Lawrence Berkeley National Laboratory

LBL Publications

Title

Novel candidate taxa contribute to key metabolic processes in Fennoscandian Shield deep groundwaters.

Permalink https://escholarship.org/uc/item/3d39s99d

Journal ISME Communications, 4(1)

Authors

Dopson, Mark Rezaei Somee, Maryam González-Rosales, Carolina <u>et al.</u>

Publication Date 2024

DOI

10.1093/ismeco/ycae113

Peer reviewed



https://doi.org/10.1093/ismeco/ycae113 Advance access publication: 23 September 2024 Original Article

Novel candidate taxa contribute to key metabolic processes in Fennoscandian Shield deep groundwaters

Mark Dopson^{1,*}, Maryam Rezaei Somee¹, Carolina González-Rosales¹, Lauren M. Lui², Stephanie Turner^{1,3}, Moritz Buck⁴,

Emelie Nilsson¹, George Westmeijer¹, Kamal Ashoor¹, Torben N. Nielsen², Maliheh Mehrshad⁴, Stefan Bertilsson⁴

¹Centre for Ecology and Evolution in Microbial Model Systems (EEMiS), Linnaeus University, 39231 Kalmar, Sweden

²Molecular Ecosystems Biology Department, Environmental Genomics and Systems Biology Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, United States

³Department of Forest Mycology and Plant Pathology, Swedish University of Agricultural Sciences, P.O. Box 7050, 75005 Uppsala, Sweden

⁴Department of Aquatic Sciences and Assessment, Swedish University of Agricultural Sciences, P.O. Box 7050, 75005 Uppsala, Sweden

*Corresponding author: Mark Dopson, Department of Biology and Environmental Sciences, Stuvaregatan 4, 39231 Kalmar, Sweden. Email: mark.dopson@lnu.se

Abstract

The continental deep biosphere contains a vast reservoir of microorganisms, although a large proportion of its diversity remains both uncultured and undescribed. In this study, the metabolic potential (metagenomes) and activity (metatranscriptomes) of the microbial communities in Fennoscandian Shield deep subsurface groundwaters were characterized with a focus on novel taxa. DNA sequencing generated 1270 de-replicated metagenome-assembled genomes and single-amplified genomes, containing 7 novel classes, 34 orders, and 72 families. The majority of novel taxa were affiliated with *Patescibacteria*, whereas among novel archaea taxa, *Thermoproteota* and *Nanoarchaeota* representatives dominated. Metatranscriptomes revealed that 30 of the 112 novel taxa at the class, order, and family levels were active in at least one investigated groundwater sample, implying that novel taxa represent a partially active but hitherto uncharacterized deep biosphere component. The novel taxa genomes coded for carbon fixation predominantly via the Wood–Ljungdahl pathway, nitrogen fixation, sulfur plus hydrogen oxidation, and fermentative pathways, including acetogenesis. These metabolic processes contributed significantly to the total community's capacity, with up to 9.9% of fermentation, 6.4% of the Wood–Ljungdahl pathway, 6.8% of sulfur plus 8.6% of hydrogen oxidation, and energy conservation via nitrate (4.4%) and sulfate (6.0%) reduction. Key novel taxa included the UBA9089 phylum, with representatives having a prominent role in carbon fixation, nitrate and sulfate reduction, and organic electron donor oxidation. These data provided insights into deep biosphere microbial diversity and their contribution to nutrient and energy cycling in this ecosystem.

Keywords: deep biosphere, metagenomics, metatranscriptomics, Candidatus

Introduction

The deep biosphere is defined as the life below the Earth's surface and is estimated to host 12%–20% of the total biomass on Earth [1, 2]. These microorganisms (along with viruses) are active and viable in groundwaters down to several kilometers (e.g. 2.8 km in the Mponeng mine, South Africa) below the Earth's surface [1, 3–9]. Due to the prevailing low nutrient and energy availability, water-bearing bedrock fractures in the terrestrial deep biosphere typically feature low cell densities [10]. The understanding of diversity and metabolic capabilities of the vast deep biosphere microbiome remains constrained by barriers to study them. Access points to observe and sample these microbiomes are limited to boreholes, mines, or subsurface laboratories, making the deep biosphere one of the least understood biomes on Earth.

The "great plate count anomaly" revealed that the majority of the Earth's microbial diversity remains uncultured [11]. The use of nucleic acid sequencing has aided investigation of this uncultured majority [12], e.g. community 16S ribosomal RNA gene sequences revealing that all major ecosystems except the human body are dominated by uncultured species [13]. In addition, assembled genomes from community DNA [14–16] and single-cell amplified genomes [12] provide an updated view of the tree of life. Omics studies have been applied to shallow groundwaters [17], geysers [18], mines [19], and deep groundwaters of the Fennoscandian Shield [10, 20, 21], revealing an impressive diversity of candidate and unknown taxa at different ranks. As many microbial species are not yet axenically cultured under laboratory conditions, efforts have been made to standardize microbial classification using genomic data rather than an axenic culture as type [22, 23]. Few validly described species have been cultured from Fennoscandian Shield deep groundwaters [24-26], whereas largescale omics analysis has recovered more than 568 genome species (operational definition of species with \geq 95% average genome nucleotide identity [20]). Additionally, widespread genome features of microbes inhabiting this ecosystem highlight the importance of biological interactions in sustaining many taxa [20], a factor that further decreases the possibility to obtain axenic cultures for taxa dependent on obligate interactions and metabolic dependencies. This is especially relevant for representatives of DPANN and Patescibacteria (previously defined as candidate phyla

© The Author(s) 2024. Published by Oxford University Press on behalf of the International Society for Microbial Ecology.

Received: 8 July 2024. Revised: 6 September 2024. Accepted: 20 September 2024

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

radiation bacteria) that encompass 30% of the recovered genome species in deep Fennoscandian Shield groundwaters [20].

The typically extreme conditions in the terrestrial deep biosphere, such as low energy and nutrient availability, select for a mixture of long-term indigenous taxa. In parallel, there are infiltrating microbes from the surface, including those that are able to adapt to this environment, such as Patescibacteria, Pseudomonadota (formerly Proteobacteria), and Epsilonbacteraeota [27], while non-viable cells appear to be rapidly degraded [28]. Compared to surface biomes that have rapid fluxes in typically light, temperature, and nutrient availability, geochemical conditions in the deep biosphere are recorded to be relatively stable for at least 15 years [21], and the energy available is generally low compared to principally photosyntheticallydriven surface biomes [29]. Despite the isolated and stable conditions, dynamics in the microbial community composition can occur over multiple years with the most dynamic changes evidenced in shallow compared to deeper groundwaters [30]. Carbon and energy sources in Fennoscandian Shield oligotrophic deep groundwaters are suggested to be via infiltration of autochthonous surface species that subsequently die (termed necromass [28]), geogenic hydrogen and carbon dioxide [31], and surface-derived recalcitrant dissolved organic matter [32]. Analyses of deep subsurface groundwaters in the Fennoscandian Shield have revealed a shared "core" microbial community suggested to have adaptations to the low carbon and energy conditions of this ecosystem [20]. These adaptations (reviewed in [33]) include a small cell size with the potential to pass through a 0.2 μ m pore size membrane typically used for cell capture [34], streamlined genomes calling for metabolite exchange and symbiotic interactions [35], and an "episodic lifestyle" in response to intermittent availability of nutrients and energy sources [20]. Despite these previous studies, further inquiries into the diversity of subsurface groundwaters in the Fennoscandian Shield continue to reveal many novel taxa for which the biological roles remain to be elucidated

While the deep biosphere is the largest biome on Earth, the vast majority of its microbial diversity remains undescribed, and consequently, their contribution to ecosystem function remains overlooked. To address this knowledge gap, this study investigated novel microbial candidate taxa from deep subsurface groundwaters in the Äspö Hard Rock Laboratory (Äspö HRL; Sweden) and Olkiluoto Island (Finland) using a combined long- and short-read metagenomic and short-read metatranscriptomic approach. The questions addressed include elucidating the metabolic potential of these novel deep subsurface populations and comprehensive mapping of their metabolic role in the community.

Materials and methods Sampling sites

Metagenomic and metatranscriptomic datasets (Supplemental Table S1 and Fig. 1) were generated from the microbial community sampled from the Äspö HRL underground facility managed by the Swedish Nuclear Fuel and Waste Management Company (SKB). Äspö HRL consists of a 3.6 km tunnel extending to ~460 m below sea level (mbsl; Lat N 57° 26′ 4″, Lon E 16° 39′ 36″). Further metagenomic datasets were generated from Olkiluoto Island on the Finnish Baltic Sea coast (Lat N 61° 14′ 31″, Lon E 21° 29′ 23″) with boreholes to depths greater than 500 mbsl operated by Posiva Oy. The Äspö HRL tunnel provides access to investigate deep microbial life in 1.8 Ga old Proterozoic crystalline bedrock of the Fennoscandian Shield (also termed

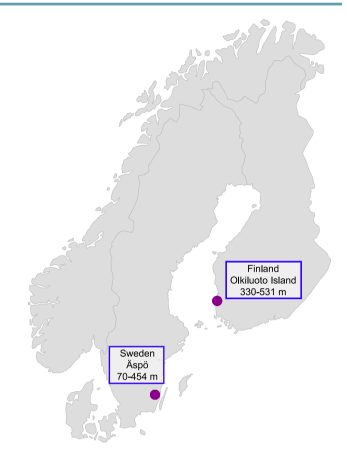


Figure 1. Map of the sampling sites at the Äspö HRL and Olkiluoto Island. Figure generated in the ggplot2, maps, and sf packages in R.

the Baltic Shield; area of the Earth's crust encompassing large parts of Norway, Sweden, Finland, and Russia [31]), with the Äspö HRL geology, hydrology, and chemistry having been previously reported [36–38]. Olkiluoto Island drillholes and tunnels penetrate Precambrian metamorphic rocks with igneous rocks as previously described [39], where the drawdown of waters can result in some mixing, but there was no evidence of oxygen infiltration.

