Aerobic microbial life persists in oxic marine sediment as old as 101.5 million years

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Sparse microbial populations persist from seafloor to basement in the slowly accumulating oxic sediment of the oligotrophic South Pacific Gyre (SPG). The physiological status of these communities, including their substrate metabolism, is previously unconstrained. Here we show that diverse aerobic members of communities in SPG sediments (4.3–101.5 Ma) are capable of readily incorporating carbon and nitrogen substrates and dividing. Most of the 6986 individual cells analyzed with nanometer-scale secondary ion mass spectrometry (NanoSIMS) actively incorporated isotope-labeled substrates. Many cells responded rapidly to incubation conditions, increasing total numbers by 4 orders of magnitude and taking up labeled carbon and nitrogen within 68 days after incubation. The response was generally faster (on average, 3.09 times) for nitrogen incorporation than for carbon incorporation. In contrast, anaerobic microbes were only minimally revived from this oxic sediment. Our results suggest that microbial communities widely distributed in organic-poor abyssal sediment consist mainly of aerobes that retain their metabolic potential under extremely low-energy conditions for up to 101.5 Ma.
Marine sediment covers ~70% of Earth’s surface. It harbors a remarkable microbial population that comprises 12–45% of total microbial biomass or ~0.6–2% of total living biomass on Earth1–4. Since the discovery of the subseafloor sedimentary biosphere5, understanding on the extent, diversity, and physiology of the microbes below the seafloor has greatly expanded6. In 2010, Integrated Ocean Drilling Program (IODP) Expedition 329 retrieved sedimentary sequences from the abyssal plain of the South Pacific Gyre (SPG), to examine subseafloor life and habitability in the lowest productivity region of the world ocean7. Although their abundance is very low, microbial cells are present throughout the entire sedimentary sequence at all the SPG sites. Their abundance (2.2 × 103 to 5.5 × 106 cells cm−3) is one to seven orders of magnitude lower than counts at the same depths in sites of ocean margins and upwelling regions8. The presence of dissolved O2, nitrate (NO3−), phosphate (PO43−), and dissolved inorganic carbon (DIC) throughout the sedimentary sequence from the seafloor to the volcanic basement indicates that cell abundance and activity are not limited by availability of electron acceptors or dissolved major inorganic nutrients from the Redfield stoichiometry of net dissolved O2 reduction to net nitrate production in the sediment, the microbial cells have been inferred to consume oxygen coupled to oxidation of marine organic matter at extremely slow rates8.

Direct evidence of the physiological nature and survival status of microbial cells in this extremely energy-poor setting is previously lacking. A cell must metabolize a certain amount of carbon relative to its own biomass before it can double its size, divide or even sustain a metabolically active state (basal power requirement9). Bioenergetic calculations indicate that the microbes in SPG abyssal clay have access to very little power10. Very low permeability (1.1–2.0 × 10−17 m2 for IODP Site U1365 4H-3 [26.6 meters below seafloor (mbsf)] and 8.9 × 10−18 m2 for IODP Site U1370 [37.5 mbsf], respectively)11, very low estimated pore size of the abyssal clay (~0.02 microns, calculated using above permeability data according to the equation shown in Tanikawa et al.12), and thick porcellaneous layers above the oldest sampled horizons appear to preclude cell migration into the sampled sediment. Consequently, the sampled communities have likely been trapped in the sediment since shortly after its deposition. The physiological status and growth potential of these buried communities and, more generally, the fractions of these energy-starved subseafloor microbes that are alive, dormant, or dead, have been essentially unknown.

To document the ecophysiology of the microbes that persist in these sedimentary habitats, we set up microaerobic incubation experiments with stable isotope-labeled substrates as tracers for microbial anabolic activities in mini-plugs of ancient (up to 101.5 Ma) pelagic clay (from IODP Sites U1365 and U1370) and relatively young (up to 13 Ma) but extremely oligotrophic calcareous nannofossil ooze (Site U1368, Supplementary Fig. 1). The results—the low relative abundance of spore formers, the rapid initial increases in incubated cell abundance, the high percentages of active cells in the incubations, and the confining nature of their sedimentary habitat—collectively suggest that microbial communities in oxic subseafloor sediment persist in metabolically active form for at least 101.5 million years.

