RESEARCH ARTICLE

Microbial diversity and mineral composition of weathered serpentine rock of the Khalilovsky massif

Irina V. Khilyas1*, Alyona V. Sorokina1*, Anna A. Elistratova1*, Maria I. Markelova1, Maria N. Siniagina1, Margarita R. Sharipova1, Tatyana A. Shcherbakova2, Megan E. D’Errico3, Michael F. Cohen4

1 Institute of Fundamental Medicine and Biology, Kazan (Volga Region) Federal University, Kazan, Russian Federation, 2 FSUE Central Research Institute of Geology of Non-metallic Mineral Resources, Kazan, Russian Federation, 3 School of Science and Technology, Sonoma State University, Rohnert Park, CA, United States of America, 4 Department of Biology, Sonoma State University, Rohnert Park, CA, United States of America

* These authors contributed equally to this work.
* irina.khilyas@gmail.com

Abstract

Endolithic microbial communities survive nutrient and energy deficient conditions while contributing to the weathering of their mineral substrate. This study examined the mineral composition and microbial communities of fully serpentinized weathered rock from 0.1 to 6.5 m depth at a site within the Khalilovsky massif, Orenburg Region, Southern Ural Mountains, Russia. The mineral composition includes a major content of serpentinite family (mostly consisting of lizardite and chrysotile), magnesium hydrocarbarnates (hydromagnesite with lesser amounts of hydrotalcite and pyroaurite) concentrated in the upper layers, and clay minerals. We found that the deep-seated weathered serpentinites are chrysotile-type minerals, while the middle and surface serpentinite mostly consist of lizardite and chrysotile types. Microbial community analysis, based on 16S rRNA gene sequencing, showed a similar diversity of phyla throughout the depth profile. The dominant bacterial phyla were the Actinobacteria (of which unclassified genera in the orders Acidimicrobiales and Actinomycetales were most numerous), Chloroflexi (dominated by an uncultured P2-11E order) and the Proteobacteria (predominantly class Betaproteobacteria). Densities of several groups of bacteria were negatively correlated with depth. Occurrence of the orders Actinomycetales, Gaiellales, Solirubrobacterales, Rhizobiales and Burkholderiales were positively correlated with depth. Our findings show that endolithic microbial communities of the Khalilovsky massif have similar diversity to those of serpentine soils and rocks, but are substantially different from those of the aqueous environments of actively serpentinizing systems.

Introduction

Microbial communities colonizing mineral surfaces are of significant evolutionary and ecological interest. Mineral surfaces are extreme eoniches, having microbiota affected by a large
number of exogenous factors such as temperature, light, salinity, high pH, deficiency and unavailability of nutrients [1]. The mineral surface microtopography, structure, chemistry and reactivity are selective factors for microbial colonization, formation of biofilms and ultimately for the whole microbial community structure. Mineral-associated microorganisms often contribute to geochemical processes including mineral formation, dissolution or deterioration [2, 3].

One enigmatic mineral considered as a potential locus for the origin of life on Earth is serpentinite [4]. Serpentinite comprising of the antigorite, lizardite, and chrysotile group minerals is an ultramafic rock mineral that occurs in all ancient orogenic belts [5] within the oceanic lithosphere at surfaces and faults [6, 7]. Formation of serpentinite occurs at relatively low temperatures through hydration of olivine-rich ultramafic rocks to serpentine, hydrogen, methane, low-molecular organic compounds [8] and, under conditions of sufficient SiO$_2$, magnetite (Fe$_3$O$_4$) [9].

In natural environments, terrestrial serpentinites (Mg,Fe,Ni,Mn,Zn)$_{2-3}$(Si,Al,Fe)$_2$O$_5$(OH)$_4$ undergo a weathering process accompanied by consistent physical and chemical changes of minerals through dissolution, decomposition and carbonation. Hydrolysis of serpentinites is accompanied by release of Mg$^{2+}$, HCO$_3^-$ and SiO$_2$, which is followed by formation of magnesium hydrocarbonates and amorphous silica oxide [10]. Serpentinite weathering is associated with increasing of alkalinity of deep-seated minerals within the geologic formation [11, 12]. We hypothesized that such changes in serpentine rocks impact the resident microbiota, which in turn could have a role in the development of serpentinite-derived infertile soils.

The Khalilovsky massif is located near the town of Novorudnyy, Gaysky District, Orenburg Region, Southern Ural Mountains, Russia (S1A and S1B Fig). The approximately 260 km$^2$ massif was deposited in the middle Paleozoic era bordered by the Uraltausky anticlinorium and greenstone synclinorium [13]. The main rock types within the massif are serpentinized lizardite and chrysotile (95–98%) with lesser amounts of clay minerals, chlorite, dolomite, calcite and magnesite [13]. The serpentine associated with the ophiolite complexes in this region are derived from harzburgite and dunite, reflecting a more melt depleted oceanic mantle [14]. Presently, the massif is characterized by rolling hills covered in some places by 1–5 cm layer of soil with grass and streams running through the deposit (S1C Fig). One of the most significant waterways of the region is the Guberlya River. The Southern Ural Mountains have an arid continental climate. The average temperature for the coldest (January) and the warmest (July) months of the year is –20˚C and +40˚C, respectively. The deposit receives an average of 181 mm of precipitation per year with a mean annual evaporation rate of 988 mm.

The purposes of this study were to (1) characterize the microbial community diversity of the fully serpentinized weathered Khalilovsky massif subsurface samples and compare them with those reported for other serpentinite-hosted systems, and (2) to perform geochemical and mineralogical analyses and determine the role, if any, of the resident microbial community in the weathering process. The results presented here show that bacterial diversity, as indicated by 16S rRNA gene sequence analysis, did not vary significantly at the phyla level with depth within our sampling site, although differences in the dominant microbial orders and genera were found. We also demonstrate that the microbial community does not appear to substantially alter its serpentine mineral substrate.
overlay, located in the central part of the Khalilovsky massif (Latitude: N 51°30' Longitude: E 58°07') in August 2017. Two duplicate samples (55.0–60.0 g) taken from opposite sides of the core were collected at the depths specified in Table 1 into sterile plastic boxes. Each sample was crushed under sterile laboratory conditions using a mortar and pestle. The mineral particle size for analysis was approximately 0.01 mm. The pH of water extracted samples ranged between pH 8.98 to pH 9.06. For the collection of serpentine from the Khalilovky massif a specific permission is not required. The Khalilovsky massif is not related to a national park or other protected areas of land of the Russian Federation.

### X-ray diffraction (XRD)

Analyses of the mineralogical samples were performed on powdered samples using a D8 Advance XRD2 Bruker diffractometer, with CuK-alpha radiation, at 40 kV, 30 mA. X-ray diffractograms were collected within the 2θ range (5–40°) with 0.05/1 sec step. Mineralogical composition was determined with advance diffract plus evaluation software (DIFFRACplus-TOPAS Software package).

### Scanning electron microscopy (SEM)

The mineralogical samples were coated with a conductive layer of Au/Pd alloy at an 80/20 ratio using a cathode sputtering method in the Quorum Q150T ES vacuum chamber. The thickness of the conductive layer was 15 nm. Measurements were conducted using a high-resolution field-emission scanning electron microscope Merlin (Carl Zeiss). Imaging of the surface morphology was carried out with 5 kV acceleration voltage of the primary electrons and 300 pA probe current for minimal impact on the object of research.

Elemental analysis was conducted using an energy dispersive spectrometer (EDS) AZtec X-Max (resolution 127 eV) under 20 kV acceleration voltages of the electron probe and 10 mm working distance in order to avoid minimal errors in the microprobe analysis. The probing depth was approximately 1 μm.

### Transmission electron microscopy (TEM)

Analysis of samples was carried out using a Hitachi HT7700 Exalens transmission electron microscope. To prepare the sample 5 μl of the mineralogical suspension was placed on a Formvar/carbon 3 mm coated copper grid and dried at room temperature. After drying, the grid

### Table 1. Sample X-Ray diffraction mineral identification and percent composition in the serpentine rock core taken at the Khalilovsky massif*

| Sample depth (m) | Serpentinite (%) | Hydrotalcite-type and pyroaurite group minerals (%) | Hydromagnesite (%) | Clay minerals (%) | Calcite (%) |
|-----------------|-----------------|-----------------------------|-------------------|-----------------|------------|
| 0.1             | 72.3 ±2.5       | 10.8 ±1                     | 15.1 ±1           | 1.8 ±0.5        | -          |
| 0.85            | 82.1 ±2.0       | 6.7 ±0.7                    | 9.3 ±0.8          | 2.0 ±0.5        | -          |
| 1.6             | 79.0 ±2.1       | 9.6 ±0.8                    | 9.2 ±0.8          | 2.2 ±0.5        | -          |
| 2.35            | 72.1 ±2.5       | 12.2 ±1                     | 15.5 ±1           | 1.2 ±0.5        | -          |
| 3.1             | 84.0 ±1.6       | 7.8 ±0.8                    | 8.2 ±0.8          | -               | -          |
| 3.85            | 81.4 ±2.1       | 11.3 ±1                     | 4.9 ±0.6          | 2.5 ±0.5        | -          |
| 4.6             | 81.6 ±2.2       | 9.8±1                       | 7.3 ±0.7          | 1.3 ±0.5        | -          |
| 5.35            | 87.5 ±2.0       | 6.6 ±0.7                    | 3.7 ±0.6          | 1.5 ±0.5        | 0.8±0.2    |
| 6.1             | 89.0 ±1.7       | 5.6 ±0.6                    | 4.0 ±0.6          | 1.4 ±0.5        | -          |
| 6.85            | 93.5 ±1.1       | 4.1 ±0.6                    | -                 | 2.3 ±0.5        | -          |

*Values are the mean ± range of duplicate samples of the core taken at a given depth.

https://doi.org/10.1371/journal.pone.0225929.t001
was transferred into the transmission electron microscope. Analysis was conducted at an accelerating voltage of 100kV in TEM mode.

**DNA extraction, 16S rRNA gene amplification and sequencing**

To extract DNA 10 g of crushed rock from each depth sample was submersed in 5 - 10 mL DNase-RNase-free water for one week at room temperature. The mineral suspension was subjected to three liquid N₂ freeze/thaw cycles[15], lysozyme/Proteinase K treatment, followed by a phenol-chloroform extraction step. DNA washing and precipitation was performed using 80% cold ethanol. DNA was resuspended in the DNase-RNase-free water[15]. The quantity of the extracted DNA was measured using a Qubit 2.0 Fluorometer (Invitrogen, Carlsbad, CA).

Bacterial 16S rRNA gene amplicons (~ 500-bp fragments) were produced using the 337F (5’-CCTACGGGNGGCWGCAG-3’) and 805R (5’-GACTACHVGGGTATCTAATCC-3’) primers[16]. Archaeal 16S rRNA gene fragments were amplified with the forward primer 344F (5’-ACGGGYGCAGCAGGCGCGA-3’) and the reverse primer 806R (5’-GGACTACVSGGGTATCTAAT-3’)[17]. Reaction products were purified using Ampure XP paramagnetic beads (Beckman Coulter Inc., USA). The purified PCR products were amplified with Illumina primers containing the unique barcodes. The pooled multiplex reactions were loaded on an Illumina MiSeq instrument with the 2x300 v3 reagent mix.

**Analysis of the specific functional genes in the serpentinite-associated microbial communities**

The mcrA gene encoding methyl-coenzyme M reductase (MCR) and hydA gene encoding hydrogenase were used to investigate the potential methanogenic activity and hydrogen oxidation in the mineralogical samples using specific primers mcrA-F (5’-GGTGGTGTMGGATTCA CACARTAYGCWACAGC-3’) and mcrA-R (5’-TTTATTGCRTAGTTWGGRAGT-3’), and FeFe27F (5’-GCHGAYMTBACHATWATGARAGA-3’) and FeFe27R (5’-GCNGCYTCCCATDA CDDCDCNGT-3’), respectively[18, 19].

**Bioinformatics analysis**

Sequencing of 16S rRNA gene V3-V4 variable regions was performed on the Illumina MiSeq platform in 2x300bp mode. Reads were further processed and analyzed using the QIIME software, version 1.9.1 (http://qiime.org/) [20] according to protocols. Before filtering, there were 66287 read pairs per sample on average. Paired-end reads were initially merged and then processed to remove low quality and chimeric sequence data. The rarefaction step was performed to reduce sequencing depth heterogeneity between samples. After quality filtering, chimera filtering and rarefying, we analyzed 24442 joined read pairs. Sequences were clustered into operational taxonomic units (OTU) based on the 97% identity threshold (open reference-based OTU picking strategy), the latest SILVA database v.132, 13-12-17 [21] was used. Each OTU was required to contain at least 2 sequences for inclusion in the final OTU list. The number of observed OTUs varied from 1702 to 3578. To characterize the richness and evenness of the bacterial community, alpha diversity indices were calculated using Chao1, Shannon and Simpson metrics (S2 Table). Similarities between microbial compositions of samples were evaluated using the beta diversity characteristics, which were estimated using weighted and unweighted Unifrac measures with further non-metric multidimensional scaling (nMDS) visualization.

Raw reads are deposited in the ENA under Project ID PRJEB29885 in the fastq format (https://www.ebi.ac.uk/ena/data/view/PRJEB29885).
Results and discussion

Mineral morphology and geochemical composition

Rocks containing the serpentine group of minerals are present in all orogenic belts worldwide, typically forming large massifs along continental margins and mélangé, faults and shear zones. The geochemical, mineralogical and microbiological features of serpentinites attract much interest among researchers in the field of geology, mineralogy, geochemistry and microbiology. The process of weathering results in serpentinites of different mineral maturities. Mineral rocks of the Khalilovsky massif are represented by massive and coarsely fractured lizardite and chrysotile with small amounts of bastite [13]. We observed two distinct serpentine zones of the massif: (1) the soil-grass serpentine ecosystem; and (2) the solid rocky nutrient poor serpentine ecosystem from which we obtained our core (S1 Fig).

We collected duplicate samples at given depths from 0.1 m to 6.85 m from a single core drilled at a site presently undergoing erosion and weathering (Table 1). This weathered serpentine has a distinct greenish grey color and fine to medium-grained texture, which is distinct from rocky serpentinites that are characterized by a dark greenish grey color and dense homogenization (S2 Fig). The mineralogical composition of the rock core samples was evaluated by X-ray diffraction and found to be represented by serpentinite minerals and hydromagnesite, with a minority of hydrotalcite, pyroaurite, clay minerals and calcite (Table 1, S3 Fig). Magnesium hydrocarbonate minerals (hydrotalcite-type and pyroaurite group minerals and hydromagnesite) are known to accrete in veins and fissures formed through surface weathering[22]. Consistent with this, the content of these minerals decreased in depth from 0.1 m to 6.85 m and, conversely, the proportion of serpentinite group minerals increased in the deep-seated layers. Calcium carbonates (e.g. calcite) were negligible or completely absent throughout.

Serpentine rocks include a range of major minerals, such as lizardite, chrysotile, antigorite and other minorities characterized by different texture and structure[12]. Scanning electron microscopy (SEM) and energy dispersive X-ray spectrometry (EDS) was used to characterize the surface morphology and determine the chemical phases of serpentinites collected from different depths. A plate-like morphology, characteristic of lizardite[23], with a minor inclusion of cylindrical fibers was observed in serpentinite collected from 0.1 m depth (Fig 1A). Plates and unoriented fibers, often arranged in parallel strands, were characteristic of the serpentinite collected from the 3.1 m depth (Fig 1B). The deep-seated serpentinites (6.85 m in depth) displayed well oriented, needle-shaped hollow fibers, or nanotubes, that are characteristic of chrysotile [23] (Fig 1C). The sizes of the particles observed in this study ranged from less than 1 μm to greater than 200 μm. As determined by EDS, oxygen, magnesium, silicon and iron were the predominant elements in the mineralogical samples (S4 Fig). Additionally, the transition metals nickel and chromium were detected in the upper layers of the serpentine deposit, only nickel was found in the middle layers, but both elements were absent in the deep-seated serpentinites (S4 Fig). Transmission electron microscopy (TEM) was applied to investigate the crystal structure of serpentinites. The irregular shaped plates and hollow cylinders with different thicknesses were typical for all serpentinites (S5 Fig).

Microbial community composition

We characterized the microbial diversity of the weathered serpentine minerals sampled from the core obtained from a solid rocky zone of the Khalilovsky massif. The composition of microbial communities was similar throughout the 0.1 m to 6.85 m depth profile at the phylum/class level. Among the 31 different phyla found, Actinobacteria, Proteobacteria and Chloroflexi were the most abundant phyla in all of the mineralogical samples (Fig 2).
The dominant phylum *Actinobacteria* throughout the core depths was represented primarily by an unclassified genus within the order *Acidimicrobiales*, comprising 11.0% to 38.0% of all OTU sequences, and members of the order *Actinomycetales* (*Nocardioidaceae* family), comprising 9.9% to 21.5% of OTUs (Fig 3, S3 Table). The phenotypic basis for the success of these particular groups within this habitat remains to be determined. The phylum *Proteobacteria* was represented mainly by the classes *Betaproteobacteria* (*Comamonadaceae*, *Oxalobacteraceae* and *Delftia*), *Gammaproteobacteria* (*Enterobacteriaceae* family, *Serratia* genus), and *Deltaproteobacteria* (*Myxococcales* order, *Haliangiaceae* family). Members of the genus *Delftia* are known to be distributed in the microbial communities of a deep subsurface thermal aquifer system (Siberia, Russia), a deep subsurface crystalline rock the Pyhäsalmi mine (Finland), and ornamental limestone from buildings [33–35]. Among the phylum *Chloroflexi* there was an abundance of an uncultured P2-11E order of bacteria.

**Microbial community composition correlating with depth**

Non-metric Bray-Curtis dissimilarity analysis (at a 97% similarity level) showed that microbial communities of the serpentine core samples collected from the different depths were closely related (Fig 4). Similar results were obtained using PCoA analysis, which explained 37.16% of the observed variation (Fig 4B).

Mineral porosity and water circulation are expected to be important factors influencing the microbial community of the Khalilovsky massif. Weathering-induced cracking at shallower depths as well as the fibrous tubular-type serpentinite nanostructures contribute to rock porosity [12] that should in turn positively impact serpentine-associated microflora since microbial colonization can only occur within fissures of rocks. Statistically significant correlations of the OTU abundances from the rock core relative to their distribution according to depth are presented in Table S4 Table. The orders *Acidimicrobiales*, *Gaiellales*, *Solirubrobacterales*, *Gemmatimonadetes*, *Rhizobiales*, *Burkholderiales*, P2-11E and unidentified members related to the GAL15 phylum positively correlated with depth. Additionally, several orders show a negative correlation with depth. Precipitation of hydromagnesite from magnesium-rich alkaline water percolating through fissures can occur without microbial mediation [36] but extracellular polymeric substances of adherent biofilms can accelerate such precipitation[37]. Whether the microbes residing in the serpentine rock fissures have contributed to the hydromagnesite deposition cannot by ascertained from our results.

**Comparison to other serpentine-related microbial communities**

To date, investigations have largely focused on microbial communities hosted in aqueous actively serpentinizing systems, enriched hydrogen and methane [24, 26–32]. However, relatively limited publications have reported on microbial communities associated with fully serpentinized rocks. Our results indicate that the microbial community of the mature serpentine Khalilovsky massif are more similar to serpentine mineral surface-associated (Fig 2A) than aqueous phase-associated microbial communities (Fig 2B & 2C).

These findings are consistent with those of Daae et al. [25] who found the groundwater microbial communities to be substantially different from the fracture-coating microbial communities within partially serpentinized dunite from the Leka ophiolite, Norway (Fig 2A & 2B). The *Actinobacteria* phylum, similar to our findings, predominated in the mineral fractions.
Fig 2. Comparative analysis of mineral- and aqueous-associated serpentine-hosted bacterial communities according to Phylum. (A) Community profiles of the Khalilovsky massif compared to those associated with serpentine minerals in other terrestrial environments. (B, C) Profiles of selected aqueous-associated planktonic communities from (B) terrestrial environments and (C) seawater serpentine- and ultramafic-hosted systems [24–32].

https://doi.org/10.1371/journal.pone.0225929.g002
Fig 3. A heat map of the bacterial genera found in serpentine rock core samples collected at different depths. Colors indicate percent relative abundance of 74 genus level taxonomic groupings (i.e. 97% OTU sequence identity) that occur within the major phyla (each representing \textasciitilde 0.5\% of all OTUs) detected at different depths. Names of the most highly represented genera are given.

https://doi.org/10.1371/journal.pone.0225929.g003
taken from 15 and 160 cm in depth, whereas in the fluid passing through minerals were dominated by members of Betaproteobacteria.

Some progress has been made in characterizing microbial communities associated with serpentinite-derived soils. A study of bacterial community structure by 16S rRNA gene clone library analysis found the dominant bacterial phyla in serpentine soil to be Actinobacteria, Acidobacteria, Alphaproteobacteria, Verrucomicrobia, Green-nonsulfur-related bacteria and Gemmatimonadetes [28], whose members are known to be well adapted to low moisture soils. We found Gemmatimonadetes consistently at frequencies ranging up to 6% in all of our mineral samples from the Khalilovsky massif (S3 Table).

Endolithic microbial communities of the heavily serpentinized open-pit chrysotile asbestos Jeffrey mine (Québec, Canada) were dominated by Proteobacteria, Cyanobacteria and Planctomycetes [29].

Analysis of ultramafic serpentinite soils including harsh and lush serpentines of the McLaughlin Reserve in the Northern Coast Ranges of California revealed that Proteobacteria and Actinobacteria were two abundant phyla in the microbial communities of minerals taken from a depth of 7.5 cm [26]. Among Actinobacteria, the orders Acidimicrobiales and Gaiellales were relatively abundant.

Firmicutes, a ubiquitous bacterial phylum inhabiting a wide range of ecosystems, is more highly represented in actively serpentinizing systems. Within the deep-seated weathered serpentinite of the arid Khalilovsky massif, the capacity to survive desiccation conferred by endospores may be responsible for the status of Bacillus as the dominant genus for this phylum (S3 Table). Acceleration of serpentinite dissolution by a Bacillus species under laboratory conditions was postulated to occur through the action of secreted organic acids and ligands [38].

Fig 4. Clustering analyses performed based on Bray-Curtis dissimilarity distances of prokaryotic OTU sequence abundance. (A) A hierarchical cluster tree; and (B) principal coordinate analysis (PCoA) scatter plot. PCoA1 and PCoA2 explained 25.7% and 11.46% of the observed variation. Black and grey-colored circles indicate the depth of the microbial communities.

https://doi.org/10.1371/journal.pone.0225929.g004
Thus, *Bacillus* as well as other bacteria that secrete such compounds could have a role in initiating and widening fissures within serpentine rock, leading to soil formation.

Similar to our findings at the Khalilovsky massif, Archaea were not found on the rocks from the Jeffery mine [29] and are proportionally less represented on mineral surface communities within other serpentinized systems [25, 26] relative to aqueous-associated communities of actively serpentinizing fields [8, 24, 31, 32, 39–41]. Consistent with the lack of Archaea, we did not detect the methyl coenzyme-M reductase (*mcrA*) gene, which is a common indicator of methane production within actively serpentinizing environments [8]. Similarly, we were not able to amplify from our samples the *hydA* gene, a marker of bacterial hydrogen metabolism in actively serpentinizing environments [8, 27, 42]. Furthermore, the H$_2$-oxidizing genus *Hydrogenophaga* did not exceed 0.26% of the sequences in any given core sample (S3 Table), whereas they reached up to 58% of the total bacterial community in samples collected from a chimney of the PHF[27].

**Conclusions**

The results of this study demonstrate the texture, microstructure, geochemical composition and microbial diversity within nutrient-poor, fully serpentinized weathered rock found at the Khalilovsky massif, Russia. SEM analysis revealed plate and fibrous tubular structures characteristic of lizardite and chrysotile, respectively, of the serpentinites collected from different depths. Geochemical analysis showed increasing serpentine-type mineral content and decreasing content of magnesium hydrocarbonates with depth, indicative of weathering-induced fissuring closer to the surface. 16S rRNA phylogenetic analysis of the endolithic microbial communities demonstrated a similar distribution of phyla throughout the 0.1 m to 6.85 m sampled depth, with *Actinobacteria*, and *Proteobacteria* being the predominant phyla followed by *Chloroflexi* and *Acidobacteria*. The basis for the dominance among the *Actinobacteria* of a single unclassified genus of *Acidimicrobiales* deserves further study. The Khalilovsky massif rock core microbial communities were substantially different from aqueous serpentine environments and relatively close to those of fully serpentinized rock surfaces and soils, notably in regard to the predominance of *Actinobacteria*.

**Supporting information**

**S1 Fig.** The Khalilovsky massif location and sample collection site. (A) The Gaysky District, Orenburg Region, Southern Ural Mountains, Russia (B) The Khalilovsky massif (C) The serpentinite rolling hills.

(PDF)

**S2 Fig.** Photographs of serpentinite minerals before (A) and after crushing (B).

(PDF)

**S3 Fig.** Representative X-ray diffraction (XRD) patterns of rock core samples collected at (A) 0.1 m, (B) 3.1 m and (C) 6.85 m in a depth. HM–Hydromagnesite, Sp–Serpentinites, HT–Hydrotalcite, Py–Pyroaurite.

(PDF)

**S4 Fig.** Backscattered electron images (BSE) with elemental spectra of the serpentinites collected at (A) 0.1 m, (B) 3.1 m and (C) 6.85 m in depth.

(PDF)
S5 Fig. Transmission electron micrographs (TEM) of serpentinites collected at (A) 0.1 m, (B) 3.1 m and (C) 6.85 m in a depth.

S6 Fig. Rarefaction curve of OTUs (operational taxonomic units) for 20 rock core samples.

S1 Table. The geochemical composition of serpentinites at the Khalilovsky massif. \% = C \times 10^n, C–content of elements, n–degree.

S2 Table. The diversity indices correlated with species richness (Shannon’s index), species diversity (Simpson’s index) and evenness and species richness estimators (Chao1).

S3 Table. Taxonomic assignments for all 16S rRNA tag sequences from rock core samples of the Khalilovsky massif, Russia.

S4 Table. Spearman’s rank correlation coefficients between relative abundance of OTUs and depth of serpentine minerals.

Acknowledgments
Microscopy studies were carried out at the Interdisciplinary Center of Analytical Microscopy of Kazan Federal University. The authors are grateful to Dr. N.I. Naumkina for carrying out the X-ray diffraction analysis and the Interdisciplinary Center for Collective Use (ID RFME-FI59414X0003) sponsored by Ministry of Education and Science of the Russian Federation.

Author Contributions
Conceptualization: Irina V. Khilyas, Tatyana A. Shcherbakova, Michael F. Cohen.
Data curation: Irina V. Khilyas, Tatyana A. Shcherbakova, Michael F. Cohen.
Formal analysis: Irina V. Khilyas, Alyona V. Sorokina, Maria I. Markelova, Tatyana A. Shcherbakova.
Funding acquisition: Margarita R. Sharipova.
Investigation: Alyona V. Sorokina, Anna A. Elistratova, Maria N. Siniagina.
Resources: Irina V. Khilyas, Margarita R. Sharipova.
Supervision: Margarita R. Sharipova, Michael F. Cohen.
Writing – original draft: Irina V. Khilyas, Tatyana A. Shcherbakova, Megan E. D’Errico, Michael F. Cohen.

References
1. Brazelton WJ, Morrill PL, Szponar N, Schrenk MO. Bacterial communities associated with subsurface geochemical processes in continental serpentinite springs. Appl Environ Microbiol. 2013; 79(13):3906–16. Epub 2013/04/16. https://doi.org/10.1128/AEM.00330-13 PMID: 23584766; PubMed Central PMCID: PMC3697581.
2. Douglas S, Beveridge TJ. Mineral formation by bacteria in natural microbial communities. FEMS Microbiol Ecol. 1998, 26(2):79–88. https://doi.org/10.1111/j.1574-6941.1998.tb00494.x
1. Gadd GM. Metals, minerals and microbes: geomicrobiology and bioremediation. Microbiology. 2010; 156(Pt 3):609–43. Epub 2009/12/19. https://doi.org/10.1099/mic.0.037143-0 PMID: 20019082.

2. Sleep NH. Geological and geochemical constraints on the origin and evolution of life. Astrobiology. 2018; 18(9):1199–219. Epub 2018/08/21. https://doi.org/10.1089/ast.2017.1778 PMID: 30124324.

3. Garzanti E, Doglioni C, Vezzoli G, Ando S. Orogenic belts and orogenic sediment provenance. J Geol. 2007; 115(3):315–34. https://doi.org/10.1086/512755

4. Bach W, Früh-Green GL. Alteration of the oceanic lithosphere and implications for seafloor processes. Elements. 2010; 6(3):173–8. https://doi.org/10.2113/gselements.6.3.173

5. Schrenk MO, Brazelton WJ, Lang SQ. Serpentinitization, carbon, and deep life. Rev Mineral Geochem. 2013; 75(1):575–606. https://doi.org/10.2138/rmg.2013.75.18

6. Brazelton WJ, Thornton CN, Hyer A, Twing KI, Longino AA, Lang SQ, et al. Metagenomic identification of active methanogens and methanotrophs in serpentinite springs of the Voltri Massif, Italy. PeerJ. 2017; 5:e2945. Epub 2017/02/06. https://doi.org/10.7717/peerj.2945 PMID: 28149702; PubMed Central PMCID: PMC5274519.

7. Evans BW, Hattori K, Baronnet A. Serpentine: What, Why, Where? Elements. 2013; 9(2):99–106. https://doi.org/10.2113/gselements.9.2.99

8. Gannoun A, Tessalina S, Bourdon B, Orgeval J-J, Birkle JL, Allègre CJ. Re–Os isotopic constraints on the genesis and evolution of the Dergamish and Ivanovka Cu (Co, Au) massive sulphide deposits, south Ural, Russia. Chem Geol. 2003; 196:193–207. https://doi.org/10.1016/S0009-2541(02)00413-8

9. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nature Methods. 2010; 7:335. https://doi.org/10.1038/nmeth.f.303 PMID: 20383131

10. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nature Methods. 2010; 7:335. https://doi.org/10.1038/nmeth.f.303 PMID: 20383131

11. Brazelton WJ, Ludwig KA, Sogin ML, Andreishcheva EN, Kelley DS, Shen C-C, et al. