Statistical Analysis of Community RNA Transcripts between Organic Carbon and Geogas-Fed Continental Deep Biosphere Groundwaters

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ABSTRACT Life in water-filled bedrock fissures in the continental deep biosphere is broadly constrained by energy and nutrient availability. Although these communities are alive, robust studies comparing active populations and metabolic processes across deep aquifers are lacking. This study analyzed three oligotrophic Fennoscandian Shield groundwaters, two “modern marine” waters that are replenished with organic carbon from the Baltic Sea and are likely less than 20 years old (171.3 and 415.4 m below sea level) and an extremely oligotrophic “thoroughly mixed” water (448.8 m below sea level) of unknown age that is composed of very old saline and marine waters. Cells were captured either using a sampling device that rapidly fixed RNA under in situ conditions or by filtering flowing groundwater over an extended period before fixation. Comparison of metatranscriptomes between the methods showed statistically similar transcript profiles for the respective water types, and they were analyzed as biological replicates. Study of the small subunit (SSU) rRNA confirmed active populations from all three domains of life, with many potentially novel unclassified populations present. Statistically supported differences between communities included heterotrophic sulfate-reducing bacteria in the modern marine water at 171.3 m below sea level that has a higher organic carbon content than do largely autotrophic populations in the H2- and CO2-fed thoroughly mixed water. While this modern marine water had signatures of methanogenesis, syntrophic populations were predominantly in the thoroughly mixed water. The study provides a first statistical evaluation of differences in the active microbial communities in groundwaters differentially fed by organic carbon or “geogases.”

IMPORTANCE Despite being separated from the photosynthesis-driven surface by both distance and time, the deep biosphere is an important driver for the earth’s carbon and energy cycles. However, due to the difficulties in gaining access and low cell numbers, robust statistical omics studies have not been carried out, and this limits the conclusions that can be drawn. This study benchmarks the use of two separate sampling systems and demonstrates that they provide statistically similar RNA transcript profiles, importantly validating several previously published studies. The generated data are analyzed to identify statistically valid differences in active microbial community members and metabolic processes. The results highlight contrasting taxa and growth strategies in the modern marine waters that are influenced by recent infiltration of Baltic Sea water versus the hydrogen- and carbon dioxide-fed, extremely oligotrophic, thoroughly mixed water.

KEYWORDS deep biosphere, groundwaters, metatranscriptomes, protein-coding RNA, rRNA

Citation Lopez-Fernandez M, Broman E, Simone D, Bertilsson S, Dopson M. 2019. Statistical analysis of community RNA transcripts between organic carbon and geogas-fed continental deep biosphere groundwaters. mBio 10:e01470-19. https://doi.org/10.1128/mBio.01470-19.

Editor Mark J. Bailey, CEH-Oxford
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Received 6 June 2019
Accepted 9 July 2019
Published 13 August 2019
The deep biosphere is the largest biome on earth, where the continental subsurface alone hosts up to $6 \times 10^{29}$ cells from all three domains (1). Deep life has been demonstrated as active by, e.g., “viable/dead” PCR amplification (2), “omics” (3–5), and video evidence (6). A previous study at the Swedish Nuclear Fuel and Waste Management Company (SKB)-operated Åspö Hard Rock Laboratory (Åspö HRL) used a specially designed sampling device to fix cells under in situ conditions to ensure that RNA transcripts were unaffected by sampling procedures (3). In contrast, other studies used cell capture from flowing groundwater on filters over several days prior to fixation (see, e.g., reference 4). However, it is unknown if extended capture times alter the RNA transcript profile.

The extreme oligotrophy in the continental deep biosphere can limit cell numbers to $10^1$ to $10^7$ cells/ml (1), while Åspö HRL groundwaters contain $10^5$ to $10^6$ cells/ml (7). Due to the difficulty of obtaining deep biosphere samples and the large water volume needed to extract sufficient RNA for sequencing, no omics studies have provided sufficient replicates for valid statistics.

In this study, we combined RNA transcript data from the sampling device (3) and from cells captured over several days on filter holders to evaluate if the two methods are comparable (see File S1 in the supplemental material). Additionally, we statistically analyzed gene transcript counts pertaining to active microbial taxa and their metabolic processes between groundwaters of various ages and origins.

