Estimation of microbial metabolism and co-occurrence patterns in fracture groundwaters of deep crystalline bedrock at Olkiluoto, Finland

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Abstract

The microbial diversity in oligotrophic isolated crystalline Fennoscandian Shield bedrock fracture groundwaters is great but the core community has not been identified. Here we characterized the bacterial and archaeal communities in 12 water conductive fractures situated at depths between 296 m and 798 m by high throughput amplicon sequencing using the Illumina HiSeq platform. A sequencing depth of up to $1.2 \times 10^6$ reads per sample revealed that up to 95% and 99% of the bacterial and archaeal sequences obtained, respectively, belonged to only a few common species, i.e. the core microbiome. However, the remaining rare microbiome contained over 3 and 6 fold more bacterial and archaeal taxa. Several clusters of co-occurring rare taxa were identified, which correlated significantly with physicochemical parameters, such as salinity, concentration of inorganic or organic carbon, sulphur, pH and depth. The metabolic properties of the microbial communities were predicted using PICRUSt. The approximate estimation showed that the metabolic pathways included commonly fermentation, fatty acid oxidation, glycolysis/gluconeogenesis, oxidative phosphorylation and methanogenesis/anaerobic methane oxidation, but carbon fixation through the Calvin cycle, reductive TCA cycle and the Wood-Ljungdahl pathway was also predicted. The rare microbiome is an unlimited source of genomic functionality in all ecosystems. It may consist of remnants of microbial communities prevailing in earlier environmental conditions, but could also be induced again if changes in their living conditions occur. In this study only the rare taxa correlated with any physicochemical parameters. Thus
these microorganisms can respond to environmental change caused by physical or biological factors that may lead to alterations in the diversity and function of the microbial communities in crystalline bedrock environments.

1 Introduction

Identifying and understanding the core microbiome of any given environments is of crucial importance for predicting and assessing environmental change both locally and globally (Shade and Handelsman, 2012). In a previous study (Bomberg et al., 2015) we showed by 454 amplicon sequencing that the active microbial communities in Olkiluoto deep subsurface were strictly stratified according to aquifer water type. Nevertheless, more rigorous sequencing efforts and more samplings have shown that an archaeal core community consisting of the DeepSea Hydrothermal Vent Euryarchaeotal Group 6 (DHVEG-6), ANME-2D and Terrestrial Miscellaneous Group (TMEG) archaea may exists in the anaerobic deep groundwater of Olkiluoto (Miettinen et al., 2015). The bacterial core groups in Olkiluoto deep groundwater include at least members of the Pseudomonadaceae, Comamonadaceae and Sphingomonadaceae (Bomberg et al., 2014; 2015; Miettinen et al., 2015). The relative abundance of these main groups varies at different depths from close to the detection limit to over 90% of the bacterial or archaeal community (Bomberg et al., 2015; Miettinen et al., 2015). However, both the archaeal and the bacterial communities contain a wide variety of smaller bacterial and archaeal groups, which are distributed unevenly in the different water conductive fractures.

The rare biosphere is a concept describing the hidden biodiversity of an environment (Sogin et al., 2006). The rare biosphere consists of microbial groups that are ubiquitously distributed in nature but often present at low relative abundance and may thus stay below the limit of detection. Due to modern high throughput sequencing techniques, however, the hidden diversity of rare microbiota has been revealed. These microorganisms are the basis for unlimited microbial functions in the environment and upon environmental change specific groups can readily activate and become abundant. Access to otherwise inaccessible nutrients activate specific subpopulations in the bacterial communities within hours of exposure (Rajala et al., 2015) and enrich distinct microbial taxa at the expense of the original microbial community in the groundwater (Kutvonen, 2015). Mixing of different groundwater layers due to e.g. breakage of aquifer boundaries and new connection of separated aquifers may cause the microbial community to change and activate otherwise dormant processes. This has
previously been shown by Pedersen et al. (2013), who indicated increased sulphate reduction activity when sulphate-rich and methane-rich groundwater mixed. The stability of deep subsurface microbial communities in isolated deep subsurface groundwater fractures are assumed to be stable. However, there are indications that they may change over the span of several years as slow flow along fractures is possible (Miettinen et al., 2015; Sohlberg et al., 2015).

The microbial taxa present in an environment interact with both biotic and abiotic factors. Co-occurrence network analyses and metabolic predictions may help to understand these interactions. Barberan et al. (2012) visualised the co-occurrence networks of microbial taxa in soils and showed novel patterns connecting generalist and specialist species as well as associations between microbial taxa. They showed that specialist and generalist microbial taxa formed distinct and separate correlation networks, which also reflected the environmental settings. Metagenome predicting tools allows us to estimate microbial metabolic functions based on NGS microbiome data. Using the PICRUSt tool (Langille et al., 2013) Tsitko et al. (2014) showed that oxidative phosphorylation was the most important energy producing metabolic pathway throughout the 7 m depth profile of an Acidobacteria-dominated nutrient poor boreal bog. Cleary et al. (2015) showed that tropical mussel-associated bacterial communities could be important sources of bioactive compounds for biotechnology. This approach is nevertheless hampered by the fact that only little is so far known about uncultured environmental microorganisms and their functions and the PICRUSt approach is best applied for human microbiome for which it was initially developed (Langille et al., 2013). However, metagenomic estimations may give important indications of novel metabolic possibilities even in environmental microbiome studies.

