Methyl-compound use and slow growth characterize microbial life in 2-km-deep subseafloor coal and shale beds

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The past decade of scientific ocean drilling has revealed seemingly ubiquitous, slow-growing microbial life within a range of deep biosphere habitats. Integrated Ocean Drilling Program Expedition 337 expanded these studies by successfully coring Miocene-aged coal beds 2 km below the seafloor hypothesized to be “hot spots” for microbial life. To characterize the activity of coal-associated microorganisms from this site, a series of stable isotope probing (SIP) experiments were conducted using intact pieces of coal and overlying shale incubated at in situ temperatures (45 °C). The 30-month SIP incubations were amended with deuterated water as a passive tracer for growth and different combinations of 13C- or 2H-labeled methanol, methylamine, and ammonium added at low (micromolar) concentrations to investigate methotrophy in the deep subseafloor biosphere. Although the cell densities were low (50–2,000 cells per cubic centimeter), bulk geochemical measurements and single-cell–targeted nanometer-scale secondary ion mass spectrometry demonstrated active metabolism of methylated substrates by the thermally adapted microbial assemblage, with differing substrate utilization profiles between coal and shale incubations. The conversion of labeled methylamine and methanol was predominantly through heterotrophic processes, with only minor stimulation of methanogenesis. These findings were consistent with in situ and incubation 16S rRNA gene surveys. Microbial growth estimates in the incubations ranged from several months to over 100 y, representing some of the slowest direct measurements of environmental microbial biosynthesis rates. Collectively, these data highlight a small, but viable, deep coal bed biosphere characterized by extremely slow-growing heterotrophs that can utilize a diverse range of carbon and nitrogen substrates.

Significance

Microbial cells are widespread in diverse deep subseafloor environments; however, the viability, growth, and ecophysiology of these low-abundance organisms are poorly understood. Using single-cell–targeted stable isotope probing incubations combined with nanometer-scale secondary ion mass spectrometry, we measured the metabolic activity and generation times of thermally adapted microorganisms within Miocene-aged coal and shale bed samples collected from 2 km below the seafloor during Integrated Ocean Drilling Program Expedition 337. Microorganisms from the shale and coal were capable of metabolizing methylated substrates, including methylamine and methanol, when incubated at their in situ temperature of 45 °C, but had exceedingly slow growth, with biomass generation times ranging from less than a year to hundreds of years as measured by the passive tracer deuterated water.

Advances in deep-sea drilling technology and analytical methods for cored samples have led to the discovery of microbial life in a range of deep subseafloor habitats, from Earth’s most oligotrophic sediments (1) to its largest aquifer (2). To probe the limits for microbial life, Integrated Ocean Drilling Program (IODP) Expedition 337 sampled a deeply buried [2 km below seafloor (kmbsf)] coal bed system of terrestrial origin and low thermal alteration at Site C0020. In addition to increased depth, temperature, and organic carbon input from coal beds, Site C0020 lacked the dominant electron acceptors (oxygen and sulfate) that have been the focus of other deep-biosphere–focused IODP expeditions (3, 4). Initial results from Expedition 337 showed cell abundances no longer tracked the global depth trend based on previous cruises (5), suggesting that life-limiting conditions may have been reached at these extreme depths (6). The coal bed, however, contained comparatively high concentrations of microbial cells, 10- to 1,000-fold higher than in the adjacent shale and sandstone lithologies, which is a feature common to lignite-containing terrestrial sediments, where higher organic substrate concentrations sourced from the lignite can sustain higher cell numbers (7, 8). Recovered 16S rRNA gene diversity from the coal beds revealed an assemblage that phylogenetically resembled modern terrestrial environments (e.g., peat or forest soil), which was interpreted as representing indigenous microorganisms within Miocene-aged coal, rather than contaminants or overprinting of commonly observed marine sediment microbes (6).

While the concept of coal beds hosting microbial life for millions of years has existed for almost a century (9), microbial activity and metabolic potential through time remain largely unknown in these systems. Measuring activity in the deep biosphere requires a balance between stimulating life enough to observe a metabolic signal and overstimulation of the system such that experimental conditions no longer approximate in situ conditions (10). Subseafloor microbial metabolic activity, such as
bacterial sulfate reduction, is often interpreted through geochemical profiles or microcosm incubations to determine rates of substrate utilization (11–14). Other approaches include traditional (13) and subseafloor-adapted (15) cultivation-based methods, analysis of actively transcribed genes (16), and biomolecule degradation modeling (17) from recovered core materials. These bulk methods have been valuable in developing a picture of the deep subseafloor biosphere characterized by viable, but extremely slow-growing, microorganisms with a range of metabolisms. There are limitations to these methods in some subseafloor habitats like the Shimokita coal bed, off Japan, where biomass is extremely low (1–1,000 cells per cubic centimeter), and bulk analyses may lack the sensitivity to detect microbial activity and growth. Stable isotope probing (SIP) coupled to single-cell–targeted nanometer-scale secondary ion mass spectrometry (NanoSIMS) analysis is well suited for direct activity measurements in low-biomass environments and has been applied in select deep subseafloor biosphere studies (18). This technique additionally offers a unique perspective on the cell-to-cell variation in growth and metabolic potential of co-occurring microorganisms in nature.

Here we used SIP-NanoSIMS to determine the viability and metabolic capability of microorganisms associated with the Shimokita Peninsula coal bed and surrounding lithologies since burial over 20Mya. We focused specifically on methyl substrate metabolism, which could be hypothetically sourced from coal, biomass, or hydrocarbons, since rates of microbial growth in long-term, high-temperature microcosm experiments. Taking advantage of the ability to measure three different stable isotope tracers in parallel with NanoSIMS, we applied combinations of 13C- and 15N-labeled substrates in tandem with deuterated water (2H2O) as a passive tracer for growth. The use of 2H2O with NanoSIMS is an effective method for measuring single-cell microbial biosynthesis with high sensitivity, capable of detecting extraordinarily slow rates of growth (19) predicted to be characteristic of subseafloor biosphere ecosystems (20). This tracer has been successfully combined with lipid biomarker and NanoSIMS analyses for assessing rates of growth in pure culture and mammalian microbiome studies (19, 21, 22), soils (23), and seabed methane seep environments (24, 25), but it has not yet been applied to study microbial growth in low-biomass, deep subsurface ecosystems. This multiple-isotope SIP-NanoSIMS study of microbial life in deep subseafloor environments recovered evidence for methyl substrate metabolism in deep coal and shale beds and exceedingly slow growth within this low-abundance, high-temperature microbial community.