Cell capture from groundwaters

A portion of the community nucleic acids from planktonic and biofilm cells captured from the various groundwaters included in this investigation were previously published [20, 21, 40-42]. Further metagenomes and metatranscriptomes were generated for this study according to the previously described methods (Table 1) plus for nanopore sequencing in Supplementary File S1. Briefly, planktonic microbial cell fractions for metagenomes were captured by flushing stagnant water from the borehole to ensure a representative sample was obtained, and then connecting a high-pressure holder containing a 0.1 μ m filter and allowing water to pass through the filter under in situ temperature and pressure. The filters were then aseptically collected before flash freezing in liquid nitrogen and stored at -80°C until processing. In addition, biofilm cells were collected in flow cells containing solid support directly attached to the boreholes, where the groundwater was allowed to flow under in situ temperature and pressure before opening the vessels for cell recovery [40]. Finally, cells for community RNA transcripts were captured on 0.1 μ m filters under in situ temperature and pressure, followed by directly flash freezing in liquid nitrogen and storage at -80°C [41]. The

Table 1. Details of the nucleic acid sequencing used in this study and not published in the previous iteration of the FSG [20].

Designation	Borehole	Nucleic acid	Platform	Depth (mbsl)	Metagenome/-transcriptome accession numbers (NCBI)
AHRL-69.4	KR0015B	DNA	Illumina NovaSeq	69.4	SRR26275512
			Oxford Nanopore		SRR27925880
		RNA	Illumina NextSeq		SRR26275508
AHRL-200.6	SA1420A	DNA	Illumina NovaSeq	200.6	SRR26275511
		RNA	Illumina NextSeq		SRR26275507
AHRL-281.6	SA2074A	DNA	Illumina NovaSeq	2817	SRR26275510
AHRL-345.0	SA2600A	DNA	Illumina NovaSeq	345.0	SRR26275509
		RNA	Illumina NextSeq		SRR26275506

DNA/RNA extraction method for each sample and sequencing platform is provided in Supplementary Table S1.

Bioinformatic analyses

Deep groundwater metagenomes (57 samples) were captured from the Äspö HRL (KR0015B, KF0069A01, SA1229A, SA1420A, SA2074A, KA2198, SA2600A, KA3105-4, and KA3385A) and Olkiluoto Island (KR_11, KR_13, and KR_46) boreholes (metadata and a map of the Äspö HRL boreholes are previously published [10, 20, 21]) with details given in Table 1, Supplementary File S1, and Supplementary Table S1. Detailed bioinformatic methods are provided in Supplementary File S1. Briefly, Fennoscandian Shield genome (FSG) metagenomic datasets were quality-checked and trimmed by Trimmomatic [43] and then assembled using MEGAHIT [44]. Contigs ≥ 2 kb were binned using MetaBAT2 [45], genome quality assessed via CheckM [46], and taxonomy assigned using GTDB-tk [47]. Single-cell amplified genomes were assembled and quality checked as previously explained [20]. MAGs/SAGs were de-replicated based on similarity thresholds using fastANI by mOTUlizer [48]. This resulted in 112 representative MAGs/SAGs that were denoted as unclassified by GTDB-tk at the family level and above that were selected for further analysis and were assigned unique codes based on their origin with corresponding accession numbers given in Supplementary Table S2. Recovered MAGs/SAGs were metabolically annotated via the METABOLIC tool in community and genome-level modes [49]. The abundance of 112 candidate MAGs/SAGs in each trimmed metagenome and metatranscriptome samples was calculated by CoverM (v.0.6.1) using TPM as the normalization method, with further details available in Supplementary File S1.

Results and discussion

The present study contributed new long- and short-read metagenome (13 samples from 4 boreholes) and short-read metatranscriptome (9 samples from 3 boreholes) sequence data from planktonic cells in Äspö HRL deep subsurface groundwaters (Table 1). This resulted in 1.15 TB of metagenomic data that were combined with previously published sequencing results [20] to enable reconstruction of 2295 MAGs and SAGs with \geq 50% completeness and \leq 5% contamination. These MAGs/SAGs were de-replicated into 1270 representative assembled genomes, from which novel taxa identified according to taxonomic assignment of the genome taxonomy database (GTDB) were selected for further analyses.

Number and abundance of novel FSG taxa

While the updated FSG did not contain any novel phyla, there were 12 de-replicated representative bacterial MAGs/SAGs assigned to seven novel candidate classes, 46 representative MAGs/SAGs in 34 novel candidate orders, 98 MAGs/SAGs in 72 novel candidate families, 335 MAGs/SAGs in 230 novel candidate genera, and 626 MAGs/SAGs in 463 novel candidate species (Fig. 2 and Supplementary Table S2). Analogously, the genomes contained novel archaea, with four de-replicated representative archaeal MAGs assigned to separate candidate orders, 14 MAGs in 13 candidate families, 95 MAGs in 65 candidate genera, and 168 MAGs in 115 candidate species (Fig. 3 and Supplementary Table S2). The high number of previously undescribed taxa in the FSG were in line with the existing view of the deep biosphere as an untapped reservoir of biological novelty [17]. The FSG encompasses groundwaters of differing depths, ages, and sources from two sites on either side of the Baltic Sea, which exhibit a greater microbial diversity as compared to other Fennoscandian Shield [50], North American [51], and South African [52] groundwaters greatly dominated by a single taxa. However, the factors controlling the large differences in worldwide deep groundwater diversities have not been elucidated.

The bacterial phyla with the highest number of novel MAGs/SAGs at the levels of class, order, and family was Patescibacteria with 27 representatives, followed by Planctomycetota (8 MAGs/SAGs), Omnitrophota (7 MAGs/SAGs), Margulisbacteria plus Elusimicrobiota (both 5 MAGs/SAGs), and Desulfobacterota (4 MAGs/SAGs). Patescibacteria comprise a large percentage of the bacterial phylogenetic diversity [14, 53] and have been extensively identified in near-surface and deep groundwaters [14, 17]. The Patescibacteria are suggested to infiltrate groundwaters due to their high tendency of being exported from deep soil horizons along with streamlined genome features that support a simple fermentative lifestyle that enable them to survive in low carbon and energy conditions [54]. The Omnitrophota have been identified in the deep terrestrial biosphere at the DeMMO site, South Dakota [55]. This taxa is also adapted to the deep biosphere by having a small cell size and reduced genomes, even if various lineages code for assorted respiration processes for energy conservation (e.g. dissimilatory nitrite reduction plus sulfur and iron oxidation [55]) and gene clusters indicative of symbiotic lifestyles [56]. The Margulisbacteria were also identified from shallow suboxic/anoxic groundwater, where the majority of the community was suggested to lack genes coding for complete metabolic pathways [17]. The Elusimicrobiota were originally defined as endosymbionts [57], while subsequently described free-living groundwater populations are metabolically diverse [58]. Finally, the Desulfobacterota are known to populate anaerobic groundwaters [59]. The novel archaeal taxa were affiliated to Thermoproteota (5 MAGs/SAGs) and Nanoarchaeota (2 MAGs/SAGs). The recently described Candidatus Methanodesulfokores washburnensis from the Thermoproteota has been identified in the Pennsylvania, USA, deep subsurface [60]



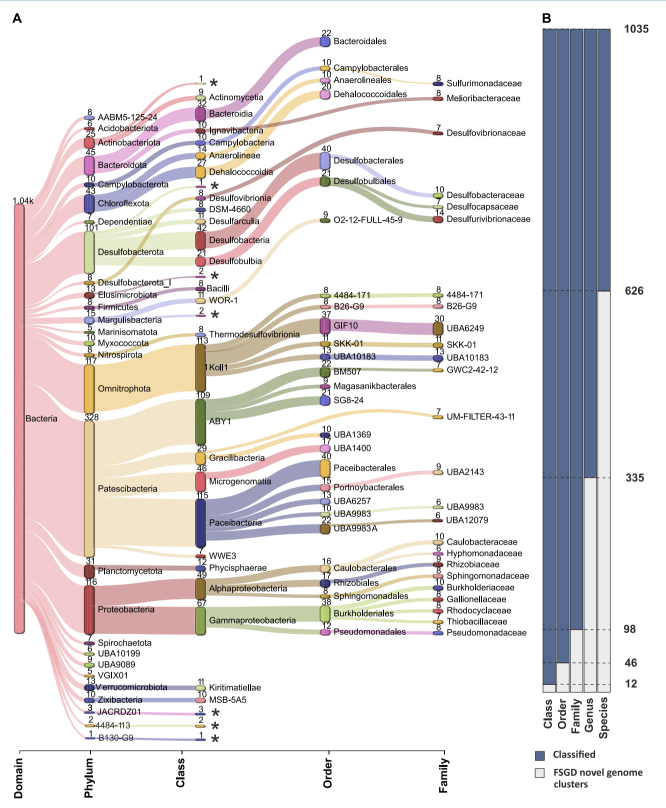


Figure 2. Taxonomic profile information of FSG bacterial metagenome assembled genomes (MAGs) and single amplified genomes (SAGs). (A) Distribution of MAGs/SAGs in different phylogenetic levels, with the top 25 taxa at each level. Each number indicates the number of cluster MAGs/SAGs in that taxonomic level. Novel candidate classes marked with an asterisk. (B) Classified/unclassified number of MAGs/SAGs by GTDB-tk in each taxonomic level.