Using extensive high throughput amplicon sequencing in this study we aimed to identify the core microbiome in the deep crystalline bedrock fractures of Olkiluoto Island and also to identify the rare microbiome. We aimed to show the interactions between the taxa of the rare biosphere and the surrounding environmental parameters in order to validate the factors that determine the distribution of the rare taxa. Finally, we aimed to estimate the prevailing metabolic activities that may occur in the deep crystalline bedrock environment of Olkiloto, Finland.
2 Materials and methods

2.1 Background

The Olkiluoto site has previously been extensively described (Posiva, 2013) and is only briefly described here. The Island of Olkiluoto situating on the west coast of Finland has approximately 60 drillholes drilled for research and monitoring purposes. Studies on the chemistry and microbiology of the groundwater have been on-going since the 1980s. The groundwater is stratified with a salinity gradient extending from fresh to brackish water to a depth of 30 m and the highest salinity concentration of 125 g L⁻¹ total dissolved solids (TDS) at 1000 m depth (Posiva, 2013). The most abundant salinity causing cations are Na⁺ and Ca²⁺ and anions Cl⁻. Between 100 and 300 m depths, the groundwater originates from ancient (pre-Baltic) seawater and has high concentrations of SO₄²⁻. Below 300 m the concentration of methane in the groundwater increases and SO₄²⁻ is almost absent. A sulphate-methane transition zone (SMTZ), where sulphate-rich fluid replaces methane-rich fluid, is located at 250 – 350 m depth. Temperature rises linearly with depth, from ca. 5 – 6 °C at 50 m to ca. 20 °C at 1,000 m depth (Ahokas et al., 2008). The pH of the groundwater is slightly alkaline throughout the depth profile. Multiple drillholes intersect several groundwater-filled bedrock fractures, including larger hydrogeological zones such as HZ20 or HZ21 (Table 1). The bedrock of Olkiluoto consists mainly of mica-gneiss and pegmatitic granite type rocks (Kärki & Paulamäki, 2006).

This study focused on 12 groundwater samples from water conductive fractures situated at between 296 m and 798 m below sea level bsl and originating from 11 different drillholes in Olkiluoto (Figure 1). The samples represented brackish sulphate waters and saline waters (as classified in Posiva, 2013). The samples were collected between December 2009 and January 2013 (Table 1). The physicochemical parameters of the groundwater samples have been reported by Miettinen et al. (2015), but have for clarity been collected here (Table 1).

2.2 Sample collection

The collection of samples occurred between December 2009 and January 2013 (Table 1) as described previously (Bomberg et al., 2015; Miettinen et al., 2015; Sohlberg et al., 2015). The samples were obtained from 11 different permanently packered or open drillholes equipped
with removable inflatable packers. The position and direction of the drillholes are indicated in Figure 1. Shortly, in order to obtain indigenous fracture fluids, the packer-isolated fracture zones were purged by removing stagnant drillhole water by pumping for a minimum of four weeks before the sample water was collected. The water samples were collected directly from the drillhole into an anaerobic glove box (MBRAUN, Germany) via a sterile, gas-tight poly acetate tube (8 mm outer diameter). Microbial biomass DNA extraction was concentrated from 1000 mL samples by filtration on cellulose acetate filters (0.2 µm pore size, Corning) by vacuum suction inside the glove box. The filters were immediately extracted from the filtration funnels and frozen on dry ice in sterile 50 ml cone tubes (Corning). The frozen samples were transported on dry ice to the laboratory where they were stored at -80°C until use.

2.3 Nucleic acid isolation

Community DNA was isolated directly from the frozen cellulose-acetate filters with the PowerSoil DNA extraction kit (MoBio Laboratories, Inc., Solana Beach, CA), as previously described (Bomberg et al., 2015). Negative DNA isolation controls were included in the isolation protocol. The DNA concentration of each sample was determined using the NanoDrop 1000 spectrophotometer.

2.4 Estimation of microbial community size

The size of the microbial community was determined by epifluorescence microscopy of 4',6 diamidino-2-phenylindole dihydrochloride (DAPI) (Sigma, MO, USA) stained cells as described in Purkamo et al. (2013). The size of the bacterial population was determined by 16S rRNA gene targeted quantitative PCR (qPCR) as described by Tsitko et al. (2014) using universal bacterial 16S rRNA gene-targeting primers fD1 (Weisburg et al., 1991) and P2 (Muyzer et al., 1993), which specifically target the V1- V3 region of the bacterial 16S rDNA gene. The size of the archaeal population in the groundwater was determined by using primers ARC344f (Bano et al., 2004) and Ar744r (reverse compliment from Barns et al., 1994) flanking the V4-V6 region of the archaeal 16S rRNA gene.

The qPCR reactions were performed in 10µL reaction volumes using the KAPA 2 × Syrb® FAST qPCR-kit on a LightCycler480 qPCR machine (Roche Applied Science, Germany) on white 96-well plates (Roche Applied Science, Germany) sealed with transparent adhesive
seals (4titude, UK). Each reaction contained 2.5 µM of relevant forward and reverse primer and 1 µL DNA extract. Each reaction was run in triplicate and no-template control reactions were used to determine background fluorescence in the reactions.