Materials and Methods

SIP Incubation Preparation. IODP Expedition 337 operations commenced July 26 and continued through September 30, 2012, on the drilling research vessel Chikyu. Utilizing riser drilling, a sedimentary sequence was recovered down to 2,466 mbsf at Site C0020 Hole A (41°10′36″N, 124°12′02″E) in a water depth of 1,180 m off the Shimokita Peninsula, Japan. The drilled sequence transitioned from open marine (youngest; late Pliocene, ~5 Ma) to terrestrial (oldest; late Oligocene, ~30 Ma) with depth. Models for maximum temperature reached by Expedition 337 coring report 63.7 °C (26). Shipboard sedimentological, geochemical, and microbiological data and methods are available through IODP publications (27). Additional coal petrography is available in a report by Gross et al. (28).

A total of 52 incubation amendment conditions were prepared onboard to interrogate a range of potential deep biosphere metabolic strategies, and then incubated back in the laboratory at temperatures approximating in situ measurements (27). In this study, incubations from shale (core BL4, 1,606 mbsf, 37 °C incubation temperature), coal (core 15R3, 1,921 mbsf, 45 °C incubation temperature), and mixed (homogenized mixture from multiple cores 19R1, 19R5, 19R7, 20R3, 20R3, 20R8, 20R3, 25R1, 25R2, and 25R3, 1,950–1,999 mbsf, 45 °C incubation temperature) with methanol, methane, and ammonium substrate additions were analyzed. Age estimates of these samples are early to middle Miocene (6). In situ temperatures ranged from 38–48 °C at these sample depths, with pressures of ~30 MPa (27). Two coal beds were included in these incubations: a shallower coal-only sample deposited under more marine-influenced conditions (~1,921 mbsf, core 15R3) and a deeper coal bed deposited under more limnic conditions included in the mixed lithology sample (~2,000 mbsf, cores 24R3 and 25R1). Cores used for incubations were prepared by removal of outer drill-fluid-contaminated layers by means of a sterile ceramic knife as soon as possible after core recovery and stored at 4 °C until incubation preparation, while maintaining an anaerobic atmosphere during the entire process. For preparation of the SIP incubations, the interior portion of the core was manually crushed into centimeter-sized pieces under sterile, anaerobic conditions and distributed evenly into sterile 50-mL glass vials with butyl rubber stoppers and screw caps (Nichidenrika-Glass Co. Ltd.). Vials were flushed with argon and pressurized to 1 atm argon headspace. Sterile C-, N-, and S-free media (1% PBS, 30 g/L NaCl, 12 g/L MgCl2, and 3 g/L KCl) was prepared anaerobically with 2H2O (20 atom % (at. %) 2H2O). This value (20 at. % 2H2O) was selected as the highest level of enrichment with little to no effect on the activity of microorganisms in pure culture (21, 22). Time point 1, time point 2, and autoclaved treatments were prepared for each substrate condition. Time point 1 incubations lasted for 6 mo, while time point 2 and autoclaved treatments were maintained at the in situ incubation temperature for 2.5 y. Due to low levels of activity ascertained from geochemical measurements, all NanoSIMS analyses were conducted on time point 2 and autoclaved samples. Amendments and incubation conditions for the methyl-substrate substrate utilized in this study are listed in Table 1. Equimolar amounts of substrate (30 μmol C, 1.5 mmol final; 3 μmol N, 0.15 mmol final) were added across incubation conditions at 50 at. % enrichment (Cambridge Isotopes). Hydrogen was added as 5 mL of 100% H2 overpressure to incubations (~0.15 atm H2). A full list of the additional incubation conditions prepared onboard is provided in cruise methods (29). Alkalinity (34.39–9.88 mM) and ammonium (2.80–1.83 mM) concentrations from formation fluid samples collected onboard exceed concentrations of C and N amendments (30). Concentrations of methylamine (0.05 mM) and methanol (1 mM) measured from another study of lignite coal (31) also suggested our substrate additions were environmentally relevant.

After 30 months of incubation (March 2014), all treatments were sampled for geochemical analyses before preparation for NanoSIMS. Three milliliters of headspace gas was removed to a vial filled with 0.1 M NaOH for methane analysis and amended with 1 mL of liquid culture media from Table 1. Incubations were sonicated (Bioruptor UCD-250; COSMO BIO) in an ice bath for 20 cycles of 30 s at 200 W on and 30 s off, and (ii) samples were incubated in hydrofluoric acid after initial sonication, rather than after first-density gradient separation. The cell detection limit was determined by no-sample–added controls run in parallel with samples. The same cell enumeration method was used for incubations and initial shipboard samples.

Cells were stained with SYBR Green I [1:40 dilution of SYBR Green in 10 mM Tris and 1 mM EDTA (pH 7.5)], fixed with 4% paraformaldehyde (PFA), and then subjected to two washes, incubating in 3X PBS for 6 h and then for 2 h after each wash. Samples were centrifuged (3,500 × g), and supernatant was decanted after each wash. PFA-fixed samples were stored in 50% ethanol, 3X PBS. The other half of the sample was preserved in glycerol (70% glycerol, 100 mM Tris, and 10 mM EDTA; Bigelow Single Cell Genomics Center preservation protocol), frozen by a cell alive system (CAS, ABI Corporation, Ltd.), and stored at –80 °C (33).