and is capable of both methanogenesis and sulfate reduction that makes it potentially possible to perform anaerobic oxidation of methane within cellular boundaries [61]. Based upon a single transcript being represented for the ORFs in each of the MAGs/SAGs with a TPM value >100, 89 out of the 112 novel taxa were considered to be active in at least one groundwater. Furthermore, when plotting the same metatranscriptomes as transcripts per million (TPMs), the low transcript numbers were

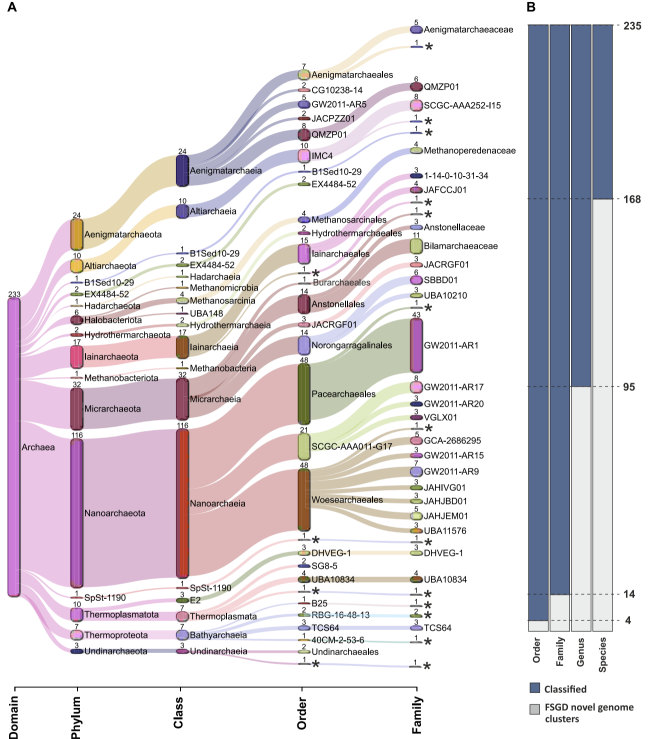


Figure 3. Taxonomic profile information of FSG archaeal metagenome assembled genomes (MAGs) and single amplified genomes (SAGs). (A) Distribution of MAGs/SAGs in different phylogenetic levels, with the top 25 taxa at each level. Each number indicates the number of cluster MAGs/SAGs in that taxonomic level. Novel candidate classes marked with an asterisk. (B) Classified/unclassified number of MAGs/SAGs by GTDB-tk in each taxonomic level

rounded to zero, such that 30 out of the 112 novel taxa were also considered as active (Fig. 4 and Supplementary Table S3). This supports previous studies in the Fennoscandian Shield showing a broad active community in Äspö HRL [41, 42, 62] and South African [5] groundwaters.

Clustering of the novel classes, orders, and families based on relative abundances and incidence (rather than phylogeny)

in selected Äspö HRL and Olkiluoto Island sites revealed four representative MAGs/SAGs (cluster I) almost ubiquitously present across the investigated Fennoscandian Shield groundwaters (Fig. 4 and Supplementary Table S3). This cluster included two novel orders (AHRL-171.3-O3 and AHRL-448.4-O2) and a novel family (AHRL-448.4-F2) from the UBA9089 phylum that have solely been identified from groundwaters [20, 63] plus a novel

Α



Figure 4. Mapping metagenomic (A) and metatranscriptomic (B) reads against novel FSG taxa at the levels of class, order, and family. (A) Average of the abundance (log₁₀ of TPM) of 112 novel taxa in groundwater samples from Äspö HRL (AHRL) and Olkiluoto Island (OI). OI-528.7 (borehole OL-KR46) was not included as no novel taxa at family or above was detected in this groundwater. (B) Average of the RNA transcript abundance (log₁₀ of TPM) of 112 novel taxa in selected Äspö HRL groundwaters (no data for OI boreholes were available).

Desulfobacterales family (AHRL-415.2-F1) [64]. The three UBA9089 phylum MAGs/SAGs were also highly represented in the Äspö HRL metatranscriptomes, particularly in the AHRL_345.0 groundwater. In contrast, AHRL-415.2-F1 RNA transcripts were only present in AHRL_200.6.

Clusters II and III (four and three MAGs/SAGs, respectively) were also mostly distributed across the investigated groundwaters with particularly high representation in the shallowest Olkiluoto Island groundwater for cluster II. Cluster II MAGs/SAGs included the novel order OI-330.5-O1 of the UBA10199 phylum suggested to have streamlined genomes [65], the AHRL-345.0-F1 family from the RPQK01 order in the *Acidobacteriota*, and AHRL-345.0-O1 attributed to the heterotrophic and obligatory anaerobic *Anaerolineae* (Chloroflexota phylum) [66]. Cluster III MAGs/SAGs included AHRL-200.6-F18 from the *Portnoybacterales* (*Patescibacteria*) that was also active in the AHRL_200.6 groundwater.

All novel FSG MAGs/SAGs from cluster IV were solely identified in the AHRL.69.4 groundwater except for AHRL-69.4-O12 within the UBA6262 phylum that was also identified in the OI.366.7 groundwater. These included AHRL-69.4-O5 from the ABY1 class (Patescibacteria) reported to contain ultrasmall representatives that actively replicate in groundwaters [67]. Members of the Patescibacteria are enriched in aquifer sediments, where they largely exhibit a surface-attached lifestyle. This preference for surface attachment is attributed to higher environmental stability, greater access to nutrients, and proximity of putative symbiotic partners as compared to being freeliving and planktonic [67]. Nevertheless, previous work has demonstrated that the Fennoscandian Shield early-stage biofilm formation can be attributed to autotrophic and diazotrophic populations, including the Thiobacillus genus, while Patescibacteria are suggested to attach only after the biofilm has formed [40, 68]. Further FSG MAGs/SAGs in this cluster included a novel family (AHRL-69.4-F21) in Veblenbacterales (Patescibacteria), an order containing other representatives identified in a shallow aquifer [17]; a novel family (AHRL-69.4-F24) in the order H5-PLA8 from the Planctomycetota; a Desulfobacterota class (AHRL-69.4-C1); a novel family (AHRL-69.4-F26) within the class Methylacidiphilales suggested to feature aerobic and acidophilic representatives [69]; and a novel Xenobia order (AHRL-69.4-O2) from the Eremiobacterota of which some orders are facultative anaerobes [70]. Of the cluster IV MAGs/SAGs, AHRL-69.4-F2 and AHRL-69.4-F4 were identified as having significant numbers of RNA transcripts in the AHRL.69.4 groundwater. Both these novel archaeal families were from the Bathyarchaeia (Thermoproteoata) that encode various one-carbon utilization strategies, including methane oxidation [71].

MAG cluster V was ubiquitously represented in the Äspö HRL AHRL.69.4 and AHRL.200.6 boreholes but rare in the other groundwaters. The MAGs/SAGs in this cluster included seven aligning with the Patescibacteria, of which AHRL-69.4-F13 and AHRL-69.4-F14 (both order JAACEG01) were also active in the AHRL.69.4 borehole. In addition, AHRL-69.4-F3 (Bathyarchaeia order RBG-16-48) and Planctomycetota AHRL-69.4-O10 were active in this groundwater.

The remaining MAG/SAG clusters VI–X included novel taxa identified in several of the Äspö HRL groundwaters but again were rare in Olkiluoto. These included the *Omnitrophota* family AHRL-200.6-F12 suggested to feature traits, such as being ultra-small and host-dependent to survive low carbon and energy conditions [56] and a novel *Pacearchaeales* taxa at family level (AHRL-200.6-F3) belonging to *Nanoarchaeota* DPANN archaea [12]. Co-occurrence analysis for populations identified in groundwaters of varying depths suggested the order *Pacearchaeales* from the

Nanoarchaeota were putative symbiotic partners of Patescibacteria [72]. The active novel taxa from these clusters were solely identified in groundwater AHRL.200.6 and included AHRL-200.6-O12 from the Patescibacteria class UBA1384 previously suggested to contribute to a New Zealand groundwater ultra-small prokaryote RNA transcript-based activity [67]; AHRL-200.6-F14 from the ABY1 class; the novel family AHRL-200.6-F10 from the Margulisbacteria [17]; and a Patescibacteria family AHRL-200.6-F17 previously suggested to lack genes for amino acid or nucleotide biosynthesis [53].