The qPCR conditions consisted of an initial denaturation at 95 °C for 10 minutes followed by 45 amplification cycles of 15 seconds at 95 °C, 30 seconds at 55 °C and 30 seconds at 72 °C with a quantification measurement at the end of each elongation. A final extension step of three minutes at 72 °C was performed prior to a melting curve analysis. This consisted of a denaturation step for 10 seconds at 95 °C followed by an annealing step at 65 °C for one minute prior to a gradual temperature rise to 95 °C at a rate of 0.11 °C s⁻¹ during which the fluorescence was continuously measured. The number of bacterial 16S rRNA genes was determined by comparing the amplification result (Cp) to that of a ten-fold dilution series (10¹-10⁷ copies µL⁻¹) of Escherichia coli (ATCC 31608) 16S rRNA genes in plasmid for bacteria and a dilution series of genomic DNA of Halobacterium salinarum (DSM 3754) for archaea. The lowest detectable standard concentration for the qPCRs was 10² gene copies/reaction. Inhibition of the qPCR by template tested by adding 2.17×10⁴ plasmid copies containing fragment of the morphine-specific Fab gene from Mus musculus gene to reactions containing template DNA as described in Nyyssönen et al. (2012). Inhibition of the qPCR assay by the template DNA was found to be low. The average Crossing point (Cp) value for the standard sample (2.17×10⁴ copies) was 28.7 (± 0.4 sd), while for the DNA samples Cp was 28.65 - 28.91 (± 0.03-0.28 sd). Nucleic acid extraction and reagent controls were run in all qPCRs in parallel with the samples. Amplification in these controls was never higher than the background obtained from the no template controls.

2.5 Amplicon library preparation

This study is part of the Census of Deep Life initiative, which strives to obtain a census of the microbial diversity in deep subsurface environment by collecting samples around the world and sequencing the 16S rRNA gene pools of both archaea and bacteria. The extracted DNA samples were sent to the Marine Biological Laboratory in Woods Hole, MA, USA, for preparation for HiSeq sequencing using the Illumina technology. The protocol for amplicon library preparation for both archaeal and bacterial 16S amplicon libraries can be found at http://vamps.mbl.edu/resources/faq.php. Shortly, amplicon libraries for completely overlapping paired-end sequencing of the V6 region of both the archaeal and bacterial 16S rRNA genes were produced as previously described (Eren et al., 2013). For the archaea,
primers A958F and A1048R containing Truseq adapter sequences at their 5’ end were used, and for the bacteria primers B967F and B1064R for obtaining 100 nt long paired end reads (https://vamps.mbl.edu/resources/primers.php). The sequencing was performed using a HiSeq 1000 system (Illumina).

2.6 Sequence processing and analysis

Contigs of the paired end fastq files were first assembled with mothur v 1.32.1 (Schloss et al., 2009). Analyses were subsequently continued using QIIME v. 1.8. (Caporaso et al., 2010). Only sequences with a minimum length of 50 bp were included in the analyses. The bacterial and archaeal 16S rRNA sequences were grouped into OTUs (97% sequence similarity) using both the open reference and closed reference OTU picking strategy and classified using the GreenGenes 13_8 16S reference database (DeSantis et al., 2006). The sequencing coverage was evaluated by rarefaction analysis and the estimated species richness and diversity indices were calculated. For comparable α- and β-diversity analyses the data sets were normalized by random subsampling of 17,000 sequences/sample for archaea and 140,000 sequences/sample for bacteria. Microbial metabolic pathways were estimated based on the 16S rRNA gene data from the closed OTU picking method using the PICRUSt software (Langille et al., 2013) on the web based Galaxy application (Goecks et al., 2010; Blankenberg et al., 2010; Giardine et al., 2005). The predicted KO numbers were plotted on KEGG pathway maps (http://www.genome.jp/kegg/) separately for the bacterial and archaeal predicted metagenomes, with a threshold of a minimum of 100 genes in total estimated from all samples. The sequence data has been submitted to the Sequence Read Archive (SRA, http://www.ncbi.nlm.nih.gov/sra) under study SRP053854, Bioproject PRJNA275225.

2.7 Statistical analyses

Non-metric multi-dimensional scaling plots using Chord’s similarity index were calculated separately for the archaeal and bacterial communities using PAST3 (Hammer and Harper, 2001). The samples were also hierarchically clustered based on community similarity using the UPGMA clustering with Bray-Curtis similarity index and 100 bootstrap repeats with PAST3. A co-occurrence network was calculated using the Gephi software (Bastian et al., 2009) using the Fruchtermann-Feingold layout with the betweenness centrality algorithm for identifying microbial taxa with numerous connections and the Louvain method (Blondel et al., 2008) for identifying closely associated groups of microbes. The calculations were based
on Spearman’s rank correlation calculations obtained by the out.association command in
mothur and only pairs with \( r > 0.6 \) and \( p < 0.01 \) were included in the network analysis and
nodes with a degree range of less than 10 were excluded from the graph. The \( p \) values were
not corrected and the Spearman’s rank correlations were only used to form pairs between taxa
for the network visualization.

3  Results

3.1  Microbial community size

The total number of microbial cells detected by epifluorescence microscopy of DAPI stained
cells was between \( 2.3 \times 10^4 \) and \( 4.2 \times 10^5 \) cells mL\(^{-1}\) groundwater (Figure 2, Table 1). The
concentration of bacterial 16S rRNA gene copies mL\(^{-1}\) varied between \( 9.5 \times 10^3 \) and \( 7.0 \times 10^5 \)
and that of the archaea \( 2.6 \times 10^1 \) and \( 6.3 \times 10^4 \) (Figure 2, Table 1).