One milliliter of liquid and ~1 g of sediment chips were subsampled by pipette and sterile cell culture loop, respectively, from the PFA-fixed sample. Cell separation, microscopy, and sorting procedures followed the method of Morono et al. (32), with the following modifications to improve cell recovery: (i) Samples were sonicated (Bioruptor UCD-250; COSMO BIO) in an ice bath for 20 cycles of 30 s at 200 W on and 30 s off, and (ii) samples were incubated in hydrofluoric acid after initial sonication, rather than after first-density gradient separation. The cell detection limit was determined by no-sample–added controls run in parallel with samples. The same cell enumeration method was used for incubations and initial shipboard samples. Cells were stained with SYBR Green I [1:40 dilution of SYBR Green in TE (10 mM Tris and 1 mM EDTA)] and sorted following the flow cytometry protocol of Morono et al. (32). Sorted cells were concentrated directly from the flow cytometer into NanoSIMS vials, and coated with indium tin oxide (ITO) as described by Morono et al. (18) and Inagaki et al. (36). ITO coating on polycarbonate membranes (Isopore GTB02500; Millipore) was prepared by a sputtering deposition technique at Astelltech

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Sample Preparation for DNA and iTag Processing. Cells for gDNA extraction were recovered from unfixed samples stored in glycerol using a modified version of the NanoSIMS protocol by omitting detergent and methanol treatments and only shaking (60 min at 500 rpm using a Shake Master; Bio Medical Science) before centrifugation. Nycodenz was also omitted from the gradient separation solution to prevent PCR inhibition (linear gradients of 10%, 30%, and 67% sodium polytungstate solutions were used instead). After separation, the staining and flow cytometry sorting protocol was identical to NanoSIMS processing, with the exception that the cells were collected into a UV sterilized 0.5-mL PCR tube rather than on the ITO filter. Sorted cells were stored at −20 °C until DNA extraction. A proteinase K digestion mixture consisting of 20 mM Tris–HCl (pH 8.0), 5 mM MgCl₂, 2 mM DTT, 4 U/mL heat-labile double-strand specific DNase (HL-dsDNase; ArcticZymes), and 0.4 mg/mL proteinase K was utilized for cell lysis. Before proteinase K application, potential endogenous DNA contamination in the proteinase K mixture was removed by digestion using HL-dsDNase. The HL-dsDNase extracellular DNA digestion was carried out at 25 °C for 2 h, followed by an incubation step at 60 °C for 1 h. The sorted cells and negative controls (sheath fluid) were then lysed using the pretreated proteinase K. One-third volume of the HL-dsDNase–treated proteinase K solution was added to one volume of sheath fluid, which contained the sorted cells (total volume varied by cell sort time). Proteinase K digestion was performed at 60 °C for 8 h, followed by proteinase K inactivation at 95 °C for 10 min. The resultant DNA extract was then used directly for first-round PCR amplification with universal primers targeting the V4 region of the 16S rRNA gene (U515F: 5′-[Biotin]-GTGCCAGCMGCAGCGGAG-3′; U806R: 5′-YITCGTCACTGGGAGGCAGCAG-3′). 4 μL of the PCR reaction was loaded onto a 1% agarose gel and stained with ethidium bromide. Locations of the PCR amplicons were determined by gel electrophoresis and then indexed and barcoded in the second PCR assay. The iTag sequencing was performed on a MiSeq Illumina platform following.

Table 1. Geochemical and cell enumeration for SIP-NanoSIMS incubations

<table>
<thead>
<tr>
<th>Lithology</th>
<th>13C source, 15N source</th>
<th>DIC, mM</th>
<th>δ13C-DIC, ‰</th>
<th>CH4/Ar, ‰</th>
<th>δ13C-CH4, ‰</th>
<th>δ3H-H2, ‰</th>
<th>Cells, g (rock)</th>
<th>Total cells</th>
<th>Percent active</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shale - 8L4 (1,606 mbsf)</td>
<td>— — — —</td>
<td>0.36 nm</td>
<td>-0.48 nm</td>
<td>0.3 nm</td>
<td>0.3 nm</td>
<td>851</td>
<td>1,548</td>
<td>5E + 05</td>
<td>1E + 05</td>
</tr>
<tr>
<td>— MeAm.</td>
<td>Y 0.08 0.10 0.66</td>
<td>-1.24 0.2 0.3</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
<td>153</td>
<td>5,873</td>
<td>2E + 04</td>
<td>6E + 05</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.30 3.21</td>
<td>4.62 0.1</td>
<td>0.3 nm</td>
<td>0.1 nm</td>
<td>1,203</td>
<td>0.95</td>
<td>11.92</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.51 0.72</td>
<td>8.32 0.2</td>
<td>0.1 nm</td>
<td>0.1 nm</td>
<td>26,171</td>
<td>1,664</td>
<td>3E + 06</td>
<td>2E + 05</td>
<td></td>
</tr>
<tr>
<td>MeOH.</td>
<td>— 0.28</td>
<td>0.40</td>
<td>-2.86</td>
<td>0.1</td>
<td>0.2 nm</td>
<td>26,171</td>
<td>1,664</td>
<td>3E + 06</td>
<td>2E + 05</td>
</tr>
<tr>
<td>MeOH</td>
<td>— 0.74 0.69</td>
<td>128.37 0.3</td>
<td>0.1 nm</td>
<td>0.2 nm</td>
<td>539</td>
<td>7,503</td>
<td>5E + 04</td>
<td>1E + 06</td>
<td>78</td>
</tr>
<tr>
<td>Coal - 15R3 (1,921 mbsf)</td>
<td>— — — —</td>
<td>0.16 nm</td>
<td>-10.33 nm</td>
<td>0.1 nm</td>
<td>0.2 nm</td>
<td>60</td>
<td>60</td>
<td>nm</td>
<td>nm</td>
</tr>
<tr>
<td>— MeAm.</td>
<td>Y 0.06 0.07</td>
<td>-6.00</td>
<td>-3.77</td>
<td>1.1</td>
<td>5.5</td>
<td>42.8</td>
<td>-46.1</td>
<td>-170.4</td>
<td>69</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.04 0.23</td>
<td>16.19</td>
<td>-7.20</td>
<td>0.8</td>
<td>8.6</td>
<td>-57.2</td>
<td>-55.7</td>
<td>-198.2</td>
<td>21*</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.02</td>
<td>0.03</td>
<td>88.77</td>
<td>20.10</td>
<td>0.8</td>
<td>4.5</td>
<td>-42.3</td>
<td>-55.0</td>
<td>-144.1</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.11</td>
<td>0.03</td>
<td>108.06</td>
<td>55.79</td>
<td>1.0</td>
<td>8.1</td>
<td>-58.0</td>
<td>-58.8</td>
<td>-158.2</td>
</tr>
<tr>
<td>MeOH</td>
<td>— 0.06 0.08</td>
<td>19.09</td>
<td>-6.46</td>
<td>2.4</td>
<td>5.4</td>
<td>-49.7</td>
<td>-52.9</td>
<td>-96.5</td>
<td>32*</td>
</tr>
<tr>
<td>MeOH</td>
<td>— 0.07</td>
<td>0.05</td>
<td>-15.14</td>
<td>26.67</td>
<td>1.2</td>
<td>3.8</td>
<td>-57.5</td>
<td>-56.6</td>
<td>bd</td>
</tr>
<tr>
<td>Mixed - 19:25R (1,950–2,000 mbsf)</td>
<td>— — — —</td>
<td>0.16 nm</td>
<td>4.29 nm</td>
<td>0.2 nm</td>
<td>0.2 nm</td>
<td>22,200</td>
<td>22,200</td>
<td>nm</td>
<td>nm</td>
</tr>
<tr>
<td>— MeAm.</td>
<td>Y 1.48 1.38</td>
<td>5.63</td>
<td>3.35</td>
<td>0.2</td>
<td>1.4</td>
<td>11</td>
<td>32*</td>
<td>24*</td>
<td>8E + 02</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.90 1.97</td>
<td>5.72</td>
<td>6.85</td>
<td>0.2</td>
<td>0.1</td>
<td>20</td>
<td>43</td>
<td>7E + 02</td>
<td>1E + 03</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.98 0.97</td>
<td>38.45</td>
<td>37.73</td>
<td>0.1</td>
<td>0.5</td>
<td>0.8</td>
<td>37*</td>
<td>6E + 02</td>
<td>5E + 02</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 1.73</td>
<td>1.79</td>
<td>48.63</td>
<td>43.57</td>
<td>0.1</td>
<td>0.1</td>
<td>0.8</td>
<td>42</td>
<td>130</td>
</tr>
<tr>
<td>MeOH</td>
<td>— 0.15</td>
<td>1.76</td>
<td>6.32</td>
<td>14.87</td>
<td>0.2</td>
<td>2.2</td>
<td>16*</td>
<td>271</td>
<td>5E + 04</td>
</tr>
<tr>
<td>MeOH</td>
<td>— 1.83</td>
<td>1.96</td>
<td>9.42</td>
<td>14.87</td>
<td>0.2</td>
<td>2.2</td>
<td>14*</td>
<td>2,692</td>
<td>5E + 04</td>
</tr>
<tr>
<td>MeAm.</td>
<td>— 0.11</td>
<td>0.03</td>
<td>108.06</td>
<td>55.79</td>
<td>1.0</td>
<td>8.1</td>
<td>0.16</td>
<td>0.16</td>
<td>nm</td>
</tr>
</tbody>
</table>

Geochemical summary of incubations by lithology and substrates added (MeAm., methylamine (tan); MeOH, methanol (gray); Am., ammonium), with H2O only (no C or N added) provided as a background sample, after 864 d incubation. Addition of hydrogen to incubation is indicated by Y (yes). Samples are in paired columns or N added) provided as a background sample, after 864 d incubation. Addition of hydrogen to incubation is indicated by Y (yes). Samples are in paired columns.
the methods outlined by Hoshino and Inagaki (33). Resultant paired-end reads were processed and assigned taxonomy using QIIME (36) and a custom version of the UNITE, NR9, and SILVA databases (37, 38) optimized for marine sedimentary microorganisms. The total sequences and relative abundance of most abundant taxa are provided in Dataset S1, and all taxa are provided in Dataset S2.

Results

Evaluation of Incubation Cellular Abundances and Sample Purity

Stringent contamination monitoring during Expedition 337 suggested sandstone lithologies were the most susceptible to drilling mud contamination (27). Therefore, only shale and coal lithologies were used in this study. While cells were detected in the drilling fluids [2.66 × 10⁶ cells per milliliter (30)] and have the potential to contaminate downstream analyses (6), two lines of evidence suggest exogenous cell contamination was minimal in our long-term stable isotope incubation study. First, molecular screening of the 16S rRNA sequences recovered from our 37 °C and 45 °C incubations was distinct from that of sequences associated with common drilling mud contaminants from Expedition 337 (6). Second, cultivation attempts to enrich for bacteria (39) or fungi (40) from drilling fluids were unsuccessful, suggesting that contaminant cells had low viability and were unlikely to overgrow the indigenous microorganisms in our moderately thermophilic, low-nutrient coal and shale incubations. The low microbial viability of exogenous microorganisms may be due to the chemical composition of the drilling mud, containing bentonite and detergent chemicals, as well as the extreme physical conditions experienced during circulation of the high pH (10.5–11) drilling fluids between the 2-kmbfs drilling formation and the ship, with repeated cycling between oxic and anoxic conditions and temperatures ranging from 2–60 °C.

Cell abundances were calculated from concentrates after cell separation from the rock matrix (Table 1). Cell numbers measured at the end of the incubations, including no amendment controls, were all elevated relative to cell concentrations calculated from shipboard-processed samples, reported as 7 cells per gram of shale, 200 cells per gram of coal, and 942 cells per gram of mixed lithology (6). The elevated cell concentration in the incubated no-amendment controls suggested that the addition of water alone could have been enough to stimulate growth in this system using substrates native to the coal and shale. Microbial cells stained with SYBR Green I appeared as small cocci (single or doublets) or rods, with the exception of the mixed incubation, which also contained filamentous microorganisms (Fig. S1). The range of microbial cell sizes determined from NanoSIMS ion images yielded median cell lengths ranging from 0.7–1.2 μm in shale, 0.3–0.9 μm in coal, and 0.4–1.0 μm in the mixed incubations, consistent with previous reports from deep subsea floor sediments (41).