Results and discussion

Single-cell anabolic activity with 13C-, 15N-labeled substrates. Single cell-targeted ion imaging analysis of 6986 individual cells by NanoSIMS showed that aerobic microbes from every sample of this oligotrophic oxic sediment actively take up isotope-labeled carbon and nitrogen substrates, even from sediment as old as 101.5 Ma13 (U1365 9H-3) (Figs. 1 and 2b, c). The number of cellular regions of interests (ROIs) analyzed per incubation sample ranged from 9 to 210 (Supplementary Data 1). Incorporation of carbon and nitrogen from organic compounds and ammonium was generally seen in the ROIs from all of the microaerobic (~3.3% O2 in the headspace) incubations, with bimodal distribution in the degree of substrate incorporation (Fig. 2b). While initial cell counts were extremely low, ranging from 102 to 3 × 103 cells cm−3 (Fig. 2a)8, the microbes in the oxic sediment responded rapidly to the substrate injection in microaerobic incubations, increasing total numbers by orders of magnitude and taking up labeled carbon and nitrogen within 68 days of incubation.

In contrast to the microaerobic incubations, incubations of ancient (65.5 Ma13) oxic abyssal clay from Site U1370 conducted without adding O2 showed only marginal incorporation of selected substrates with minimal growth of biomass (Supplementary Fig. 2). Although we did not supply additional O2, the oxic sediment contained ~1.5 µM of O2 in its interstitial water1, which introduced 16.4 nmol of O2 in the incubation vial and reached an equilibrated concentration of not more than 14.6 nM in the sediment. Although the slight increase of biomass at the early stage of the U1370 incubations may have been aerobic microbial growth, there was not enough O2 to sustain aerobic growth for the entire period of incubation. Although the sediment contained abundant dissolved nitrate (~46 µM) and sulfate (~27 µM) in its interstitial water1, we did not see intensive growth of biomass in later stages of the incubation. This result suggests that metabolic activity hardly revived and did not persist in the absence of added O2 for microbial communities that have been exposed to dissolved O2 for 65.5 million years. This, in turn, suggests that long-time exposure to dissolved O2 over geological time may extinguish anaerobic microorganisms including anaerobic spore-formers (or at least make them hard to revive).

The microbial metabolic response to microaerobic inoculation conditions was heterogeneous; the extent and rate of labeled isotope incorporation varied considerably from one ROI to another (Fig. 2b, c), and from one labeled substrate to another, similarly to the substrate incorporations seen in anaerobic incubations of deep subseafloor coalbed sediment14. Of the labeled substrates tested, heterotrophic substrates generally supported biomass growth. However, in the case of nannofossil-bearing clay of U1368 2H-5 sediment, the amino acid-supplemented incubation showed biomass increase only at day 21 of incubation and afterward decreased to the level of original cell abundance, indicating the loss of increased biomass at a later stage of incubation. We observed only marginal autotrophic incorporation of bicarbonate (0–23.6 atom%, in average 0.8 atom%, Fig. 2b), consistent with primary reliance on organic matter as carbon source for biomass production.

Rates of substrate uptake and biomass generation. The response to substrate addition, in general, was faster (on average, 3.09 times, Table 1) and more extensive for nitrogen incorporation than for carbon incorporation (Fig. 2c). Ammonium, which was added as a nitrogen source in all incubations except for the incubation with 13C and 15N-labeled amino acids, was generally incorporated quickly into cellular biomass. The preferred incorporation of supplemented ammonium over nitrate, which originally existed at ~40 µM in all of the sediment samples5, was predictable given higher energetic costs of reducing nitrate to the oxidation state of N in amino acids15. Although ammonia assimilation requires energy to be incorporated into amino acids15,16, its incorporation was observed even under the condition supplemented with only ammonium without a carbon source. In addition, the high ammonia incorporation and biomass increase during those incubations suggests at least a minor contribution of chemolithoautotrophic ammonia oxidizers17.
The growth rates of microbes were calculated based on the rates of biomass growth and substrate incorporation (Fig. 3a, Table 1, and Supplementary Data 2). The biomass-based specific growth rates were generally higher than the substrate-based biomass generation rates. This difference suggests that the active microbial communities utilized carbon and nitrogen compounds in the sediment other than the supplemented isotope-labeled substrates for their biomass growth. Although the total organic carbon content in the sediment is very low (0.01–0.02 weight percent), it exists in bioavailable form but protected under in situ conditions16. This amount of total organic carbon is enough to sustain the observed growth of microbes without an additional carbon source; it corresponds to potential microbial biomass19 of >10^10 cells cm^-3, exceeding the original microbial biomass by 7–8 orders of magnitude, and the post-incubation biomass by 4 orders of magnitude.