3.2  Sequence statistics, diversity estimates and sequencing coverage

The number of bacterial v6 sequence reads from the 12 samples varied between \( 1.4 - 7.8 \times 10^5 \)
reads, with a mean sequencing depth of \( 2.9 \times 10^5 \) (\( \pm 1.8 \times 10^5 \) standard deviation)
reads/sample (Table 2). The archaeal v6 sequence reads ranged from \( 0.17 - 12.1 \times 10^5 \) reads
with a mean sequencing depth of \( 4.1 \times 10^5 \) (\( \pm 3.5 \times 10^5 \) standard deviation) reads/sample. The
numbers of observed operational taxonomic units (OTUs) represented on average 82.6\% (\( \pm \)
12.5\%) of the Chao1- and 78.1 \% (\( \pm \) 13.4\%) of the ACE-estimated numbers of bacterial
OTUs (Table 2ab). The archaeal communities were slightly better covered, with on average
88.5\% (\( \pm \) 11.5\%) of the Chao1 and 84.8\% (\( \pm \) 12.6\%) of the ACE estimated number of OTUs
detected. Shannon diversity index \( H' \), calculated from 140,000 and 17,000 random sequence
reads per sample for the bacteria and archaea, respectively, was high for both bacterial and
archaeal communities. High \( H' \) values and climbing rarefaction curves (Figure S1) indicated
high diversity in the microbial communities in the different deep groundwater fracture zones
of Olkiluoto. The bacterial \( H' \) was on average 13 (\( \pm \)0.74), ranging from 11 to 14 between the
different samples. The archaeal \( H' \) was on average 11 (\( \pm \)1.2) ranging from 9 to 12 between
the samples.
3.3 Microbial communities

From the bacterial v6 sequences 49 different bacterial Phyla were detected (Appendix 1). These phyla included 165 bacterial classes, 230 orders, 391 families and 651 genera. The greatest number of sequences, between 21.83% and 47.94% per sample, clustered into an undetermined bacterial group (Bacteria, Other), which may be due the fact that sequences of poorer quality may be difficult to classify, especially as the sequences are short.

Only 31 of the identified genera represented at least 1% of the bacterial sequence reads in any sample (Figure 3a).

The archaea were represented by two identified phyla, the Euryarchaeota and the Crenarchaeota (Appendix 2). These included 21 classes, 38 orders, 61 families and 81 genera. Between 4.7% and 35.0% of the archaeal sequences of each sample were classified to unassigned Archaea, with a general increase in unassigned archaeal sequences with increasing depth. 15 archaeal genera were present at a minimum of 1% relative abundance in any of the samples (Figure 3b).

The bacterial core community, i.e. the taxa detected in all the tested samples, constituted of 95 out of 651 identified bacterial genera (Appendix 3). These genera accounted for 80.78 – 95.81% of all the bacterial sequence reads in the samples. The archaeal core community consisted of 25 of the 81 identified genera and accounted for 95.05 – 99.75% of the total number of sequence reads in each sample (Appendix 4).

3.4 Environmental parameters driving the microbial communities

The microbial community profiles of the different samples were clustered in a UPGMA tree (Figure 4). The samples were loosely clustered according to depth with the deeper samples generally more associated with each other and the samples from shallower depths associating with each other. A similar trend was seen in the NMDS plots (Figures 6a and b), although the bacterial communities clustered the samples more tightly into three groups compared to the archaeal communities. No clear environmental factor was identified to drive the communities. However, the deepest bacterial communities were affected by the increasing salinity and the communities from the shallower depths were affected by the concentration of sulphides and the alkalinity, according to the NMDS plots. The archaeal communities, on the other hand, were affected by the concentration of sulphate, sulphur, sulphide, iron, bicarbonate and magnesium at 415 m and 510 m depth.
3.5 Co-occurrence network

The co-occurrence network (Figure 6, Figure S2) indicated specific bacterial taxa are central to the whole microbial community. In the network the size of the nodes indicates centrality (small node = low centrality, big node = high centrality) and the colour indicates the degree of connections (colour scale blue-green-yellow-red indicates increasing degree). Numerous microbial groups with specific functions, such as sulphate and sulphur reduction (e.g. *Desulfomonile*, *Desulfobacteraceae*, *Desulfovibionales*, *Desulfurispora*, *Planctomycetes*), oxidation of reduced sulphur compounds (e.g. *Sulfuricurvum*, *Sulfurimonas*, *Thiohalorhabdales*, *Thiobacterales*, *Sulfobacillaceae*), methylo trophy (*Methylophaga*, *Methylosinus*, *Methylococcales*, *Methyloversatilis*), nitrogen cycling (e.g. *Nitrospira*, *Rhizobiaceae*), syntrophic bacteria (e.g. *Syntrophaceae*, *Syntrophobacteraceae*) showed relatively high degrees of centrality and number of connections in the network. However, the majority of the taxa with the highest degree of centrality were heterotrophic bacteria capable of fermentation, such as the Elusimicrobia, *Exiguobacterium*, *Gordonia*, *Planctomycetes*, and taxa capable of degradation of recalcitrant organic molecules, such as *Kordiimonadales*.

3.6 Predicted metabolic functions of the deep subsurface microbial communities

The putative metabolic functions of the microbial communities at different depth was predicted using the PICRUSt software, which compares the identified 16S rRNA gene sequences to those of known genome sequenced species thereby estimating the possible gene contents of the uncultured microbial communities. The analysis is only an approximation, but may give an idea of the possible metabolic activities in the deep biosphere. In order to evaluate the soundness of the analysis a nearest sequenced taxon index (NSTI) for each of the bacterial and archaeal communities was calculated by PICRUSt. An NSTI value of 0 indicates high similarity to the closest sequenced taxon while NSTI=1 indicates no similarity. The NSTI of the bacterial communities at different depths varied between 0.045 in sample OL-KR44 and 0.168 in sample OL-KR13 (Figure 7). The NSTI for archaea were much higher ranging from 0.141 in sample OL-KR9 at depth of 432 m and 0.288 in OL-KR44. This indicates that the metagenomic estimates are only indicative. The estimated microbial metabolism did not differ noticeably between the different depths (Figure 8a and b). The most important predicted metabolic pathways included membrane transport in both bacterial and archaeal communities. The most common pathways for carbohydrate metabolism were the butanoate, propionate, glycolysis/gluconeogenesis and pyruvate metabolism pathways for the
bacteria and glycolysis/gluconeogenesis and pyruvate metabolism pathways for the archaea (Figure 9). Glucose is converted into pyruvate and further to Acetyl-CoA by both bacteria and archaea. The bacterial community may produce and utilize acetate. Both the bacterial and archaeal communities fix carbon via the Wood-Ljungdal (WL) reverse Citric acid cycle (rTCA) and Calvin pathways. Methane is produced from methylamines, CO₂ and methanol by the methanogenic archaea. Based on the predicted metagenomes the bacterial community is not able to oxidize methane or hydrolyze methanol, but the methylotrophs present may use formic acid and trimethylamines.