Metabolic potential for carbon and nitrogen fixation

Autotrophic carbon fixation encoded in the novel classes, orders, and families of the FSG included the Wood–Ljungdahl pathway (37 out of the 112 novel MAGs/SAGs affiliated to 21 phyla) along with ten MAGs/SAGs from seven phyla carrying genes coding for the reverse TCA cycle (Fig. 5 and Supplementary Fig. S1). Genes encoding the Wood-Ljungdahl pathway were present in 6 out of the 7 Omnitrophota, 3 out of the 4 Desulfobacterota, all three UBA9089, 3 of the 5 Elusimicrobiota, both Myxococcota and VGIX01 (formerly Candidatus Eisenbacteria [17]), and 1 Altiarchaeota MAGs/SAGs. The presence of the Wood-Ljungdahl pathway has previously been reported for the Omnitrophota [56], Desulfobacterota plus UBA9089 [20], Elusimicrobiota [58], Myxococcota [73], VGIX01 [17], and Altiarchaeota [74]. This pathway is often prevalent in the deep biosphere (e.g. South Africa and USA [19, 75]), as it is favored by anaerobic populations close to the thermodynamic limits of life and requires strict anoxia [76]. In addition, genes coding for the reverse TCA cycle were identified in all three UBA10199, 2 out of 4 of Desulfobacterota, and a single MAG from the Undinarchaeota, Altiarchaeota, and Thermoplasmatota. This confirmed a previous report that both Desulfobacterota and UBA9089 use the reverse TCA cycle and conserve energy from sulfate reduction [77].

Nitrogen fixation potential was identified in three of the novel MAGs/SAGs, namely two out of four *Desulfobacterota* and one of the two Myxococcota. However, the novel order AHRL-69.4-O12 assigned to the previously reported diazotrophic phylum UBA6262 [78] lacked apparent genes for nitrogen fixation, potentially narrowing the metabolic versatility of this deep biosphere candidate taxa or that the genes were not assembled in the incomplete MAG. Nitrogen fixation is common in some terrestrial deep biosphere environments and has been reported in the Horonobe Underground Research Laboratory in Japan [79].

Metabolic potential for oxidation of inorganic electron donors

Genes encoding electron donors that were potentially capable of supporting a lithotrophic lifestyle included sulfide oxidation (as represented by fccB and sqr) and sulfur oxidation (sdo and sor) that were present in one and 50 of the 112 novel MAGs/SAGs, respectively. The novel taxa encoding genes for sulfur oxidation included all four Desulfobacterota MAGs/SAGs, all three UBA9089 MAGs/SAGs, and one Margulisbacteria MAG/SAG (AHRL-200.6-O7). Margulisbacteria were previously identified as mediating sulfate/sulfite reduction [20, 59, 63], and these data suggested that the novel FSG MAG/SAG was able to carry out both the oxidative and reductive steps in sulfur cycling [80]. While Desulfobacterota encompass sulfate-reducing populations, the presence of sulfur oxidation genes also expands the metabolic repertoire of this phylum. While Patescibacteria are often considered to have simple fermentative lifestyles encoded by a reduced genome (e.g. Saccharimonadia [81]), the Portnoybacterales have been implicated in sulfur

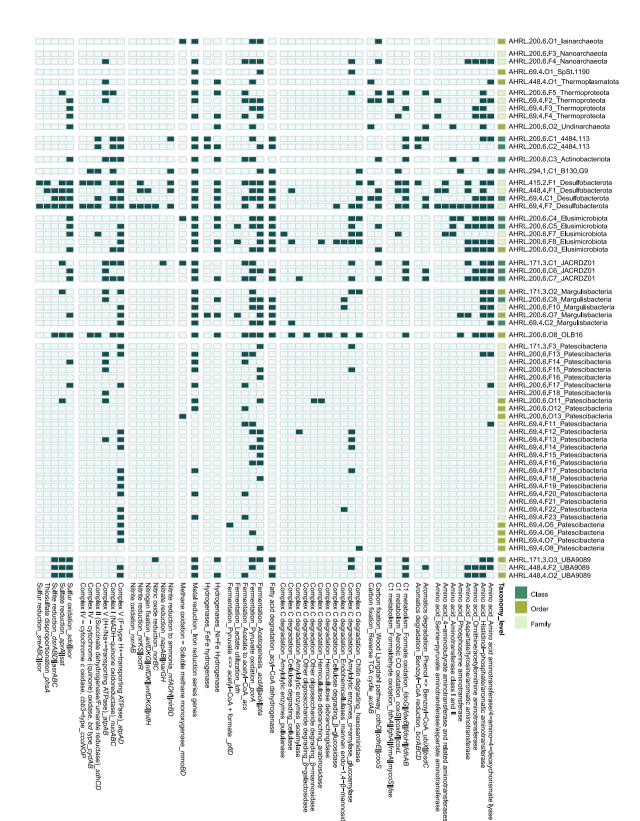


Figure 5. Presence of selected metabolic traits in a sub-set of the novel FSGs (full version of the figure is available in Supplementary Fig. S3). Certain categories, such as "Complex carbon degradation_Hemicellulose debranching," that are made up of several individual genes (and consequently data lines) in Supplementary Fig. S3 have been amalgamated for space considerations. Presence is indicated by dark color and absence light (left). The rank of novel taxa is indicated on the right.

cycling [17] even if the novel AHRL-200.6-F18 family (belonging to the *Portnoybacterales*) lacked genes for sulfur oxidation. In contrast, the novel families AHRL-200.6-F17 and AHRL-200.6-F14 MAGs/SAGs (*Paceibacterales* and SC72 orders, respectively) contained genes for sulfur oxidation, suggesting several *Patescibacteria* lineages have a broader metabolic repertoire than simple fermentative processes and highlights the importance of investigating their underexplored diversity and metabolic potential.

Another potential electron donor is hydrogen, which can be produced by, e.g. water radiolysis and serpentinization. Hydrogen is consumed by respiratory or bi-directional Ni/Fe hydrogenases [82], such as group 1 (10 MAGs/SAGs), group 3c (23 MAGs/SAGs), and group 3abd (5 MAGs/SAGs), which were identified in 26 phyla and thereby clearly demonstrating that capacity for hydrogen oxidation is a common trait across these novel taxa. Hydrogen oxidation is reported in the Fennoscandian Shield subsurface [83], and MAGs/SAGs with these genes included the previously reported hydrogen oxidizers *Margulisbacteria* [17], *Desulfobacterota* [84], and RUG730 that was formerly classified as Elusimicrobium [85]. Hydrogen-dependent terrestrial deep biosphere environments have been reported from around the globe (e.g. [75, 86]) with its importance suggested to increase with depth from the surface (reviewed in [2]).

Metabolic potential for oxidation of organic electron donors

An additional source of nutrients and energy in deep groundwater is necromass [28], which here is broadly defined as polymers and dissolved organic constituents of biogenic origin either from dead cells or excreted compounds produced by, e.g. fermentation (Fig. 5 and Supplementary Fig. S1). Processes related to polymer hydrolysis, amino acid use, and fermentation were present among the novel taxa that lacked other apparent energy acquisition mechanisms (all but two of the novel Patescibacteria and all the 4484-113, Aenigmatarchaeota, GCA–001730085, and Verrucomicrobiota phyla). Nevertheless, some of these taxa are known to be sustained by symbiotic interactions for life in deep groundwaters, e.g. compensating for gaps in their metabolic capacity, while concomitantly providing excreted metabolites to their symbiotic partners [20, 87, 88]. Among the novel taxa, certain phyla, such as Planctomycetota, UBP15, Sumerlaeota, Chloroflexota, and OLB16, featured an abundance of biopolymer degrading pathways for hydrolysis of chitin and carbohydrates. The presence of these pathways in the novel taxa suggested capacity to use and assimilate necromass in this environment. However, it should be noted that many key genes involved in heterotrophic utilization of these compounds can have other roles in the cell, including recycling of cellular materials also in autotrophs. A further pathway potentially involved in necromass utilization is degradation of the aromatic amino acid phenylalanine incorporating anaerobic aromatic degradation mediated by bcrABCD that was limited to five of the novel taxa, such as 4484-113, one Chloroflexota, and one Desulfobacterota MAG/SAG.

Metabolic potential for methane oxidation

Methane is an additional relevant electron donor in deep biosphere habitats, with methanotrophy enabled by the *mmoBD* genes. These genes were present in eight of the novel taxa of MAGs/SAGs, including two MAGs/SAGs from the *Planctomycetota*, one MAG from the *Patescibacteria*, SZUA–182, JACRDZ01, *Iainar chaeota*, *Elusimicrobiota*, and AABM5–125–24. The identification of methanotrophy in, e.g. the *Patescibacteria* widens the previously recognized genetic capability of affiliated populations. Active anaerobic oxidation of methane has previously been reported for the Fennoscandian Shield deep subsurface and is potentially coupled to sulfur cycling [89].

Metabolic potential for energy conservation via electron transport

Energy conservation can be achieved via nitrate reduction to nitrite enabled by napAB/narGH that were present in five MAGs/SAGs, or via nitrite reduction to ammonia enabled by nrfADH/nirBD present in nine MAGs/SAGs, and finally via denitrification enabled by nirKS/octR found in three MAGs/SAGs. These nitrogen reduction processes were highly represented in the Desulfobacterota novel lineages as previously described [90]. A further energy conservation pathway is the reduction of sulfur-(sreABC/sor) present in 2 MAGs/SAGs, sulfite-(dsrABD/asrABC) present in 12 MAGs/SAGs, and sulfate (aprA/sat) present in 23 MAGs/SAGs. These MAGs/SAGs included sulfate/sulfite/sulfur reduction in the four, three, and two Desulfobacterota MAGs/SAGs, respectively. In addition, sulfate/sulfite reduction was present in all three UBA9089 MAGs/SAGs, the single UBP15 plus OLB16 MAGs/SAGs (formerly Candidatus Omnitrophica [12]), and both Myxococcota MAGs/SAGs. Sulfate reduction has previously been reported for the Desulfobacterota [59], UBA9089 [20, 63], and OLB16 (Ca. Omnitrophica) [56]. Sulfate/sulfite/sulfur reduction is commonly suggested to be an important energy conservation method in the terrestrial deep biosphere [20, 21, 80]. Energy conservation can also be mediated by ferric iron [91] and manganese [92] reduction that was present in 90 of the novel MAGs/SAGs, although it remains uncertain whether these genes mediated electron transfer. Finally, while one of the three novel Thermoproteota AHRL-200.6-F5 could reduce sulfate, it was not suggested to be capable of methanogenesis as described for Ca. Methanodesulfokores washburnensis [61].

Electron transport via complexes I, II, and IV was common among the novel taxa, with the exception of, e.g. Patescibacteria, which have been reported as having an energy conservation predominantly via fermentation [87]. In contrast, one or both of the two ATPase complexes (F- and V-types) were also broadly present in fermenters, such as Patescibacteria (19 out of 27 MAGs/SAGs). The ability to produce an F-type ATPase has previously been reported for the cultured Patescibacteria (Saccharibacteria) Southlakia epibionticum [93]. Such ATPases can operate in reverse by pumping out protons to sustain a proton motive force, but this was considered less likely in S. epibionticum, where instead it is suggested to use the ATPase in pH homeostasis. However, it was unclear which of these potential roles the ATPase plays in the deep groundwater populations. An alternative electron transport system (among other functions of this complex) is the RNF complex that was present in 41 of the 112 novel taxa (based on the presence of at least three out of the six mf genes), including two each of the Desulfobacterota and Elusimicrobiota, one of the two Nanoarchaeota, all the JACRDZ01, JAFGOL01, Margulisbacteria, Omnitrophota, and Ca. Eisenbacteria VGIX01 (Supplementary Fig. S2). The RNF complex was widely prevalent among the novel taxa as well as the deep biosphere community, suggesting it may act as a basal electron transport system that requires less energy than complexes I-IV, which could potentially be an adaptation to the deep groundwater milieu. For instance, the RNF complex is utilized for autotrophic CO₂ reduction [77] at the "thermodynamic limit of life" [94].

Metabolic potential for energy conservation via fermentation

In the absence of thermodynamically favorable electron acceptors, fermentation emerges as a feasible and widespread

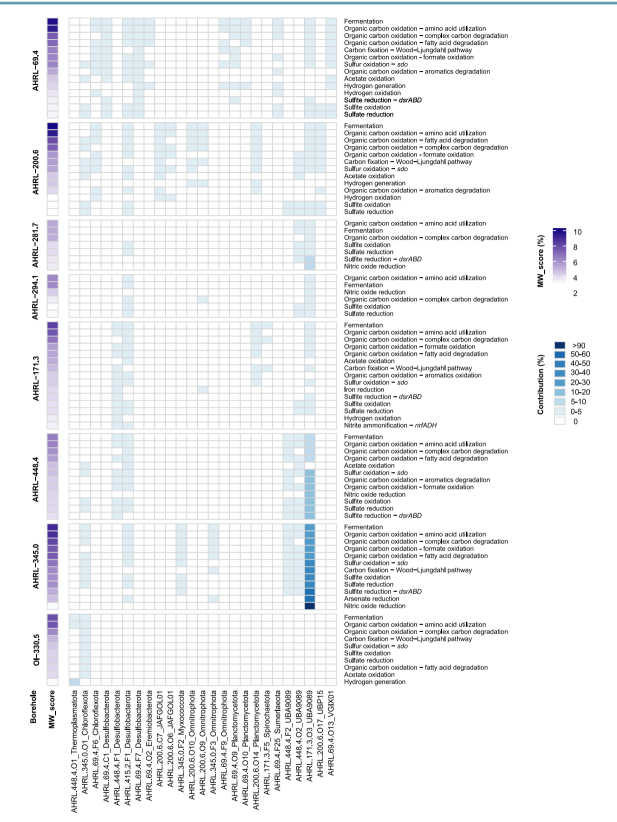


Figure 6. Contribution of selected novel taxa in functions with MW score \geq 3 (MAGs with zero percent contribution are not shown) and the metabolic processes they contribute to. Borehole OI-366.7 that lacked any MAGs/SAGs contributing to the metabolisms was omitted. No contribution was observed between 60% and 89%.

metabolic strategy for organic carbon degradation (e.g. in South Dakota, USA, groundwaters [55]). In accordance with this, the capacity for several fermentative pathways (e.g. acetogenesis, lactic acid fermentation, and pyruvate oxidation based upon

their potential to contribute to fermentation pathways while alternative pathways for the putative fermentation genes were incomplete in that MAG/SAG) was detected across many of the novel taxa with the potential for production of both acetate via the Wood-Ljungdahl pathway and hydrogen via Fe/Fe hydrogenases. The hydrogen would then likely act as an electron donor for pathways, such as sulfate reduction [63] and methanogenesis [95], since high concentrations of fermentation products are unlikely to accumulate in low carbon and energy conditions. Additionally, oxidation of one-carbon fermentation products, such as formate and formaldehyde, was prevalent among the novel taxa except for representatives of the *Patescibacteria*, *Margulisbacteria*, *Elusimicrobiota*, and *Nanoarchaeota*. These fermentation products could also play a role in reciprocal metabolic partnerships in deep groundwater biomes [20].

Community-level contribution of novel taxa to metabolism

The community-scale metabolic capacity of each borehole (expressed as a percentage) for different metabolic functions was calculated by the METABOLIC tool [49], via a metric called metabolic weight score (MW-score; Supplementary Fig. S3). To calculate the MW-score for each function, the coverage values of genomes containing that function are summed and normalized by the overall coverage of the function [49]. A higher MW-score indicates more frequently shared functions and their increased abundance in the microbial community in each borehole. Furthermore, the contribution percentage of each MAG to each metabolic module function is determined, providing insights into the relative contributions of different microbial groups.

For the entire community in each groundwater type, these included fermentation (\leq 9.9%), amino acid utilization (\leq 9.5%), complex carbon (\leq 8.9%), aromatic degradation (\leq 5.6%), and fatty acid degradation (\leq 8.0%), pointing at one of the main nutrient sources in FSG groundwater (necromass). Of the carbon fixation processes, the Wood–Ljungdahl pathway was most prevalent (\leq 6.4%) except for borehole Ol-366.7, where the Calvin–Benson–Bassham cycle (7.5%) was the dominant autotrophic process. Furthermore, the novel taxa contributed to the oxidation of lithotrophic electron donors, including sulfur species (\leq 6.7%) and hydrogen (\leq 8.6%). Finally, the novel taxa also contributed significantly to energy conservation via electron transport that included nitrate (\leq 4.2% for *napAB*) and sulfate (\leq 6.0%) reduction.

Among the novel taxa, AHRL-171.3-O3, AHRL-448.4-O2, and AHRL-448.4-F2 MAGs/SAGs from the UBA9089 phylum strongly contributed to important metabolic functions in three Äspö HRL boreholes (Fig. 6 and Supplementary Table S4). For instance, the combined contribution of AHRL-171.3-O3 and AHRL-448.4-O2 in borehole AHRL-345.0 accounted for 91.9% of the total community's nitric oxide reduction; 59.5% plus 51.4% of the total community's sulfite and sulfate reduction; 50.1% of the total community's carbon fixation by the Wood-Ljungdahl pathway; and 51.4% plus 45.5% of the total community's sulfite and sulfur oxidation, respectively. In addition to the novel UBA9089, the AHRL-345.0-O1 MAG/SAG attributed to the Anaerolineae contributed e.g. an additional 3.2% each to the total AHRL-345.0 borehole community's Wood-Ljungdahl pathway, sulfite oxidation, and sulfate reduction MW-scores, suggesting this groundwater was dominated by the two UBA9089 and one Chloroflexota novel taxa.

Other influential novel MAGs/SAGs contributing to the MWscores included AHRL-448.4-O1 from the Thermoplasmatota contributing 6.1% of the hydrogen generation in groundwater OI-330.5, along with AHRL-69.4-O10 (*Planctomycetota*) and AHRL-69.4-O13 (VGIX01), contributing 2.7% and 1.7 to groundwater AHRL-69.4, respectively. Cryptic cycling of hydrogen in the AHRL-69.4 groundwater may have been partially mediated by AHRL-69.4-F6 (*Chloroflexota*) contributing 1.7% of the AHRL-69.4 hydrogen oxidation capacity. Finally, *Desulfobacterota* (AHRL-415.2-F1) was also suggested to play a key role in various metabolic processes in groundwater AHRL-200.6, including fermentation (0.8%), five classes of organic carbon oxidation (totaling 5.0%), acetate oxidation (1.8%), sulfur and sulfite oxidation (1.4% and 2.6%), and sulfate reduction (2.6%).

Conclusions

The high number of novel FSG taxa corroborates the deep terrestrial biosphere as a reservoir of undiscovered microbial diversity that encodes a wide range of metabolic strategies to survive and significantly contribute to different processes in the ecosystem. These active novel taxa contributed to the metabolic potential for fermentation, amino acid utilization, complex carbon, aromatics, and fatty acid degradation, while expanding the metabolic capabilities of representatives in several phyla. The prevalence of the RNF complex in many novel taxa suggested that this may be an adaptation for energy conservation in low carbon and energy conditions. Community-level analysis revealed key contributions, particularly from representatives of the UBA9089 phylum, in biogeochemical processes. This exploration unveiled novel taxa with intriguing metabolic capabilities, offering valuable insights into deep biosphere microbial diversity and functional potential in emphasizing their significance in Fennoscandian Shield groundwater nutrient and energy cycling.

Acknowledgements

The Swedish Nuclear Fuel and Waste Management Co (S.K.B.) is acknowledged for access to the Äspö HRL and Sicada database. Magnus Ståhle is acknowledged from sampling of Äspö HRL groundwaters and preparation of nucleic acids for sequencing. 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 (NAISS 2023/22-893, 2023/6-261, and 2024/5-56). The computations were enabled by resources provided by the National Academic Infrastructure for Supercomputing in Sweden (NAISS) and the Swedish National Infrastructure for Computing (SNIC) at UPPMAX, Uppsala University partially funded by the Swedish Research Council through grant agreements no. 2022-06725 and no. 2018-05973. A portion of the nucleic acid sequencing included in this study was carried out at by the U.S. Department of Energy Joint Genome Institute, a DOE Office of Science User Facility, is supported under Contract No. DE-AC02-05CH11231. Funding for open-access publishing provided by Linnaeus University.

Author contributions

M.D., T.N.N., and S.B. conceived the study; S.T. and G.W. collected samples and G.W. and K.A. extracted nucleic acids; L.M.L. extracted nucleic acids and prepared sequencing libraries; M.R.S., C.G., M.M., M.B., and E.N. analyzed the data sets; and M.D., M.M., and S.B. drafted the manuscript with input from all authors. All authors have read and approved the manuscript before submission.

Correspondence and requests for materials should be addressed to Mark Dopson.

Supplementary material

Supplementary material is available at ISME Communications online.

Conflicts of interest

None declared.

Funding

The study was supported by The Swedish Research Council (contract 2018-04311) to M.D., Wenner-Gren Stiftelserna contract UPD2021-0082 to M.D. and M.R.S., and Olle Engkvists Stiftelse contract 216-0462 to M.D. and C.G. S.B. acknowledges funding from the Swedish Research Council (grant 2017-04422) and Science for Life Laboratory. M.M. was supported by a grant from the Swedish research council Formas (grant no. 2021-00546). T.N.N. and L.M.L. were supported by the Laboratory Directed Research and Development Program of Lawrence Berkeley National Laboratory under U.S. Department of Energy (DOE) Contract No. DE-AC02-05CH11231.

Data availability

Previously released FSG MAGs can be accessed through the NCBI BioProject under the accession number PRJNA627556. SAGs are publicly available in figshare with the identifier https://doi.org/10.6084/m9.figshare.12170313 under the project name "Fennoscandian Shield genomic database (FSGD)." The additional MAGs generated for this study are deposited under the BioProject accession number PRJNA1023754 and their accession numbers are given in Table 1 and Supplementary Table S1. A R Markdown file is available in Supplementary File S2.

References

- Soares A, Edwards A, An D et al. A global perspective on bacterial diversity in the terrestrial deep subsurface. Microbiology 2023;169:001172. https://doi.org/10.1099/mic.0.001172
- Beaver RC, Neufeld JD. Microbial ecology of the deep terrestrial subsurface. ISME J 2024;18:wrae091. https://doi.org/10.1093/ ismejo/wrae091
- Overholt WA, Trumbore S, Xu X et al. Carbon fixation rates in groundwater similar to those in oligotrophic marine systems. Nat Geosci 2022;15:561–7. https://doi.org/10.1038/ s41561-022-00968-5
- Puente-Sánchez F, Arce-Rodríguez A, Oggerin M et al. Viable cyanobacteria in the deep continental subsurface. Proc Natl Acad Sci USA 2018;115:10702–7. https://doi.org/10.1073/pnas. 1808176115
- Lau MCY, Kieft TL, Kuloyo O et al. An oligotrophic deepsubsurface community dependent on syntrophy is dominated by sulfur-driven autotrophic denitrifiers. Proc Natl Acad Sci USA 2016;113:E7927–36. https://doi.org/10.1073/pnas.1612244113
- Holmfeldt K, Nilsson E, Simone D et al. The Fennoscandian Shield deep terrestrial virosphere suggests slow motion 'boom and burst' cycles. Commun Biol 2021;4:307. https://doi.org/10.1038/ s42003-021-01810-1
- Rahlff J, Turzynski V, Esser SP et al. Lytic archaeal viruses infect abundant primary producers in Earth's crust. Nat Commun 2021;12:4642. https://doi.org/10.1038/s41467-021-24803-4

- Nuppunen-Puputti M, Kietäväinen R, Purkamo L et al. Rock surface fungi in deep continental biosphere—exploration of microbial community formation with subsurface in situ biofilm trap. Microorganisms 2021;9:64. https://doi.org/10.3390/ microorganisms9010064
- Wanger G, Southam G, Onstott TC. Structural and chemical characterization of a natural fracture surface from 2.8 kilometers below land surface: biofilms in the deep subsurface. *Geomicrobiol J* 2006;23:443–52.
- Lopez-Fernandez M, Åström M, Bertilsson S et al. Depth and dissolved organic carbon shape microbial communities in surface influenced but not ancient saline terrestrial aquifers. Front Microbiol 2018;9:2880. https://doi.org/10.3389/fmicb.2018. 02880
- Staley JT, Konopka A. Measurement of in situ activities of nonphotosynthetic microorganisms in aquatic and terrestrial habitats. Ann Rev Microbiol 1985;39:321–46. https://doi.org/10.1146/ annurev.mi.39.100185.001541
- Rinke C, Schwientek P, Sczyrba A et al. Insights into the phylogeny and coding potential of microbial dark matter. Nature 2013;499:431-7. https://doi.org/10.1038/nature12352
- Lloyd KG, Steen AD, Ladau J et al. Phylogenetically novel uncultured microbial cells dominate earth microbiomes. mSystems 2018;3:e00055–18. https://doi.org/10.1128/msystems.00055-18
- Hug LA, Baker BJ, Anantharaman K et al. A new view of the tree of life. Nat Microbiol 2016;1:16048. https://doi.org/10.1038/ nmicrobiol.2016.48
- Parks DH, Rinke C, Chuvochina M et al. Recovery of nearly 8,000 metagenome-assembled genomes substantially expands the tree of life. Nat Microbiol 2017;2:1533–42. https://doi.org/10.1038/ s41564-017-0012-7
- Spang A, Saw JH, Jørgensen SL et al. Complex archaea that bridge the gap between prokaryotes and eukaryotes. Nature 2015;521: 173–9. https://doi.org/10.1038/nature14447
- Anantharaman K, Brown CT, Hug LA et al. Thousands of microbial genomes shed light on interconnected biogeochemical processes in an aquifer system. Nat Commun 2016;7:13219. https:// doi.org/10.1038/ncomms13219
- Probst AJ, Ladd B, Jarett JK et al. Differential depth distribution of microbial function and putative symbionts through sedimenthosted aquifers in the deep terrestrial subsurface. Nat Microbiol 2018;3:328–36. https://doi.org/10.1038/s41564-017-0098-y
- Magnabosco C, Ryan K, Lau MC et al. A metagenomic window into carbon metabolism at 3 km depth in Precambrian continental crust. ISME J. 2016;10:730–41. https://doi.org/10.1038/ ismej.2015.150
- Mehrshad M, Lopez-Fernandez M, Sundh J et al. Energy efficiency and biological interactions define the core microbiome of deep oligotrophic groundwater. Nat Commun 2021;12:4253. https://doi. org/10.1038/s41467-021-24549-z
- Wu X, Holmfeldt K, Hubalek V et al. Microbial metagenomes from three aquifers in the Fennoscandian shield terrestrial deep biosphere reveal metabolic partitioning among populations. ISME J 2016;10:1192–203. https://doi.org/10.1038/ismej.2015.185
- Hugenholtz P, Chuvochina M, Oren A et al. Prokaryotic taxonomy and nomenclature in the age of big sequence data. ISME J 2021;15:1879–92. https://doi.org/10.1038/s41396-021-00941-x
- Hedlund BP, Chuvochina M, Hugenholtz P et al. SeqCode: a nomenclatural code for prokaryotes described from sequence data. Nat Microbiol 2022;7:1702–8. https://doi.org/10.1038/ s41564-022-01214-9
- 24. Motamedi M, Pedersen K. Desulfovibrio aespoeensis sp. nov., a mesophilic sulfate-reducing bacterium from deep groundwater

at Äspö Hard Rock Laboratory, Sweden. Int J Syst Bacteriol 1998;**48**: 311–5. https://doi.org/10.1099/00207713-48-1-311

- Kotelnikova S, Macario AJL, Pedersen K. Methanobacterium subterraneum sp. nov., a new alkaliphilic, eurythermic and halotolerant methanogen isolated from deep granitic groundwater. Int J System Evol Microbiol 1998;48:357–67.
- Kalyuzhnaya MG, Khmelenina VN, Kotelnikova S et al. Methylomonas scandinavica sp.nov., a new methanotrophic psychrotrophic bacterium isolated from deep igneous rock ground water of Sweden. Syst Appl Microbiol 1999;22:565–72. https://doi. org/10.1016/S0723-2020(99)80010-1
- 27. Westmeijer G, Mehrshad M, Turner S et al. Connectivity of terrestrial deep biosphere microbial biomes with surface communities. *Comm Biol* 2022;**5**:37. https://doi.org/10.1038/ s42003-021-02980-8
- Lopez-Fernandez M, Broman E, Wu X et al. Investigation of viable taxa in the deep terrestrial biosphere suggests high rates of nutrient recycling. FEMS Microbiol Ecol 2018;94:fiy121. https://doi. org/10.1093/femsec/fiy121
- Jørgensen BB, Andrén T, Marshall IPG. Sub-seafloor biogeochemical processes and microbial life in the Baltic Sea. Environ Microbiol 2020;22:1688–706. https://doi.org/10.1111/1462-2920. 14920
- Hubalek V, Wu X, Eiler A et al. Connectivity to the surface determines diversity patterns in subsurface aquifers of the Fennoscandian Shield. ISME J 2016;10:2447–58. https://doi. org/10.1038/ismej.2016.36
- Hallbeck L, Pedersen K. Characterization of microbial processes in deep aquifers of the Fennoscandian Shield. Appl Geochem 2008;23:1796–819. https://doi.org/10.1016/j.apgeochem.2008.02. 012
- Osterholz H, Turner S, Alakangas LJ et al. Terrigenous dissolved organic matter persists in the energy-limited deep groundwaters of the Fennoscandian Shield. Nat Commun 2022;13:4837. https:// doi.org/10.1038/s41467-022-32457-z
- Lever MA, Rogers KL, Lloyd KG et al. Life under extreme energy limitation: a synthesis of laboratory- and field-based investigations. FEMS Microbiol Rev 2015;39:688–728. https://doi. org/10.1093/femsre/fuv020
- Luef B, Frischkorn KR, Wrighton KC et al. Diverse uncultivated ultra-small bacterial cells in groundwater. Nat Commun 2015;6:6372. https://doi.org/10.1038/ncomms7372
- Kantor RS, Wrighton KC, Handley KM et al. Small genomes and sparse metabolisms of sediment-associated bacteria from four candidate phyla. mBio 2013;4:11. https://doi.org/10.1128/ mBio.00708-13
- Mathurin FA, Astrom ME, Laaksoharju M et al. Effect of tunnel excavation on source and mixing of groundwater in a coastal granitoidic fracture network. Environ Sci Technol 2012;46: 12779–86. https://doi.org/10.1021/es301722b
- Laaksoharju M, Gascoyne M, Gurban I. Understanding groundwater chemistry using mixing models. Appl Geochem 2008;23: 1921–40. https://doi.org/10.1016/j.apgeochem.2008.02.018
- Smellie JAT, Laaksoharju M, Wikberg P. Äspö, SE Sweden

 a natural groundwater-flow model derived from hydrogeological observations. J Hydrol 1995;172:147–69. https://doi. org/10.1016/0022-1694(95)02720-A
- Andersson J, Ahokas H, Hudson JA, Olkiluoto Site Description, 2006. 2007; Posiva; Olkiluoto; ISBN 978-951-652-151-3.
- 40. Wu X, Pedersen K, Edlund J et al. Potential for hydrogen-oxidizing chemolithoautotrophic and diazotrophic populations to initiate biofilm formation in oligotrophic, deep terrestrial

subsurface waters. Microbiome 2017;**5**:37. https://doi.org/10. 1186/s40168-017-0253-y

- Lopez-Fernandez M, Broman E, Simone D et al. Statistical analysis of community RNA transcripts between organic carbon and 'geogas' fed continental deep biosphere groundwaters. mBio 2019;10:e01470–19. https://doi.org/10.1128/mBio.01470-19
- 42. Lopez-Fernandez M, Simone D, Wu X *et al.* Metatranscriptomes reveal all three domains of life are active, but are dominated by bacteria in the Fennoscandian crystalline granitic continental deep biosphere. *mBio* 2018;**9**:e01792–18.
- Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 2014;30:2114–20. https://doi.org/10.1093/bioinformatics/btu170
- Li D, Liu CM, Luo R et al. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. Bioinformatics 2015;31:1674–6. https:// doi.org/10.1093/bioinformatics/btv033
- Kang DD, Li F, Kirton E et al. MetaBAT 2: an adaptive binning algorithm for robust and efficient genome reconstruction from metagenome assemblies. *PeerJ* 2019;**7**:e7359. https://doi. org/10.7717/peerj.7359
- Parks DH, Imelfort M, Skennerton CT et al. CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. *Genome Res* 2015;25:1043–55. https:// doi.org/10.1101/gr.186072.114
- 47. Chaumeil P-A, Mussig AJ, Hugenholtz P et al. GTDB-Tk v2: memory friendly classification with the genome taxonomy database. *Bioinformatics* 2022;**38**:5315–6. https://doi.org/10.1093/ bioinformatics/btac672
- Buck M, Mehrshad M, Bertilsson S. mOTUpan: a robust Bayesian approach to leverage metagenome-assembled genomes for core-genome estimation. NAR Genom Bioinform 2022;4:4. https:// doi.org/10.1093/nargab/lqac060
- Zhou Z, Tran PQ, Breister AM *et al*. METABOLIC: high-throughput profiling of microbial genomes for functional traits, metabolism, biogeochemistry, and community-scale functional networks. *Microbiome* 2022;**10**:33. https://doi.org/10.1186/s40168-021-012 13-8
- Westmeijer G. Microbial Life Deep Underground: From Anaerobic Cultures to Reconstructed Genomes, in Biology and Environmental Sciences. Växjö, Sweden: Linnaeus University, 2024.
- Chapelle FH, O'Neill K, Bradley PM et al. A hydrogen-based subsurface microbial community dominated by methanogens. Nature 2002;415:312–5. https://doi.org/10.1038/415312a
- Chivian D, Brodie EL, Alm EJ et al. Environmental genomics reveals a single-species ecosystem deep within earth. Science 2008;**322**:275–8. https://doi.org/10.1126/science.1155495
- Brown CT, Hug LA, Thomas BC et al. Unusual biology across a group comprising more than 15% of domain bacteria. Nature 2015;523:208–11. https://doi.org/10.1038/nature14486
- Herrmann M, Wegner C-E, Taubert M et al. Predominance of Cand. Patescibacteria in groundwater is caused by their preferential mobilization from soils and flourishing under oligotrophic conditions. Front Microbiol 2019;10:1407. https://doi.org/10.3389/ fmicb.2019.01407
- Momper L, Casar CP, Osburn MR. A metagenomic view of novel microbial and metabolic diversity found within the deep terrestrial biosphere at DeMMO: a microbial observatory in South Dakota, USA. Environ Microbiol 2023;25:3719–37. https:// doi.org/10.1111/1462-2920.16543
- 56. Seymour CO, Palmer M, Becraft ED et al. Hyperactive nanobacteria with host-dependent traits pervade Omnitrophota. Nat