The studied groundwaters were two modern marine waters (MM-171.3 and MM-415.2) that are replenished from the Baltic Sea and have a residence time of $\sim 20$ years and a “thoroughly mixed” water (TM-448.4) that is composed of different waters of multiple origins and unknown age (3, 7, 8). Cells were captured, and community RNA was extracted and sequenced according to File S1. The small subunit (SSU) rRNA sequences (File S2) were annotated against the SILVA database and normalized as relative abundances (File S3). The MM-415.2 filter holder metatranscriptomes only had two replicates and thus cannot be statistically compared to the others. However, this groundwater was clearly different in both its SSU and protein-coding RNA (pcRNA) transcripts (Fig. 1) and is discussed in File S4. Nonmetric multidimensional scaling (NMDS) of SSU rRNA transcript beta diversity suggested that the three water samples were statistically different in their microbial communities (permutational multivariate analysis of variance [PERMANOVA] 9,999 permutations, $P = 0.0011$; Fig. 1). Previous analysis of the sampling device (SD) TM-448.4-4 sample showed it was different from the SD TM-448.4-3 sample, as it had likely been recently exposed to an electron donor (3). Repetition of the NMDS without this outlier altered the significance between the three groundwaters ($P = 0.004$). Without TM-448.4-4, the grouping supports the notion that (i) the two methods give highly similar RNA transcript patterns and, therefore, sampling with filter holders over several days is valid, and (ii) in the absence of periodic availability of an electron donor (as for the SD TM-448.4-4 sample [3]), the deep biosphere communities were stable for a minimum of 2 years.

SSU rRNA-based phylogeny from all analyzed metatranscriptomes showed that a broad range of phyla from all three domains of life were viable and had protein-synthesizing potential (3) (Fig. 1). It also reinforced that the deep biosphere contains a large relative proportion of active candidate phyla from all three domains (e.g., Patescibacteria) along with many unclassified sequences. Statistically valid differences between the MM-171.3 and TM-448.4 groundwaters included sulfate-reducing bacteria (SRB) with Desulfobulbaceae in the MM-171.3 groundwater compared to Desulfobacteraceae and Desulfurivibrio in the TM-448.4 groundwater (File S5). This confirms that sulfur compound reduction is prevalent (see, e.g., references 9 and 10) with the predominantly organoheterotrophic SRB Desulfobulbaceae (11) in the MM-171.3 groundwater compared to autotrophic Desulfurivibrio spp. (12) in the ultraoligotrophic TM-448.4 water. In addition, increased 16S rRNA gene transcripts in the TM-448.4 groundwater that aligned within the Syntrophus genus demonstrated that syntrophy is likely to be an important survival strategy in these oligotrophic groundwaters (13).

Analysis of pcRNA transcripts identified 973 unique prokaryote genes (File S6). The NMDS analysis also showed that the community-level transcription profiles were...
FIG 1 (A and B) Taxonomic annotation of the SSU rRNA (A) and protein-coding RNA (B) sequences showing stacked bars of the taxonomic phyla and Proteobacteria classes (β-proteobacteria shown separately) with a relative abundance of >0.1% for the modern marine (MM-171.3 and MM-415.2) and thoroughly mixed (TM-448.4) groundwaters. Rare taxa with a relative abundance of <0.1% are given as "other phyla." (C and D) NMDS Bray-Curtis dissimilarity (beta diversity) plots based on the SSU rRNA at the lowest taxonomic level that could be assigned to the SILVA database using the Ribosomal Database Project classifier (C) and a second NMDS without the SD TM-448.4 outlier (D). (E and F) NMDS Bray-Curtis plots based on the annotated transcripts (i.e., UniProtKB identifiers) with an E value of <10 and TPM of >100 from the full data set (E) and without the SD TM-448.4 outlier (F). The sampling methods are filter holders (FH) and sampling device (SD). Cand, Candidatus.
statistically different ($P = 0.002$; Fig. 1), and further removal of the SD TM-448.4-4 outlier gave a $P$ value of 0.004. Altogether, 410 prokaryotic genes had significant differential expression between the MM-171.3 and TM-448.4 groundwaters (false-discovery rate [FDR] $< 0.05$; E value $< 0.001$). Transcripts encoding tricarboxylic acid (TCA) cycle ($mdh$, $fumC$, and $sucC$) and ATP synthase ($atpAG$) proteins had higher transcript counts in the MM-171.3 groundwater, while increased TM-448.4 transcripts encoded, e.g., ribosomal (e.g., $rpmB$, $rpsB$, and $rplC$) and stress/repair (e.g., $dfx$, $recGN$, $cspAB$, $clpPX$, $dnaK$, and $hspC4$) proteins. Additionally, a qualitative comparison of the SD TM-448.4-4 outlier (3) with the other three replicates suggested that this outlier had more transcripts involved with, e.g., replication and metabolic processes. Overall, most overexpressed transcripts were seen in the MM-171.3 groundwater, robustly demonstrating that this community was actively growing while the TM-448.4 populations were in “metabolic standby” (3).

The metabolic process with the greatest number of statistically different MM-171.3 groundwater transcripts was methanogenesis from CO$_2$ ($fwdC$, $mtrACDEH$, $mcrABCG$; right) (A), as well as protein-coding RNA transcripts assigned to sulfate-reducing bacteria taxa (left) and genes attributed to sulfate reduction (cytochrome $c_3$, $rd2$, $aprA$, $dsiA$, and $dsiAB$; right) (B).

**FIG 2** (A and B) Average of the significantly different (false-discovery rate [FDR] $< 0.05$; E value $< 0.001$) transcripts per million sequences (TPM) for the modern marine (MM-171.3) and thoroughly mixed (TM-448.4) groundwaters for protein-coding RNA transcripts assigned to methanogenic taxa (left) and genes attributed to methanogenesis from CO$_2$ ($fwdC$, $mtrACDEH$, $mcrABCG$; right) (A), as well as protein-coding RNA transcripts assigned to sulfate-reducing bacteria taxa (left) and genes attributed to sulfate reduction (cytochrome $c_3$, $rd2$, $aprA$, $dsiA$, and $dsiAB$; right) (B).
dissimilatory sulfate-reducing genes **aprA**, **dsrA**, and **dsvAB** attributed to *Desulfovibrio* spp. were increased in the TM-448.4 groundwater (Fig. 2 and File S8). The importance of syntrophy was also further demonstrated by pcRNA transcripts in both the MM-171.3 and TM-448.4 waters attributed to *Syntrophus aciditrophicus* that grows alongside H₂ utilizers (17) predominantly present in the TM-448.4 groundwater (File S9). Finally, earlier observations of cyanobacteria in ancient deep terrestrial groundwaters (18, 19) were confirmed by increased *Synechocystis* pcRNA transcripts in the TM-448.4 water, also demonstrating their viability in these habitats.

This work presents for the first time a statistically robust omics study of deep subsurface crystalline rock groundwaters with different depths and geochemical characteristics. We conclude that cell capture over several days does not alter RNA transcript profiles compared to rapid *in situ* fixation in this extremely oligotrophic environment. Importantly, this analysis of the two methods validates published studies that have used capture times prior to RNA fixation over the several days needed to obtain sufficient biomass for biomolecule extraction from low-cell-density deep groundwaters. The similarity of the data obtained by the two methods was likely due to the long-term and stable oligotrophic conditions in the respective groundwaters. These novel findings also provide evidence on how the differences in active communities and metabolic processes are influenced by organic carbon versus geogas-fed modern marine and thoroughly mixed groundwaters, respectively. This benchmarking of deep biosphere metatranscriptome analyses paves the way for future and still-needed exploration of the living deep biosphere in a statistically sound way.

**Data availability.** The raw sequence data are available in the NCBI Sequence Read Archive BioProject numbers PRJNA400688 and PRJNA541524 for the sampling device and filter holders, respectively.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.01470-19.

- **FILE S1**, DOCX file, 0.1 MB.
- **FILE S2**, XLSX file, 0.1 MB.
- **FILE S3**, XLSX file, 0.2 MB.
- **FILE S4**, DOCX file, 0.1 MB.
- **FILE S5**, XLSX file, 0.1 MB.
- **FILE S6**, XLSX file, 0.4 MB.
- **FILE S7**, EPS file, 0.5 MB.
- **FILE S8**, XLSX file, 0.1 MB.
- **FILE S9**, EPS file, 1.9 MB.

**ACKNOWLEDGMENTS**

The Swedish Research Council (contracts 2018-04311, 2017-04422, and 2014-4398) and The Swedish Nuclear Fuel and Waste Management Company (SKB) supported the study. M.D. thanks the Crafoord Foundation (contracts 20180599 and 20130557), the Nova Center for University Studies, Research and Development, and Familjen Hellmans Stiftelse for financial support. M.D. and D.S. thank the Carl Tryggers Foundation (grant KF16:18) for financial support. S.B. acknowledges financial support from the Swedish Research Council and Science for Life Laboratory.

High-throughput sequencing was carried out at the National Genomics Infrastructure hosted by the Science for Life Laboratory. Bioinformatics analyses were carried out utilizing the Uppsala Multidisciplinary Center for Advanced Computational Science (UPPMAX) at Uppsala University (project b2013127).

M.L.-F., S.B., and M.D. conceived the study. M.L.-F. carried out the sampling and prepared nucleic acids for sequencing. E.B. and D.S. carried out bioinformatic analysis. M.L.-F., S.B., and M.D. interpreted data. M.L.-F. and M.D. drafted the manuscript that was approved by all authors.

We declare no conflicts of interest.
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