Assessing Methylated Substrate Metabolism by Bulk ¹³C and ²H Geochemical Analysis of Methane and DIC Measurements. Total methane production at the end of the incubation period was higher in coal incubations, relative to shale and mixed samples, and highest overall in nonautoclaved coal samples (Table 1). Lower, but detectable, levels of methane production were also measured in the mixed incubations. The 8¹³C–CH₄ was measured for coal incubations, which showed minimal ¹³C enrichment from the added ¹³C-methanol or methylamine (−42.3 to −58.8‰) relative in situ methane values [approximately −62‰ in both formation water and core gas sampling (6)]. Kinetic isotope effects during methane desorption could account for a portion of the methane enrichment [≥50‰ (42, 43)], but the higher ¹³C–CH₄ enrichments (−15–20‰) are unlikely explained by kinetic fractionation alone, and most likely contain some ¹³C-substrate incorporation. If all isotopic enrichment in these heaviest samples were from ¹³C substrate, it would correspond to ∼0.04% of total methane being sourced from the added ¹³C-methylamine or ¹³C-methanol. Since H₂ in the incubations is predicted to exchange with H₂O (44), any H₂-derived methane produced over the course of the incubations should also be enriched in δ¹³H–CH₄. As observed with ¹³C measurements, the values of δ¹³H–CH₄ were also minimally enriched in deuterium (~96.5 to ~198‰) relative to the in situ δ¹³H–CH₄ [approximately −200‰ in both formation water and core gas sampling (6)]. The most enriched δ¹³H–CH₄ was observed in incubations amended with ¹³C-methanol + ¹⁵N-ammonium + H₂ (~96.5‰). Isotope mass balance indicates this level of enrichment should at most account for <0.007% of the total methane recovered in the incubation. Therefore, both isotopic methods demonstrate consistently low, but detectable, levels of biogenic methane production in the coal incubations, with carbon isotope data supporting the presence of some methylotrophic methanogenesis.

Methylotrophic methanogenesis conversion of the ¹³C-labeled methanol or methylamine should also lead to the production of ¹³C-DIC in our incubations. Although shale incubations had overall higher DIC concentrations than incubations with only coal, DIC concentrations varied widely within each lithology. Methanol-amended shale incubations produced enriched ¹³C-DIC values, with the highest values observed in the ¹³C-methanol + ¹⁵N-ammonium + H₂ amendment (+430.0‰). In contrast, methylamine shale incubations did not show any appreciable enrichment (−2.86 to +8.32‰). The opposite trend was observed in the coal incubations, where ¹³C-DIC production from ¹³C-methanol was minimal (~15.14 to +26.67‰), but some of the ¹³C-methylamine incubations (autoclaved treatment) produced enriched ¹³C-DIC (+108.06‰). The ¹³C-DIC trends with the incubations were similar to what was observed in the coal incubations, with the highest enrichment in ¹³C-methylamine amendments (+327.56‰) and lower activity observed with the addition of ¹³C-methanol (+6.32 to +44.64‰). By comparison, the in situ gaseous ¹³C-CO₂ ranged from −19 to +15‰, with the most ¹³C-enriched CO₂ recovered from the coal beds (6). Assuming steady-state production during the incubation, the highest δ¹³C-DIC enrichment in coal would correspond to 2–0.2‰ DIC per cm³−1 or 0.2–0.1% of total moles of DIC generated from ¹³C-methylamine and ¹³C-methanol, respectively. In summary, the observed minor contributions of labeled methane and DIC production in our incubations indicated the potential for both methane- and DIC-generating metabolisms (methylotrophic methanogenesis) in addition to DIC-only-generating metabolisms (methyl fermentation).

Single-Cell Substrate Uptake and Generation Time Estimates of Deep Subsea Floor Microorganisms by ¹⁵N and H₂O. Of the 12 different sample conditions prepared for each lithology, eight shale, 10 coal, and six mixed lithology incubations had cells detectible by NanoSIMS after the cell separation protocol. The number of cellular ROIs analyzed per incubation ranged from nine to 306 (Fig. 1). Due to low cellular ¹³C enrichment relative to carbon background on NanoSIMS filters, single-cell ¹³C-Carbon enrichment was within instrument error of natural abundance, and was therefore not included. Although it cannot be determined if active cells were from the regeneration of existing biomass or the generation of new cells through division, at least one instance of active cells were from the regeneration of existing biomass or the generation of new cells through division at least one instance of active cells were from the regeneration of existing biomass or the generation of new cells through division at least one instance of active cells were from the regeneration of existing biomass.
coal samples. In shale incubations, the highest enrichment in bulk δ13C-DIC and cellular 2H and 15N was seen in treatments amended with 13C-methanol + 15N-ammonium + H2. Bulk δ13C-DIC was not enriched in 13C-methylamine–amended shale incubations, yet 2H and 15N (from 15N-ammonium) enrichment was still observed in cells from these incubations. The 15N-methylamine–amended shale incubations without hydrogen also showed high 2H and 15N incorporation. Finally, mixed lithology incubations had the highest δ13C-DIC (+327‰) in the 13C-methylamine + 15N-ammonium treatment, although this incubation did not yield enough cells to analyze by NanoSIMS. From the mixed incubations that could be analyzed by NanoSIMS, the 13C-methanol amendment had the highest cellular 2H enrichment (no 15N was added to this incubation).

In summary, both methanol and methylamine treatments produced 13C-DIC in coal and mixed incubations, but only methanol produced 13C-DIC in the shale incubations. Cells enriched with 15N were recovered from both 15N-methylamine– and 15N-ammonium–amended incubations in shale and coal incubations, but only 15N-ammonium showed cellular 15N uptake for mixed lithology incubations. The 2H-enriched cells were recovered across all substrate combinations and all lithologies. While bulk δ13C-DIC suggested methanol and shale, and methylamine and coal amendments and lithologies were correlated, single-cell incorporation suggests that viability and activity in these incubations were not uniformly linked to substrate addition or lithology.

Biomass generation times were calculated from the most robust NanoSIMS cellular ROI enrichments in 2H and 15N (Materials and Methods). As such, our estimates of microbial generation times represent values for the fastest growing cells within our incubations (Fig. 3), yet still result in generation time estimates from a few months to more than 100 y. The greatest overall 2H and 15N incorporation was from coal amended with 15N-methylamine, with much longer generation times for the rest of the incubations, which all had hydrogen added. This concurs with 13C-DIC production results, where samples containing coal had depressed activity when H2 was added. This sample also had the greatest offset between 2H and 15N enrichment, where generation time estimates calculated from cellular 2H2O enrichment were shorter than 15N-nitrogen enrichment from methylamine. The mixed lithology samples amended with 15N-ammonium or 15N-methylamine as the nitrogen source had more comparable uptake of 15N and 2H, with slightly higher enrichment observed in the 15N-ammonium treatment. This suggests that while ammonium and methylamine can both be assimilated into biomass, ammonium was more readily used for anabolism compared with methylamine.

**Comparison of 16S rRNA Gene Profiles to Geochemically Predicted Metabolisms.** Microbial diversity surveys using 16S rRNA gene iTAG sequencing were generated from FAC sorting of the seven most active shale (n = 2), coal (n = 3), and mixed lithology (n = 2) SIP incubations after 2.5 y of incubation at 37 °C and 45 °C (Datasets S1 and S2). The recovered microbial diversity was broadly similar to the in situ microbial assemblage reported from the same coal and shale beds described by Inagaki et al. (6), with an abundance of anaerobic heterotrophic groups that more
Epsilonproteobacteria (Campylobacterales). At the genus level, proteobacteria, Gammaproteobacteria, and Deltaproteobacteria, bacteria, Firmicutes, Bacteroidetes, Alphaproteobacteria, Betaproteobacteria, Dehalococcoidia, Deinococcus, Verrucomicrobia (Opitutae), and with additional representatives belonging to the Chloroflexi. 

Relative to \( \text{CH}_3\text{NH}_2 \) and \( \text{NH}_4^+ \), reaching 76% of the total diversity, consistent with the dominance of thermotolerant. Thermophilic and spore-forming microorganisms were capable of persisting or growing in the long-term incubations at elevated temperatures and anoxic or microoxic conditions. The fact that the microbial taxa recovered were derived from intact FACS cells rather than environmental sample DNA extracts suggests that the diversity within the deep coal and shale SIP incubations is likely associated with viable microorganisms that were capable of persisting or growing in the long-term incubations at elevated temperatures and anoxic or microoxic conditions.

Closely resembled terrestrial and wetland microbial lineages than marine sediments, many of which are generally associated with heterotrophs described from coal bed methane systems (45–47). The microbial taxa recovered were derived from intact FACS cells rather than environmental sample DNA extracts suggests that the diversity within the deep coal and shale SIP incubations is likely associated with viable microorganisms that were capable of persisting or growing in the long-term incubations at elevated temperatures and anoxic or microoxic conditions.

The 16S rRNA gene sequences within the bacterial domain were dominant in our diversity surveys of the seven most active incubations, with rare archaeal sequences affiliated with marine benthic group B detected in only two of the shale incubations incubated at 37 °C (Dataset S2). Some recovered taxa in the 45 °C incubation were related to moderately thermophilic microorganisms [e.g., *Thermomonas* within the Gammaproteobacteria (48). *Acidothrix* within the Acinetobacteria (49)], although the majority of the lineages recovered are not known for being thermotolerant. Thermophilic and spore-forming microorganisms often show enhanced cultivation and growth after high-pressure temperature treatment (50), consistent with the dominance of Firmicutes from the autoclaved coal incubation amended with methylene and ammonium, reaching 76% of the total diversity, relative to 2–3% Firmicutes within the parallel nonautoclaved incubation (Dataset S1).

Dominant bacterial groups in the incubations included Actino bacteria, Firmicutes, Bacteroidetes, Alphaproteobacteria, Betaproteobacteria, Gammaproteobacteria, and Deltaproteobacteria, with additional representatives belonging to the Chloroflexi (Dehalococcoida), Deinococcus, Verucomicrobia (Opitutae), and Epsilonproteobacteria (Campylobacterales). At the genus level, microbial assemblages differed between samples, with no genus shared across all seven samples, and 37% of genera were found only in one of the seven samples. These differences were not statistically correlated with substrate amendment or lithology, despite bulk geochemistry analysis suggesting a preference for methanol utilization in shale and methylene utilization in coal. There were also no statistically supported differences in the recovered diversity associated with lithology between the shale, coal, and mixed samples. When observed at lower taxonomic resolution and grouped by assigned taxonomic phylum, the microbial assemblages share the same dominant phyla across all incubations (Dataset S1). Therefore, while microdiversity exists at the genus level, the dominant phyla maintained over the 2.5-y incubation period were predominantly heterotrophic microorganisms (e.g., Gammaproteobacteria were at ≥10% relative abundance in all incubations).

Within these heterotrophic bacteria, there was some correlation between relative abundance of specific microbial taxa and hydrogen addition. For example, both shale and mixed lithology samples with 13C-methanol had a higher relative abundance of Firmicutes and Alphaproteobacteria in the absence of hydrogen addition to the headspace. In comparison, parallel hydrogen-amended incubations contained a higher percentage of Actinobacteria, suggesting potential utilization. While commonly described from deep subsurface biofilm ecosystems, sequences affiliated with the Chloroflexi were only abundant (28% relative abundance) in the 13C-methanol mixed lithology incubation. Many members of the Chloroflexi have a characteristic filamentous morphology, making them likely candidates for the abundant filamentous cells recovered from this incubation (Fig. S1F).

While we did not recover sequences for methylotherrophic methanogens, some of the bacterial lineages have cultured representatives reported to use methanol and/or methylene as the sole carbon or energy source, including members of the *Nocardia* within the Actinobacteria (51), Rhizobiales [e.g., *Rhodopseudomonas acidiphilus* (52)], Desulfovibionales (*Desulfovibrio fractuovosorum*) (53), and Firmicutes (54), especially many of the thermophilic spor-forming bacilli (55, 56). Other detected bacteria may contribute to methanol and methylene production through breakdown of organic compounds, a process that has also been documented in marine sediments (57). For example, Verrucomicrobia (*Opitutus*) recovered in some of the coal and shale incubations has cultured representatives that are capable of producing methanol during pectin degradation (58). Select members of the *Clostria* also produce methanol from peroxides and methanol, as well as other compounds such as methyamines (54), while others within this class consume methanol and methoxylated compounds as sole carbon and energy sources.

**Discussion**

IODP Expedition 337 off Shimokita Peninsula explored a pristine, deeply buried, Miocene-aged coal bed system in an attempt to characterize the deep subsurface biosphere within this unique environment. The initial site characterization from this expedition, described by Inagaki et al. (6), demonstrated higher cell concentrations in the coal bed compared with surrounding lithologies but reported in situ cell counts for this site that were far below the predicted depth-based trends recorded in previous ocean drilling studies (5). Successful cultivation of a hydrogenotrophic methanogen (*Methanobacterium sp.*), combined with geochemical evidence for sulfate reducers and methanogens in isotopically enriched pyrite (14) and 13C-depleted methane, as well as detection of coenzyme F420 (6), indicated the potential for biogenic methane formation in the organic carbon- and H₂-replete coal bed environment (6). The calculated Gibbs free energies of methylotrophic methanogenesis and methyl fermentation under in situ conditions indicate that both methylotrophic methanogenesis (approximately −300 kJ/mol) and fermentation (approximately −10 kJ/mol) were possible, but methylotrophic methanogenesis is more energetically favorable. Given these energetic predictions, it was unexpected that the coal and shale SIP
experiments with $^{13}$C-labeled methylated substrates did not show significant biological methanogenic activity during the 2.5-y incubation period. Instead, our isotopic data suggested the majority of methane detected during these long-term experiments was associated with the coal before incubation and subsequently released from the sample matrix over time, with minimal contributions from methylotrophic methanogenesis via the $^{13}$C-labeled substrates (at most, 0.04% of total methane from labeled substrate). This could also explain why the autoclaved coal samples had lower methane production, as the heat and pressure treatment during autoclaving, followed by flushing with argon, may have accelerated degassing of methane compared to treatments without autoclaving. Our finding that the majority of methane produced was not enriched in $^{13}$C may be attributed to other carbon substrates native to the organic-rich coal bed. However, the methane produced in the incubations also did not show enrichment in $^2$H, which is predicted to occur during active biogenic methane production in the presence of $^2$H$_2$O, as biogenic methane production requires water-derived hydrogen (59). From incubation $^2$H-CH$_4$ values, we calculated a maximum of 0.007% of the total methane could have been biologically produced during the coal incubations from utilization of any carbon substrate available in the coal, or added methylated substrate. The combined $^{13}$C and $^2$H data point to the majority of biogenic methane recovered over the 2.5-y experiment being produced before the start of the incubation, highlighting an apparent temporal offset between in situ generation and experiment being produced before the start of the incubation, high-to-the majority of biogenic methane recovered over the 2.5-y ex-

versatile methanogens were impacted more than the comparatively metabolically versatile bacterial assemblage.

Overall, the microbial diversity recovered from our incubations was consistent with the bulk microbial diversity recovered in situ (6), including a dominance of Gammaproteobacteria, Betaproteobacteria, Alphaproteobacteria, Actinobacteria, Firmicutes, Chloroflexi, and Bacteroidetes. The combined multidisciplinary results from our stable isotope incubation experiments suggest that these deep subsurface microorganisms are metabolic generalists that appear to be resistant to the unique stressors of the in situ environment beyond what has been described of their better characterized surface relatives in culture. While total cell numbers and sample sizes were low, as expected given this environment’s inherent constraints, there were general trends in relative abundance of specific taxonomic groups aligning with hydrogen amendment inhibition (Firmicutes, Alphaproteobacteria) or stimulation (Actinobacteria), as well as observations of increased numbers of putative spore-forming Firmicutes in the autoclave-treated sample. Gibbs free energies calculated for our incubation conditions with hydrogen addition (~10-fold in situ hydrogen, +10 kJ/mol) and no hydrogen addition (near zero hydrogen approaches favorability of methanogenesis) concur with the coupling of fermentation and hydrogen addition, where incubations without hydrogen amendment should be much more favorable for fermentation. It is also notable that methanol anabolism (modeled via conversion to formaldehyde, approximately ~30 kJ/mol at standard temperature and pressure) is more energetically favorable than catabolism, which may have elevated importance in subsurface environments, where most, if not all, cellular energy is likely devoted to cell maintenance (61, 62).

The highest rate of $^{13}$C-DIC production from methyl substrate amendments in our coal incubations (2 pmol of DIC per cm$^3$ day$^{-1}$) was 25-fold faster than potential sulfate reduction rates from the same coal bed with radiolabeled sulfate (14). Using a conversion factor to translate radiolabeled sulfate rates to DIC production rates [2 mol of C from 1 mol of acetate, per 1 mol of sulfate (61)], other subsurface systems yield rates ranging from ~10$^{-4}$ fmol of C per cell per day in Peru Margin sediments (13) to shallow sediment rates of ~10$^{-2}$ fmol of C per cell per day (12). Converting $^{13}$C-DIC production in our coal incubations to cellular rates using final cell concentrations, DIC production was ~0.02 fmol of C per cell per day for both methylamine and methanol. Assuming a minimal extraction efficiency of only 10%, this would decrease rates to ~0.002 fmol of C per cell per day, which is still above Peru Margin sediment rates. Our NanoSIMS data also indicate that it is unlikely all cells are active in our incubations (Table 1), and when assuming only the active portion of cells in these coal incubations produced the labeled $^{13}$C-DIC, rates are estimated to be 30-60% higher. Accepting that both ex situ stable-isotope and radioisotope incubation methods likely caused some perturbation to the natural system, either through the addition of micromolar concentrations of carbon and nitrogen (this study) or through homogenization of the sample and shaking during $^{35}$S-sulfate incubation (14), these rates are unlikely exact representations of in situ subseafloor activity, but remain consistent with the multiple independent indications of slow rates of metabolism and growth of deep biosphere microorganisms relative to pure culture studies.