Microbiol 2023;8:727-44. https://doi.org/10.1038/s41564-022-013 19-1

- Stingl U, Radek R, Yang H et al. "Endomicrobia": cytoplasmic symbionts of termite gut protozoa form a separate phylum of prokaryotes. Appl Environ Microbiol 2005;**71**:1473–9. https://doi. org/10.1128/AEM.71.3.1473-1479.2005
- Méheust R, Castelle CJ, Matheus Carnevali PB et al. Groundwater Elusimicrobia are metabolically diverse compared to gut microbiome Elusimicrobia and some have a novel nitrogenase paralog. ISME J 2020;14:2907–22. https://doi.org/10.1038/ s41396-020-0716-1
- Murphy CL, Biggerstaff J, Eichhorn A et al. Genomic characterization of three novel *Desulfobacterota* classes expand the metabolic and phylogenetic diversity of the phylum. Environ Microbiol 2021;23:4326–43. https://doi.org/10.1111/1462-2920.15614
- Wang Y, Wegener G, Hou J et al. Expanding anaerobic alkane metabolism in the domain of archaea. Nat Microbiol 2019;4: 595–602. https://doi.org/10.1038/s41564-019-0364-2
- Diao M, Dyksma S, Koeksoy E et al. Global diversity and inferred ecophysiology of microorganisms with the potential for dissimilatory sulfate/sulfite reduction. FEMS Microbiol Rev 2023;47:fuad058. https://doi.org/10.1093/femsre/fuad058
- 62. Bell E, Lamminmaki T, Alneberg J et al. Biogeochemical cycling by a low-diversity microbial community in deep groundwater. Front Microbiol 2018;**9**:02129. https://doi.org/10.3389/fmicb.2018.02129
- Anantharaman K, Hausmann B, Jungbluth SP et al. Expanded diversity of microbial groups that shape the dissimilatory sulfur cycle. ISME J 2018;12:1715–28. https://doi.org/10.1038/ s41396-018-0078-0
- 64. Kuever J, Rainey FA, Widdel F. Desulfobacterales Ord. In: Brenner D.J. et al. (eds.), Nov, in Bergey's Manual of Systematics of Archaea and Bacteria. New York: Springer, 2005, 959.
- Singleton CM, Petriglieri F, Kristensen JM et al. Connecting structure to function with the recovery of over 1000 high-quality metagenome-assembled genomes from activated sludge using long-read sequencing. Nat Commun 2021;12:2009. https://doi. org/10.1038/s41467-021-22203-2
- 66. Yamada T, Sekiguchi Y. Anaerolineaceae, in Bergey's Manual of Systematics of Archaea and Bacteria, 1–5; John Wiley & Sons, Inc., in association with Bergey's Manual Trust; Hoboken, New Jersey.
- 67. Gios E, Mosley OE, Weaver L et al. Ultra-small bacteria and archaea exhibit genetic flexibility towards groundwater oxygen content, and adaptations for attached or planktonic lifestyles. ISME Comm 2023;3:13. https://doi.org/10.1038/ s43705-023-00223-x
- Lopez-Fernandez M, Westmeijer G, Turner S et al. Thiobacillus as a key player for biofilm formation in oligotrophic groundwaters of the Fennoscandian Shield. npj Biofilms and Microbiomes 2023;9:41. https://doi.org/10.1038/s41522-023-00408-1
- Op den Camp HJ, Islam T, Stott MB et al. Environmental, genomic and taxonomic perspectives on methanotrophic Verrucomicrobia. Environ Microbiol Rep 2009;1:293–306. https://doi.org/10.1111/ j.1758-2229.2009.00022.x
- 70. Ji M, Williams TJ, Montgomery K et al. Candidatus Eremiobacterota, a metabolically and phylogenetically diverse terrestrial phylum with acid-tolerant adaptations. ISME J. 2021;15:2692–707. https:// doi.org/10.1038/s41396-021-00944-8
- Hou J, Wang Y, Zhu P et al. Taxonomic and carbon metabolic diversification of Bathyarchaeia during its coevolution history with early earth surface environment. Sci Adv 2023;9:eadf5069. https://doi.org/10.1126/sciadv.adf5069
- 72. Chaudhari NM, Overholt WA, Figueroa-Gonzalez PA et al. The economical lifestyle of CPR bacteria in groundwater

allows little preference for environmental drivers. Environ Microbiome 2021;**16**:24. https://doi.org/10.1186/s40793-021-00 395-w

- Murphy CL, Yang R, Decker T et al. Genomes of novel Myxocococta reveal severely curtailed machineries for predation and cellular differentiation. Appl Environ Microbiol 2021;87:e0170621. https:// doi.org/10.1128/AEM.01706-21
- Adam PS, Borrel G, Gribaldo S. An archaeal origin of the Wood-Ljungdahl H4MPT branch and the emergence of bacterial methylotrophy. Nat Microbiol 2019;4:2155-63. https://doi. org/10.1038/s41564-019-0534-2
- Momper L, Jungbluth SP, Lee MD et al. Energy and carbon metabolisms in a deep terrestrial subsurface fluid microbial community. ISMEJ 2017;11:2319–33. https://doi.org/10.1038/ ismej.2017.94
- Berg IA. Ecological aspects of the distribution of different autotrophic CO₂ fixation pathways. Appl Environ Microbiol 2011;**77**:1925–36. https://doi.org/10.1128/AEM.02473-10
- Ragsdale SW. Enzymology of the Wood–Ljungdahl pathway of acetogenesis. Ann N Y Acad Sci 2008;1125:129–36. https://doi. org/10.1196/annals.1419.015
- Dong X, Zhang C, Peng Y et al. Phylogenetically and catabolically diverse diazotrophs reside in deep-sea cold seep sediments. Nat Commun 2022;13:4885. https://doi.org/10.1038/ s41467-022-32503-w
- Hernsdorf AW, Amano Y, Miyakawa K et al. Potential for microbial H₂ and metal transformations associated with novel bacteria and archaea in deep terrestrial subsurface sediments. ISME J 2017;**11**:1915–29. https://doi.org/10.1038/ismej.2017.39
- Bell E, Lamminmäki T, Alneberg J et al. Active sulfur cycling in the terrestrial deep subsurface. ISME J 2020;14:1260–72. https:// doi.org/10.1038/s41396-020-0602-x
- Lemos LN, Medeiros JD, Dini-Andreote F et al. Genomic signatures and co-occurrence patterns of the ultra-small Saccharimonadia (phylum CPR/Patescibacteria) suggest a symbiotic lifestyle. Mol Ecol 2019;28:4259–71. https://doi.org/10.1111/ mec.15208
- Søndergaard D, Pedersen CNS, Greening C. HydDB: a web tool for hydrogenase classification and analysis. Sci Rep 2016;6:34212. https://doi.org/10.1038/srep34212
- Pedersen K. Subterranean microbial populations metabolize hydrogen and acetate under in situ conditions in granitic groundwater at 450 m depth in the Äspö Hard Rock Laboratory, Sweden. FEMS Microbiol Ecol 2012;81:217–29. https://doi. org/10.1111/j.1574-6941.2012.01370.x
- Caffrey SM, Park HS, Voordouw JK et al. Function of periplasmic hydrogenases in the sulfate-reducing bacterium *Desulfovibrio* vulgaris Hildenborough. J Bacteriol 2007;189:6159–67. https://doi. org/10.1128/JB.00747-07
- Geissinger O, Herlemann DP, Mörschel E et al. The ultramicrobacterium "Elusimicrobium minutum" gen. nov., sp. nov., the first cultivated representative of the termite group 1 phylum. Appl Environ Microbiol 2009;75:2831–40. https://doi.org/10.1128/ AEM.02697-08
- Li S, Mosier D, Kouris A et al. High diversity, abundance, and expression of hydrogenases in groundwater. ISME Comm 2024;4:ycae023. https://doi.org/10.1093/ismeco/ycae023
- Castelle CJ, Brown CT, Anantharaman K et al. Biosynthetic capacity, metabolic variety and unusual biology in the CPR and DPANN radiations. Nat Rev Microbiol 2018;16:629–45. https://doi. org/10.1038/s41579-018-0076-2
- 88. Schwank K, Bornemann TLV, Dombrowski N et al. An archaeal symbiont-host association from the deep terrestrial

subsurface. ISME J 2019;**13**:2135–9. https://doi.org/10.1038/ s41396-019-0421-0

- Bell E, Lamminmäki T, Alneberg J et al. Active anaerobic methane oxidation and sulfur disproportionation in the deep terrestrial subsurface. ISME J 2022; 16:1583–93. https://doi.org/10.1038/ s41396-022-01207-w
- Langwig MV, De Anda V, Dombrowski N et al. Large-scale protein level comparison of Deltaproteobacteria reveals cohesive metabolic groups. ISME J 2022;16:307–20. https://doi. org/10.1038/s41396-021-01057-y
- 91. Garber AI, Nealson KH, Okamoto A et al. FeGenie: a comprehensive tool for the identification of iron genes and iron gene neighborhoods in genome and metagenome assemblies. Front Microbiol 2020;11:11. https://doi.org/10.3389/fmicb.2020. 00037
- 92. Wang X, Xie GJ, Tian N et al. Anaerobic microbial manganese oxidation and reduction: a critical review. Sci Total Environ 2022;822:153513. https://doi.org/10.1016/j.scitotenv.20 22.153513
- 93. Wang Y, Gallagher LA, Andrade PA et al. Genetic manipulation of Patescibacteria provides mechanistic insights into microbial dark matter and the epibiotic lifestyle. Cell 2023;186:4803–4817.e13. https://doi.org/10.1016/j.cell.2023.08.017
- Schuchmann K, Muller V. Autotrophy at the thermodynamic limit of life: a model for energy conservation in acetogenic bacteria. Nat Rev Micro 2014;12:809–21. https://doi.org/10.1038/ nrmicro3365
- Costa KC, Leigh JA. Metabolic versatility in methanogens. Curr Opin Biotechnol 2014;29:70–5. https://doi.org/10.1016/j. copbio.2014.02.012