of the vents that have the distinct subpopulations of Thermococcales is the evidence that the fluids have passed through an iron-rich habitat. It is possible that there are particular Thermococcales living in this habitat that are not common to the typical mineralogy of basalt and altered basalt that characterizes most of the hydrothermal upflow zone. Thermococcales at Axial, Fe-City, and Magnesia may be adapted to more alkaline conditions, lower temperatures, and higher Fe concentrations as compared to Thermococcales from other diffuse vents. Preliminary growth experiments with Thermococcales isolates from Axial indicate wide growth temperature ranges without any correlation to vent temperature. Future experiments will focus on gathering additional physiological data from our cultures to determine the link between relevant chemical parameters and physiological traits.

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