Both $^{15}$N-ammonium and $^2$H$_2$O have been used to assess anabolic activity and generation times of diverse microorganisms in culture and the environment (18, 19, 21, 63, 64). In the context of this single-cell–targeted multiple-isotope labeling study, the passive tracer $^2$H$_2$O was observed to be a more sensitive measure of overall anabolic activity compared with $^{15}$N assimilation from $^{15}$N-ammonium or methylamine within the deep subsurface biosphere. However, when these assimilation rates were converted to biomass generation times, the paired $^2$H-based and $^{15}$N-based generation time estimates were consistent (Fig. 3). This suggests that the most active cells were not selectively producing different
types of biomass, such as repairing lipids over generating new proteins, but were generally producing all forms of biomass. Comparison of cellular enrichment from $^{2}$H$_{2}$O incorporation across different incubations revealed a notable difference in cellular anabolic activity (generation time) in incubations supplied with $^{2}$H$_{2}$ relative to those without. $^{2}$H$_{2}$ concentrations have been reported as low as 10$^{-5}$ $\mu$M in subsurface coal beds with active hydrogenotrophic methanogenesis (65), but in situ levels were much higher in the Shimokita coal bed (1-500 $\mu$M) (6), suggesting that methanogenesis was not the dominant control over environmental hydrogen concentrations (66). In contrast to many deep biosystems that are hypothesized to be fueled by hydrogen (67), the addition of hydrogen (0.15 atm $^{2}$H$_{2}$) depressed microbial anabolic activity relative to those incubations without hydrogen for the coal incubations (Fig. 3), and suggests that fermentative microorganisms suggested by 16S rRNA gene sequencing are active in coal bed incubations. Microorganisms in the shale incubations from an overlying horizon produced the opposite trend. In this set of incubations, a larger fraction of active cells (ROIs) was detected with $^{2}$H$_{2}$ amendment, relative to the paired incubation without $^{2}$H$_{2}$, in both $^{13}$C-methylamine and $^{13}$C-methanol treatments (Fig. 1). These findings suggest the influence of $^{2}$H$_{2}$ is complex in organic-rich systems, and as observed in the deep coal horizon, $^{2}$H$_{2}$ may not always be a beneficial source of energy and, in fact, may depress microbial activity and growth.

Patterns of nitrogen source utilization (ammonia or methylamine) by the active microbial assemblage also differed between the coal and shale incubations. The $^{15}$N-ammonium incorporation was observed in the majority of incubations, independent of lithology. Methyloxyrophycrea via the microbial assimilation of $^{15}$N-methylamine, however, was predominately observed by the coal-associated microbial assemblage. It is expected that anaerobes can use ammonium as a nitrogen source, but assimilation of methylamine is less common, and the observed difference in methylamine utilization between the coal and shale incubations is notable. Supporting these observations, the diversity recovered in the coal incubations included microbial taxa affiliated with bacterial lineages shown to be capable of methylamine metabolism. These findings may point to the potential use of methylated amines in lignite coal as a source of nitrogen in this deep biosphere ecosystem.

We compared the $^{15}$N-estimated generation times from previous SIP-NanoSIMS data of $^{15}$N-ammonium incorporation from a single site to values in cells from deep marine sediments (219 mbsf) that overlay the coal and shale beds studied here (18). In the study by Morono et al. (18), $^{15}$N-ammonium–based estimates of doubling times ranged from 63 to 252 d depending on added carbon source (e.g., glucose, amino acids), whereas the fastest $^{15}$N generation times calculated from our deep coal bed incubations were ∼30-fold slower (∼20 y; Fig. 3). The study by Morono et al. (18) did not include $^{2}$H$_{2}$O precluding direct comparison; however, estimates of $^{2}$H-based generation times from coal bed microorganisms did show some overlap with the slowest rates measured within the overlying sediment microbial assemblage (1-2 y) and other marine sediment studies using $^{2}$H$_{2}$O (∼20 y, $^{2}$H-lipid incorporation rates (25)). Our inability to detect $^{13}$C-labeled biomass despite indication of carbon utilization from the bulk geochemical data also supports the findings of the study by Morono et al. (18), where nitrogen was shown to be assimilated to a greater extent than carbon by microorganisms in their sediment incubations.

Overall, the single-cell generation times from SIP-NanoSIMS data for carbon, nitrogen, and $^{2}$H$_{2}$O in our study, as well as those reported by Morono et al. (18), yield estimates that are one to two orders of magnitude faster than deep biosphere turnover times determined from bulk methods, such as amino acid race-lipid and lipid degradation (17, 65). The simultaneous use of $^{15}$N-labeled substrates in combination with the passive $^{2}$H$_{2}$O tracer offered two independent means to track single-cell anabolic activity within our incubations. Although our study focused on the most active cells in our generation time estimates, this targeted single-cell analysis illustrates the high level of heterogeneity in anabolic activity between cells within the subsurface microbial community, with the active fraction of the community ranging between no cells above the enrichment threshold (10-fold natural abundance) to 76% of the total cells with values above the threshold. This suggests that bulk methods estimating average cellular activity through the incorporation of environmental DNA cell counts are likely to underestimate microbial activity and generation times. The complementarity in these independent multiscale approaches offers an important frame of reference for setting minimum and maximum bounds for growth and metabolic activity estimates within deep subsurface ecosystems and demonstrates that even at the fringes of microbial activity, survival is not a simple story of dead or alive.

Conclusion

Despite cell abundances that were substantially lower than expected from global depth trends and sediment surface concentrations, this single-cell–targeted study successfully detected active microorganisms in terrestrial-sourced shale and coal buried at 2 kmbsf for millions of years. Incubations of intact shale and coal pieces maintained at in situ temperatures and amended with heavy metal compounds, ammonium, and $^{2}$H$_{2}$O enabled the detection of some of the slowest measurements of growth using SIP-NanoSIMS. With biomass generation times of years to hundreds of years, these rates were much faster than deep biosphere activity estimates via other bulk methods (approximately thousands of years). Although methanogenic activity could be detected via geochemical analysis, methanogenic archaea were not recovered by 16S rRNA gene iTag sequencing. Instead, heterotrophy appears to be the predominant microbial metabolism for this system, which has been found in many deep subsurface sedimentary environments (8, 68–70). Although variation in activity among incubations associated with methyl substrate amendment, presence/absence of hydrogen, high temperature/pressure (autoclave), and 16S rRNA diversity depict a heterogeneous and diverse heterotrophic community. Through this unique snapshot into a pristine coal bed, results from Expedition 337 have also opened new avenues for conceptualizing the residence time of carbon, nitrogen, and sulfur in coal deposits and global biogeochemical cycles.

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