The most abundant energy metabolic pathway in the bacterial communities was the oxidative phosphorylation (Figure S3) while for the archaea the methane metabolism was the most important (Figure 9). Utilization of propanoate and butanoate (Figure 9) by the bacterial communities as well as well covered fatty acid biosynthesis and degradation pathways indicate that the bacterial community is capable of fermentation (Figure S4a and b). Nitrate is reduced both through dissimilatory nitrate reduction to ammonia and through denitrification to nitrous oxide by the bacteria (Figure S5). In addition, nitrogen is fixed to ammonia by both archaea and bacteria. The ammonia is then used as raw material for L-glutamate synthesis (Figure S5). Sulfur metabolism was not a major pathway in either the bacterial or the archaenal communities according to the predicted number of genes. However, assimilatory sulphate reduction was indicated in both the bacterial and archaenal communities, while dissimilatory sulphate reduction and sulphur oxidation was indicated only in the bacterial communities (Figure S6).

Several amino acid synthesis pathways were predicted (Figure 8), of which the most prominent were the alanine, aspartate and glutamate synthesis, arginine and proline synthesis, cysteine and methionine synthesis, glycine, serine and threonine synthesis, phenylalanine, tyrosine and tryptophan synthesis and the valine, leucine and isoleucine synthesis pathways.

Different types of membrane transport (ABC transporters) was identified where sulphate and iron (III) were taken up by the bacteria and tungstate, molybdate, proline, zink, cobalt and nickel was taken up by both archaea and bacteria (Figure S7). The estimated number of genes for both the purine and pyrimidine metabolism was more than two times higher in the archaenal community than in the bacterial community (Figure 8a and b).
4 Discussion

The phenotypic characteristics of the Fennoscandian Shield deep subsurface microbial communities are still largely unknown although specific reactions to introduced environmental stimulants have been shown (e.g. Pedersen et al., 2013; 2014; Rajala et al., 2015; Kutvonen 2015). Nevertheless, the connection of these microbial responses to specific microbial groups is still only in an early phase. Metagenomic and gene specific analyses of deep subsurface microbial communities have revealed prominent metabolic potential of the microbial communities, which appear to be associated with the prevailing lithology and physicochemical parameters (Nyyssönen et al., 2014; Purkamo et al., 2015). It has also been shown with fingerprinting methods with ever increasing efficiency that the bacterial and archaeal communities are highly diverse in the saline anaerobic Fennoscandian deep fracture zone groundwater (Bomberg et al., 2014; 2015; Nyyssönen et al., 2012; 2014; Pedersen et al., 2014; Miettinen et al, 2015; Sohlberg et al., 2015). Nevertheless, the concentration of microbial cells in the groundwater is quite low (Figure 2, Table 1). Most of the microbial communities at different depth in Olkiluoto bedrock fractures consist of bacteria. However, at specific depths (328 m, 423 m) the archaea may contribute with over 50% of the estimated 16S rRNA gene pool (Table 1). The major archaeal group present at these depths were the ANME-2D archaea indicating that nitrate-mediated anaerobic oxidation of methane may be especially common (Haroon et al., 2013). The high abundance of archaea in Olkiluoto is special for this environment. Archaea have also been quantified from the Outokumpu deep scientific borehole (Purkamo et al., 2016), but unlike the situation in Olkiluoto the archaeal community was less than 1% of the total community at best.

Previously, using 454 amplicon sequencing, we have observed OTU numbers of approximately 800 OTUs per sample covering approximately 550 bacterial genera (or equivalent groups) and approximately 350 archaeal OTUs including approximately 80 different genera (or equivalent groups) (Miettinen et al., 2015). Miettinen et al. (2015) defined the OTUs 97% sequence homology and the number of sequence reads per sample was at most in the range of 10^4. In contrast, our sequence read numbers were 10- to 100-fold higher and the number of OTUs per sample in general 100-fold higher. This indicates that a greater sequencing depth increases the number of taxa detected from the subsurface environment and allows us a novel view of the so far hidden rare biosphere. Nevertheless, in comparison to the high number of OTUs detected the number of identified genera, 651 and 81 bacterial and
archaeal genera, respectively, seems low. On the other hand, this indicates that the sequencing
depth has been sufficient to detect most of the prokaryotic groups present. Nevertheless, the
obtained numbers of OTUs per sample in this study were huge (Table 2). This may reflect the
high level of variability in the short sequence reads of the v6 region used in this study. As
discussed by Huse et al. (2008), short sequence reads very often match several different full-
length 16S rRNA reads. As shown in our study taxonomic assignments, such as
‘Proteobacteria_other’ were common and may be due to multiple matches for the individual
sequence reads obtained in the identification step of the analysis.

In general, the microbial communities at different depth grouped loosely into clusters
according to the groundwater chemistry (Figure 5). Salinity diverged the bacterial
communities of the two deepest samples (OL-KR44 and OL-KR29) from the rest of the
samples and sulphate, sulphur and sulphide moved the more shallow samples from depths
between 296 m and 347 m to the right of the NMDS plot. Sulphate reducers were not among
the most common bacterial taxa in these samples (Figure 3), but several sulphur and sulfide
oxidizing taxa were detected, such as the *Sulfuricurvum* and members of the *Thiobacterales.*
The archaeal communities were evenly distributed throughout the NMDS plot. The archaeal
communities did not change dramatically with depth and Euryarchaeota_Other, ANME-2D
and Thermoplasma_E2 groups dominated throughout the depth profile. Previous studies on
the Finnish deep biosphere has shown that the microbial communities at different sites vary
strongly from each other. Purkamo et al. (2015) investigated the bacterial and archaeal
communities of different fracture zones of the Outokumpu deep scientific borehole and found
that the majority of the bacterial populations at depths between 180 m and 500 m depth
consist of Betaproteobacteria belonging to the *Commamonadaceae* and the archaeal
communities consist of *Methanobacteriaceae* and *Methanoregula.*

The core communities, defined as taxa present in all the studied samples, accounted for
between 80 – 97% and 95 – > 99% of the bacterial and archaeal communities, respectively.
This is a considerable frequency of common microbial taxa. Nevertheless, the number of rare
taxa detected from the sample set was 3.3 to 6.8 fold higher than the number of core taxa on
genus level. Our results agree with Sogin et al. (2006) and Magnabosco et al. (2014), who
showed that a relatively small number of taxa dominate deep-sea water and deep groundwater
habitats, respectively, but a rare microbiome consisting of thousands of taxonomically distinct
microbial groups are detected at low abundances. What this means for the functioning of the
deep subsurface is that the microbial communities have the capacity to respond and change due to changes in environmental conditions. For example, Pedersen et al. (2014) showed that by adding sulphate to the sulphate-poor but methane-rich groundwater in Olkiluoto the bacterial population changed over the span of 103 days from a non-SRB community to a community dominated by SRB. In addition, a change in the geochemical environment induced by H₂ and methane impacted the size, composition and functions of the microbial community and ultimately led to acetate formation (Pedersen et al., 2012; Pedersen, 2013; Pedersen et al., 2014). This is also in accordance to the network analysis (Figure 6), which indicated a great diversity in the metabolic functions of the most central microbial taxa detected in this environment.

The metabolic pathways predicted by PICRUSt are far from certain when uncultured and unculturable deep subsurface microbial communities are concerned. The NSTI values for both the bacterial and well as the archaeal communities were high indicating that closely related species to those found in our deep groundwater have yet to be sequenced. This is in accordance with Langille et al. (2013), who showed that environments containing a high degree of unexplored microbiota also tend to have high NSTI values. Staley et al. (2014) also showed in a comparison between PICRUSt and shot gun metagenomic sequencing of riverine microbial communities that PICRUSt may not be able to correctly assess rare biosphere functions. Nevertheless, Langille et al. (2013) showed that PICRUSt may predict the metagenomic content of a microbial community more reliably than shallow metagenomic sequencing. Thus, on higher taxonomical level common traits for specific groups of microorganisms may be revealed.

**Energy metabolism.** Deep subsurface environments are often declared energy deprived environments dominated by autotrophic microorganisms (Hoehler and Jorgensen, 2013). However, recent reports indicate that heterotrophic microorganisms play a greater role than the autotrophic microorganisms in Fennoscandian deep crystalline subsurface environments (Purkamo et al., 2015). Heterotrophic communities with rich fatty acid assimilation strategies have been reported to fix carbon dioxide on the side of e.g. fermenting activities in order to replenish the intracellular carbon pool, which otherwise would be depleted. Wu et al. (2015) also found by metagenomic analyses that fermentation was a major metabolic activity in the microbial community of Swedish deep groundwater. Our results agree with Purkamo et al. (2015) that a greater proportion of the microbial community is involved in carbohydrate and
fatty and organic acid oxidation than in fixation of inorganic carbon. Nevertheless, autotrophic carbon fixation pathways were predicted in the analysis with PICRUSt, indicating that both the archaeal and bacterial communities include autotrophic members, although these microorganisms might not be obligate autotrophs. It is also likely that heterotrophic and chemolithotrophic microorganisms coexist in the Olkiluoto deep fracture zones forming networks as shown in Figure 6 for the benefit of the whole microbial community. Such cooccurrences have been suggested by e.g. Osburn et al. (2014). It was also noted that even though evidence for methane oxidation could not be inferred from the PICRUSt predictions (no pmoA genes), the bacterial community may oxidize formate, which is in agreement with the findings reported by Wu et al. (2015).

Several carbon fixation pathways were predicted in the metagenomes, the Calvin cycle, reductive TCA (rTCA) cycle and Wood-Ljungdahl (WL) pathway. The WL-pathway is considered the most ancient autotrophic carbon fixation pathway in bacteria and archaea (Fuchs 1989, Martin et al. 2008, Berg et al. 2010; Hügler and Sievert, 2011) and was found in both the bacterial and the archaeal communities. In the archaeal community the Calvin cycle and the rTCA were especially pronounced in the samples from 296 m, 405 – 423 m and somewhat lower at 510 – 527 m depth. The bacterial communities are predicted to fix CO₂ at almost all depths with the exception of 405 m and 559 m depth. Nevertheless, our results agree with Nyyssönen et al. (2014), who showed my metagenomic analysis that the microbial communities at different depth of the Outokumpu scientific deep drill hole may fix carbon in several ways, of which the rTCA, the WL pathway and the Calvin cycle were identified. Magnabosco et al. (2016) showed that the WL pathway was the dominating form of carbon fixation in metagenomes of 3 km deep Precambrian crust biospheres in South Africa. Dong et al. (2014) also suggested that microorganisms in low-energy deep subsurface environment may have several strategies for e.g. carbon fixation, as shown in the Halomonas sulfidae, in order to access as many resources as possible. The predicted methane metabolism (methane and methyl compound consumption) and oxidative phosphorylation were equally strong in the bacterial community. Sulphur metabolism was not a common pathway for energy in either the archaeal or the bacterial communities, but bacteria with either assimilative or dissimilative sulphate reduction were present. Sulphur oxidation through the sox system was in general not predicted, but the soxD gene was predicted and oxidation of thiosulphate to sulphate may be possible (Figure S6). Nitrate is reduced both through dissimilatory nitrate reduction to ammonia and through denitrification to nitrous oxide by the bacteria. In addition, nitrogen is
fixed to ammonia by both archaea and bacteria. The ammonia is then used as raw material for L-glutamate synthesis.

Oxidative phosphorylation was one of the most prominent energy generating metabolic pathways in the bacterial community. This indicates that ATP is generated by electron transfer to a terminal electron acceptor, such as oxygen, nitrate or sulphate. In the archaeal community the oxidative phosphorylation was not as strongly indicated, but this may be due to missing data on archaeal metabolism in the KEGG database.

The main energy metabolism of the archaeal communities appeared to be the methanogenesis, especially at 296 m and 405 m. Methanogenesis was common also at all other depths except 330 m – 347 m, 415 m and 693 m – 798 m. Methane is produced from CO₂-H₂ and methanol, and from acetate, although evidence for the acetate kinase enzyme was lacking. Methanogenesis from methylamines may also be possible, especially at 296 m and 405 m. Methane oxidation using methane monoxygenases and methanol dehydrogenases does not occur in either bacterial or archaeal communities.

**Carbohydrate metabolism.** Glycolysis/glucogenesis is one of the most common carbohydrate-metabolizing pathways predicted for both the archaeal and bacterial communities (Figure 9). Pyruvate from glycolysis is oxidized to acetyl-CoA by both archaea and bacteria and used in the TCA cycle. The TCA cycle provides for example raw material for many amino acids, such as lysine and glutamate. The butanoate and propanoate metabolisms were also common in the bacterial communities, indicating fermentative metabolism and capability of fatty acid oxidation.

**Amino acid metabolism.** Non-essential amino acids, such as alanine, aspartate and glutamate are produced from ammonia and pyruvate or oxaloacetate especially in the archaeal populations. In the archaeal population proline appears to be produced from glutamate. Despite the low use of sulphate as energy source in the microbial communities sulphate and other sulphur compounds are taken up for the production of the amino acids cysteine and methionine by both the archaeal and the bacterial communities. A higher predicted relative abundance of genes involved in aromatic amino acid synthesis (phenylalanine, tyrosine, tryptophane) was seen in the archaeal than in the bacterial communities. Both the archaeal and the bacterial communities synthesise branched chained amino acids (isoleucine, leucine and valine), but only the bacteria degrade them. Especially proteobacteria have been shown to be able to use the branched chained amino acids (isoleucine, leucine and valine) and short
chained fatty acids (acetate, butyrate, propionate) as sole energy and carbon source (Kazakov et al., 2009). The branched chained amino acids function as raw material in the biosynthesis of branched chained fatty acids, which regulate the membrane fluidity of the bacterial cell. In salt stress conditions, the proportion of branch-chained fatty acids in the membranes decreases.

Membrane transport. According to the predicted metagenomes, the microbial cells transport sulphate into the cell, but do not take up nitrate. Nitrogen is taken up as glutamate but not as urea. Iron is taken up by an Fe(III) transport system and an iron complex transport system in the bacterial communities, but generally only by the iron complex transport system in archaea. However, Fe(III) transport system may also exist in the archaean communities at 405 m to 423 m depth, where also some manganese/iron transport systems could be found. Molybdate and phosphate is transported into the cell by molybdate and phosphate ATPases, respectively. Nickel is taken up mainly by a nickel/peptide transport system but also to some extent by a cobalt/nickel transport system. Zink is taken up to some extent by a zink transport system, but transport systems for manganese, manganese/iron, manganese/zink/iron, or iron/zink/copper are negligent. Ammonia is taken up by an Amt transport system.

5 Conclusions

The wide diversity of microbial groups in the deep Fennoscandian groundwater at the Olkiluoto site revealed that the majority of the microbial community present belong to only a few microbial taxa while the greatest part of the microbial diversity is represented by low abundance and rare microbiome taxa. The core community was present in all tested samples from different depths, but the relative abundance of the different taxa varied in the different samples. Specific rare microbial groups formed tight co-occurrence clusters that corresponded to different environmental conditions and these may become more abundant if the environmental conditions change. Fermentation or oxidation of fatty acids was a common carbon cycling and energy harvesting metabolic pathways in the bacterial communities whereas the archaea may either produce or consume methane. Glycolysis/gluconeogenesis was predicted to be common in both the archaean and bacterial communities. In addition both the bacterial and archaean communities were estimated to contain different common carbon fixation pathways, such as the Calvin cycle and the reductive TCA, while only the bacteria contained the Wood-Ljungdahl pathway.
Acknowledgements

The Illumina sequencing data were made possible by the Deep Carbon Observatory’s Census of Deep Life supported by the Alfred P. Sloan Foundation. Illumina sequencing was performed at the Marine Biological Laboratory (Woods Hole, MA, USA) and we are grateful for the assistance of Mitch Sogin, Susan Huse, Joseph Vineis, Andrew Voorhis, Sharon Grim, and Hilary Morrison at MBL and Rick Colwell, OSU. MB was supported by the Academy of Finland (project 261220).
References


Kutvonen, H.: Nitrogen-cycling bacteria in groundwater of the low and medium active nuclear waste repository in Olkiluoto, Finland (In Finnish) M.Sc. Thesis, Faculty of Agriculture and Forestry, University of Helsinki, Finland, 2015.


10 Wu, X., Holmfeldt, K., Hubalek, V., Lundin, D., Åström, M., Bertilsson, S. and Dopson, M. Microbial metagenomes from three aquifers in the Fennoscandian shield terrestrial deep biosphere reveal metabolic partitioning among populations. ISME J., on line.
1. Table 1. Geochemical and microbiological measurements from 12 different water conductive fractures in the bedrock of Olkiluoto, Finland.

The different drillholes are presented at the top of the table. The data is compiled from Posiva (2013) and Miettinen et al. (2015)

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Table 2a. The total number of sequence reads, observed and estimated (Chao1, ACE) number of OTUs, number of singleton and doubleton OTUs, and Shannon diversity index per sample of the bacterial 16S rRNA gene data set. The analysis results are presented for both the total number of sequence reads per sample as well as for data normalized according to the sample with the lowest number of sequence reads, i.e. 140,000 random sequences per sample.

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Number of sequence reads: 17,000 random sequences per sample.
Figure 1. Map of Olkiluoto. The boreholes used in this study are marked with a turquoise triangle and the attached black line depicts the direction of the borehole. (with courtesy of Pöyry Oy, Nov 17th, 2015 by Eemeli Hurmerinta)

Figure 2. The concentration of A) microbial cells mL\(^{-1}\) determined by epifluorescence microscopy and the estimated concentration of B) bacterial and C) archaeal 16S rRNA gene copies mL\(^{-1}\) groundwater determined by qPCR in water conductive fractures situated at different depths in the Olkiluoto bedrock.
Figure 3. The most abundant A) bacterial and B) archaeal taxa representing at least 1% of the sequence reads in any of the samples. The number in each series indicate the taxon number in the list below the figures.
Figure 4. A Bray-Curtis UPGMA cladogram clustering the studied samples according to the detected taxonomy of the microbial communities. The bacterial and archaeal community profiles were combined. The branch support values were calculated from 100 bootstrap repeats.

Figure 5. Non-metric multidimensional scaling analysis based on the A) bacterial and B) archaeal communities detected in the samples. The triplot (green) indicates directionality of the environmental variables.
Figure 6. Network of co-occurring microbial taxa based on Spearman’s rank correlation values between pairs of taxa. Correlation (R>0.7, p<0.01) between different taxa. Each circle (node) represents a taxon and the size of the node is proportional to the number of connections (Spearman correlation value) of the node. The colour of the nodes indicates degree of centrality of the taxon, with low centrality shown as blue, increasing centrality by green to yellow to orange and highest centrality as red. Taxa with less than 10% centrality range (<2 connections) were filtered out. The most prominent nodes are indicated by taxon names. In Figure S2 the names for all nodes are shown.
Figure 7. The nearest sequenced taxon index (NSTI) values calculated by PICRUSt for the bacterial (blue) and archaeal (red) communities. The NSTI value describes the sum of phylogenetic distances of each OTU to its nearest relative with a sequenced reference genome, and measures substitutions per site in the 16S rRNA gene and the weighted the frequency of the each OTU in a sample dataset. A higher NSTI value indicates greater distance to the closest sequenced relatives of the OTUs in each sample.
Figure 8. The relative abundance of predicted genes of the most abundant pathways identified in the A) bacterial and B) archaeal populations in the PICRUSt analysis. The pathways are presented according to KEGG. The samples are ordered according to depth, with OL-KR13/296m as innermost and OL-KR29/798m as the outermost sample.
Figure 9. The microbial carbon metabolism pathway according to KEGG. The predicted genes combined from all samples were plotted on the map. Green arrows indicate enzymes predicted only in the archaeal communities, red arrows indicate genes predicted only in the bacterial communities, black arrows show enzymes predicted in both the archaeal and bacterial communities and blue arrows show enzymes that were not predicted in any of the communities.
Supplementary figures

Figure S1. The rarefaction curves of observed bacterial (left pane) and archaeal (right pane) OTUs in each sample generated on sequence data normalized to 140,000 reads for bacteria and 17,000 reads for archaea.
Figure S2. Network of co-occurring microbial taxa based on Spearman’s rank correlation values between pairs of taxa. Correlation (R>0.7, p<0.01) between different taxa. Each circle (node) represents a taxon and the size of the node is proportional to the number of connections (Spearman correlation value) of the node. The colour of the nodes indicates degree of centrality of the taxon, with low centrality shown as blue, increasing centrality by green to yellow to orange and highest centrality as red. Taxa with less than 10% centrality range (<2 connections) were filtered out. The names for all taxa included in the analysis are shown.
Figure S3. Bacterial oxidative phosphorylation according to KEGG. The predicted genes from the bacterial communities belonging to the oxidative phosphorylation are shown in pink.
Figure S4. Predicted genes shown in red of the bacterial A) fatty acid degradation and B) fatty acid biosynthesis pathways, combined from all samples.
Figure S5. The predicted genes of enzymes included in the microbial nitrogen metabolism according to KEGG. Enzymes predicted from the bacterial communities are shown in red, archaeal communities in green and enzymes predicted from both archaeal and bacterial communities in green/red. Enzymes not predicted from either community are shown in blue or white.
Figure S6. The genes of ABC transporters predicted from the bacterial (pink), archaeal (green) or both (pink/green) communities. Genes not predicted in any of the communities are shown in blue.
Figure S7. The predicted genes of enzymes included in the microbial sulphur metabolism according to KEGG. Enzymes predicted from the bacterial communities are shown in red, archaeal communities in green and enzymes predicted from both archaeal and bacterial communities in green/red. Enzymes not predicted from either community are shown in blue or white.