To document the physiological status of these microorganisms, we calculated the fraction of microbial cells that originally existed in the samples. The calculation was made by using the observed active ROI fractional ratio (number of active ROI versus total ROI analyzed), increase of biomass, and ROI-based sampling depth (the numerical depth of sampling that the ROI-based analysis covered). The observed “active” ROIs ranged from 18.4 to 100% of the total ROIs analyzed, giving an average of 92%. The minimum original active fractions average 77.7%, ranging from 24.1 to 99.1% (Table 1 and Supplementary Data 2). Even in the sample of oldest sediment (U1365 9H-3, 101.5 Ma13), the revivable heterotrophic population was 39.6–99.1% of the original community. These microaerobic incubations differ notably from the previously published anaerobic incubation14 by rapidly and consistently increasing in biomass after the addition of substrates. The biomass-based doubling time (bTd) of microbes in these microaerobic incubations (Supplementary Data 2) averaged 4.9 days, ranging from 1.4 to 17.8 days. Incubations with the oldest sediment (U1365C 9H-3) showed significantly longer bTd (in average 9.8 days, pairwise t-tests, p < 0.05) than the incubations with younger sediment. The incubations with U1365C 8H-2 (95.4 Ma13) showed slightly longer but statistically insignificant bTd values than the U1368 sediment samples (Table 1).

“Inactive” ROIs, which have no detectable incorporation of supplemented substrates, represent either (i) dormant cells or (ii) dead cells (necromass). Although recycling of biomolecules from necromass is a well-known anabolic strategy in marine sediment20,21, necromass provides only a small fraction of the maintenance power demand for the microbial community in subsea floor sediment22. In our amino-acid-amended incubations, an apparent biomass decrease after an initial increase in the sample of U1368D 2H-5 (Fig. 2a) demonstrates the generation of new biomass and the remineralization after cell death (necromass) at a later incubation stage. The absence of substrate incorporation in the remaining “inactive” cells (accounting for 73.3% of total cells) at the 557-day timepoint suggested that they should have been inactive throughout the incubation period. Therefore, we infer that these cells comprise living (or at least degradation-resistant) but dormant biomass not revived by the addition of amino acids. This pattern of biomass change was not obvious or absent in the incubations with other substrates.

Comparison of 16S rRNA gene profiles to substrates. We investigated the 16S rRNA gene-based taxonomic composition of microbial cells sorted by fluorescence-activated cell sorting (FACS) from each incubation. Although pilot studies showed the community composition in anoxic sediment outside the SPG23, near-sea floor SPG sediment (<10 cmbsf)24, and SPG surface seawater25, diversity of microbial community in SPG sediment more than a meter below seafloor has not been reported. This absence of community composition information is partially due to the very low cell abundance and the nature of the sediment samples (dominated by fine-grained clay and silt with negligible sand26) that may have adsorbed released DNA from lysed cells during the DNA extraction process. For this study, we separated and sorted cells from the sediment to obtain community taxonomic composition under all of the incubation conditions, including initial sediment samples with very low biomass (~10^2 cells cm^-3) (full dataset can be found in Supplementary Data 2 and detailed methods to eliminate procedural contaminations are found in “Methods” section). The communities of all samples were dominated by bacterial sequences. Dominant bacterial groups included...
Actinobacteria, Bacteroidetes, Firmicutes, Alphaproteobacteria, Betaproteobacteria, Gammaproteobacteria, and Deltaproteobacteria (Fig. 3b, c) with a minor fraction of Chloroflexi (0–2.6%). A small fraction (~7%) of Terrestrial Hot Spring Group of thermophilic Archaea, which was recovered from sediment overlying relatively young (13.5-Ma) basement on the East Pacific Rise flank, was detected in sample U1368D 2H-5 (Supplementary Data 2). The absence of obligately autotrophic taxa and ammonia-oxidizing microorganisms confirmed the marginal incorporation of 13C-labeled bicarbonate and the dominance of heterotrophic microbial survival at extraordinarily low energy fluxes in sub-seafloor sediment are largely unexplored. Spore formation has been suggested as one of the possible mechanisms for such long-term survival of microorganisms29–31. Spore-forming lineages, however, were only a minor constituent of the microbial communities. Firmicutes related to thermophilic spore-forming microorganisms (Thermococcus32 and Carboxydothermus33) were detected at 0–12% (averaging 0.87 and 0.92%, respectively) throughout the samples and incubations. A small fraction (~2.5%) of the populations in incubations of the oldest sediment sample (U1365C 9H-3) is related to the genus Paenibacillus34, mesophilic spore-forming bacteria. This low relative abundance.

Abundance of spore formers. The mechanisms that allow microbial survival at extraordinarily low energy fluxes in sub-seafloor sediment are largely unexplored. Spore formation has been suggested as one of the possible mechanisms for such long-term survival of microorganisms29–31. Spore-forming lineages, however, were only a minor constituent of the microbial communities. Firmicutes related to thermophilic spore-forming microorganisms (Thermococcus32 and Carboxydothermus33) were detected at 0–12% (averaging 0.87 and 0.92%, respectively) throughout the samples and incubations. A small fraction (~2.5%) of the populations in incubations of the oldest sediment sample (U1365C 9H-3) is related to the genus Paenibacillus34, mesophilic spore-forming bacteria. This low relative abundance.

This is consistent with the suggestion from NanoSIMS-based results that the active microbial community utilizes native carbon and nitrogen compounds other than the substrates added to the incubation, and additionally suggest that the added substrates just stimulated the activity of microbes for their utilization of native carbon and nitrogen compounds. Rather the community composition showed dependence on incubation time, which suggests that different taxa responded to incubation at different timescales (Fig. 3c).

In general, microbial community composition in the incubations did not closely relate to the specific substrate added (Fig. 3c). This is consistent with the suggestion from NanoSIMS-based results that the active microbial community utilizes native carbon and nitrogen compounds other than the substrates added to the incubation, and additionally suggest that the added substrates just stimulated the activity of microbes for their utilization of native carbon and nitrogen compounds. Rather the community composition showed dependence on incubation time, which suggests that different taxa responded to incubation at different timescales (Fig. 3c).
of known spore formers suggests that the revival of spores (germination) was very limited in the incubations.

Methods

Materials. Sediment samples used in this study were collected by IODP Expedition 329 in the South Pacific Gyre (SPG; Supplementary Fig. 1)\(^\text{1}\). The samples collected from Site U1365 (23° 51.0377'S, 165° 38.6502’W) are zeolitic and/or metallerific pelagic clays, those from Site U1368 (27° 54.9920’N, 123° 09.6561’W) are calcareous nannofossil oozes, and those from Site U1370 (41° 51.1267’N, 153° 06.3674’W) are black metallerific clay, containing light yellowish brown clay-bearing nannofossil ooze. The water depths of these sites are respectively 5697, 3739, and 5075 meters below sea level (mbsl) at U1365, U1368, and U1370. Whole-round core samples of U1365C 8H-2 (obtained from 68.9 meters below seafloor [mbsf], 95.4 Ma), U1368D 1H-2 (1.6 mbsf), U1368D 2H-5 (14.7 mbsf), and U1370F 7H-6 (62.9 mbsf) were used for incubation experiments as described in the following section. Those samples were from horizons with no observable coring disturbance by visual core description\(^\text{35}\). Drilling fluid contamination assessment by chemical tracer revealed minimal drilling contamination of the samples (≤1 cell/g-sediment for U1365C 8H-2 and U1368D 2H-5, below detection to 0 cell/g-sediment for U1365C 8H-2 and 1368D 2H-5, below detection to 0 cell/g-sediment for U1365C 8H-2 and 1368D 2H-5)\(^\text{35}\). All Expedition 329 sediment age estimation is available in reports by Dunlea et al.\(^\text{13}\) and Alvarez Zarikian et al.\(^\text{36}\).

Incubation experiments. To identify autotrophic and heterotrophic microbial populations, as well as their potential rates for growth and substrate uptake, sterile samples were incubated with stable isotope-labeled substrates (\(^{13}\)C-glucose, \(^{13}\)C\(_2\)-acetate, \(^{13}\)C\(_3\)-propionate, \(^{13}\)C\(_4\)\(_{-}\)bicarbonate, \(^{15}\)N-amino acids mix [mixture of 20 Amino Acids], and \(^{15}\)N-ammonium). The incubation experiments were initiated onboard during expedition. Avoiding outer edges of the sediment cores (which are disturbed by visual core description\(^\text{35}\)). Drilling fluid contamination assessment by chemical tracer revealed minimal drilling contamination of the samples (≤1 cell/g-sediment for U1365C 8H-2 and U1368D 2H-5, below detection to 0 cell/g-sediment for U1365C 8H-2 and 1368D 2H-5)\(^\text{35}\). All Expedition 329 sediment age estimation is available in reports by Dunlea et al.\(^\text{13}\) and Alvarez Zarikian et al.\(^\text{36}\).
Fig. 3 Microbial growth response to the addition of substrates. a Scatter plots of microbial growth characteristics calculated from biomass (biomass-based specific growth rate, $\mu_B$ [day$^{-1}$]), carbon and nitrogen substrate (substrate-based biomass generation rates, $\mu_{SN}$ [day$^{-1}$]) for carbon and $\mu_{SN}$ [day$^{-1}$] for nitrogen, respectively. $n = 76$ samples were analyzed for the calculation of $\mu_B$ and $n = 6028$ cells were analyzed for the calculation of $\mu_{SN}$ and $\nu_B$, respectively (of totally 6253 cells analyzed, non-zero values of $\mu_B$ and $\mu_{SN}$ were used for this plot). b Relationship of community composition to addition of substrates along the time of incubation. The community compositions at “incubation time 0” show the compositions before incubation. c Non-metric multidimensional scaling (NMDS) ordination plot based on Bray–Curtis dissimilarities of community compositions at different incubation conditions and time points (stress = 0.209). Arrows represent environmental variables that were significantly ($p < 0.005$, permutations = 10000) related to the ordination. $n = 76$ samples (one for incubation time 0 and three for incubation sampling points [days 21, 68, and 557] per substrate for each sediment sample).

NanosIMS analysis of single cell-image acquisition and data processing. Cell targets were identified by fluorescence of SYBR Green I stain and marked on NanoSIMS membranes with a laser dissection microscope (LMD6000; Leica Microsystems) for ease of rediscovery on the NanoSIMS (an example is shown in Supplementary Fig. 3b). Microbial cells that incorporated stable isotope-labeled substrates were analyzed using NanoSIMS 50L (AMETEK Co. Ltd., CAMEGA RU) at the Kochi Institute for Core Sample Research, JAMSTEC. Samples on the ITO-coated polycarbonate membrane were pre-sputtered at high beam currents (30 pA/μm$^2$) before measurement. The $^{12}$C$^{15}$N$^{-}$, $^{13}$C$^{15}$N$^{-}$, $^{12}$C$^{13}$N$^{-}$ and $^{35}$S$^{-}$ secondary ions were collected and measured in parallel at a mass resolution of 8000 that is sufficient to separate $^{13}$C$^{-}$ from the $^{12}$CH$^{-}$ and $^{12}$CH$^{15}$N$^{-}$ from $^{12}$C$^{13}$N$^{-}$. Samples were measured using a 1–2 pA Cs$^+$ primary beam that was rastered over 25 × 25 μm field of a 256 × 256 pixels with a counting time of 5 ms per pixel. Recorded images and data were processed using CAMEGA WinImage software and OpenMIMS plugin$^{42}$ in Image$^{43}$ distribution of Fiji$^{44}$. Different scans of each image were aligned to correct image drift during acquisition. Final images were created by adding the secondary ion counts of each recorded secondary ion from each pixel over all scans. Intracellular carbon and nitrogen uptake from stable isotope-labeled substrates was calculated by drawing regions of interest (ROI) on CN$^{-}$ images (recognizing cells in the images) and calculating $^{12}$C$^{13}$C$^{-}$ and the $^{13}$N$^{-}/^{14}$N ratio (calculated from the $^{12}$C$^{13}$N$^{14}$/^{12}$C$^{14}$N ratio). Instrumental mass fractionation of NanoSIMS analysis was calibrated by the conversion factor.
obtained by comparing carbon and nitrogen isotopic ratios of E. coli cells of varying isotopic enrichments measured at single cells with NanoSIMS and at bulk with an elemental analyzer/isotope ratio mass spectrometer (EA/IRMS, FlashEA 1112/DeltaPlus Advantage, Thermo Fisher Scientific). Concentration of bicarbonate (DIC) in the original sample determined on board was used to calculate substrate incorporation ratio (atom %) for bicarbonate in single cells.

**Biomass and isotope calculations.** Isotope incorporation data analysis and display as violin plots of the kernel density function were done using R with the ggplot246, cowplot47, ggsci48, scales49 packages. The “active” cell ROIs, those incorporated 13C- and/or 15N-labeled substrates, were determined by their isotopic ratio exceeding the 99.7% confidence interval of background carbon and nitrogen isotopic abundance of polycarbonate membranes (1.24 atom % for 13C and 0.446 atom % for 15N). As measures for the rate of microbial biomass synthesis, we calculated the biomass-specific growth rate and the substrate incorporation-based biomass generation rates. The biomass-based specific growth rates ($\mu_b$, Eq. (1)) were calculated from the abundance of cells in the sediment samples at the start of incubation ($X_0$) and the abundance at the incubation period ($t$). The substrate incorporation-based biomass generation rates ($S$, Eqs. (2) and (3), for carbon and nitrogen substrates, respectively) were calculated from the fractional abundance of isotope label in cellular biomass, where $\mu_b$ is the biomass generation rate (encompassing both cell maintenance and generation of new cells), $t$ is the length of the incubation, $F_{\text{lab}}$ is the labeling strength, $F_1$ is the single-cell NanoSIMS measurement, and $F_{\text{nat}}$ is the natural abundance. For the calculation of $\mu_b$ and $S$, a conservative approach was used by only including ROIs with either 13C or 15N ratio above the 99.7% confidence interval of backgrounds shown above. These rate calculations assumed that the carbon and nitrogen for the biomass generation were all derived from the substrates.

$$
\mu_b = \frac{\ln X_t - \ln X_0}{t}
$$

(1)

$$
\mu_c = -\ln \left(1 - \frac{\left(\frac{F_1 - F_{\text{nat}}}{F_{\text{lab}} - F_{\text{nat}}} \right)}{t}\right)
$$

(2)

$$
\mu_N = -\ln \left(1 - \frac{\left(\frac{F_N - F_{\text{nat}}}{F_{\text{lab}} - F_{\text{nat}}} \right)}{t}\right)
$$

(3)

To document the physiological status of microorganisms in the original sediment samples, the fraction of microbes that originally existed in the sediment samples ($f_0$) was calculated from the observed active ROI fractional ratio ($f_i$), a factor of biomass increase ($A$) by following equation (Eqs. (4)–(6)).

$$
X_i = AX_0
$$

(4)

$$
X_i(1 - f_0) = X_0(1 - f_i) \quad \left[f_i < 1, X_0(1 - f_i) < X_0 \right]
$$

(5)

$$
V_0 = 1 - A(1 - f_i)
$$

(6)

If the number of ROIs was not enough to give enough sampling depth for fulfilling the Eq. (5), the $f_0$ could not be calculated and was shown as N.A.

**DNA extraction and sequencing.** Microbial cells on non-coated polycarbonate membrane were lysed using a protocol modified after Morono et al.51 at ISO Class 1 clean air environment in super-clean room of IAMSTEC-KOCHI52. Sorted cells on the membrane were lysed by three-step alkaline treatments to minimize fragmentation of extracted DNA. First, five microliters of denaturation buffer (200 mM KOH, 20 µM EDTA) was added to the tiny piece of membrane with sorted cells, incubated at room temperature for 5 min, followed by adding 5 µL of neutralization buffer (200 mM HCl, 300 mM Tris-HCl [pH 7.5]). The liquid part was recovered for the second step, which was adding another 5 microliters of denaturation buffer, heating at 70°C for 5 min, and adding 5 µL of neutralization buffer, followed by recovery of the liquid part and pooling with the first extract. The third step was adding an additional 5 µL of denaturation buffer, heating at 70°C for 10 min, and adding 5 µL of neutralization buffer, followed by recovery of the liquid part. Heating was done with an Applied Biosystems Veriti Thermal Cycler (Thermo Fisher Scientific). The lysis of cells was confirmed by checking processed membrane with fluorescence microscope after staining with SYBR Green I (Thermo Fisher Scientific). Negative control extractions (8 separate extractions) were performed by using polycarbonate membrane without sorted cells. 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’-TGY CAG CCG CGG TAA-3’, U806R: 5’-GGG GTA CCH GGG TWT GAT AT-3’). Also, PCR No-Tempiler Control reactions (PCR-NTCs) run without template DNA (6 separate reactions) were performed. First-round PCR amplicons were quantified and purified via gel electrophoresis and then indexed and barcoded in the second PCR. The DNA sequencing was performed on a MiSeq Illumina platform following the methods outlined by Hoshino and Inagaki53. The obtained sequence reads were quality trimmed and filtered by FASTQ Toolkit in Basepass (Illumina). Resultant paired-end reads were processed using mothur54 following the Mothur software package (v.1.35.0). Of Illumina MiSeq Standard Operating Procedure,55 All sequence reads from negative control extractions and PCR-NTCs, along with sequence reads from all samples were used to generate operational taxonomic units (OTUs) at 97% similarity using usearch 10.0 program.56 To remove sequences that may be of exogenous origin57, the most conservative way to remove contaminant sequences considers all taxa that are identified in the laboratory controls as contaminants. This approach has the advantage of eliminating any false-positive discoveries of novel taxa; however, it has the disadvantage of introducing false-negative results, which may be of interest. For example, microbial species that are ubiquitous in environments may be excluded from a sample. To circumvent this issue, we employed a “probabilistic” approach to examining the microbial composition based on the consistency (i.e., variance) of the relative distribution of the taxonomic assignments or OTUs across the entire set of samples. This approach compares the proportion of the OTUs present between negative controls (i.e., extracted negative controls or PCR-NTCs) and samples, and determines the significantly more abundant OTUs in negative controls (ANOVa analysis) to be identified as contaminants and excluded from the sequence assemblage. By accounting for the consistency (variance) at which any taxon or OTU is found in the negative controls or set of experimental samples, a more nuanced view of the overall data set can be achieved. For the ANOVA analysis, we used Qime58 script group_significance.py to estimate a likelihood based on observed abundances for conditions (negative controls and experimental samples). This process was conducted sequentially on extraction negative controls and PCR-NTCs, and the OTUs that have significantly higher ($p < 0.05$, ANOVA) abundance in the extraction negative controls and the NTCS than in the incubation samples were identified and removed from further analysis. This read set was taxonomically classified by using the mothur utility package with the SSURef_NR99_123_SILVA database59,60. The total sequences and relative abundance of taxa in the final sequence dataset are provided in Supplementary Data Set 2. The relative abundance of bacterial and archaeal OTUs at the genus level were exposed to non-metric multidimensional scaling with the metaMDS function of R software package vegan61. A Bray–Curtis distance matrix was used as dissimilarity index.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability.** DNA sequence data are deposited in the DNA Data Bank of Japan (DDBJ) under the accession code of DRA007733. The authors declare that the data supporting the findings of this study are available within the paper and the Supplementary Information.

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37. Author contributions

Y.M., F.I., and S.D. designed the study; S.D. and F.I. co-led IODP Expedition 329; Y.M. and F.I. carried out incubation of sediment samples with stable isotope substrates; Y.M. and T.T. prepared the samples for NanoSIMS analysis; M.I. and Y.M. conducted the NanoSIMS analyses on single cells; Y.M. and T.T. extracted the DNA; Y.M., T.A., and T.H. sequenced the DNA and analyzed the sequencing data; Y.M., S.D. and F.I. were the primary authors of the manuscript with input from all other co-authors.

38. Competing interests

The authors declare no competing interests.
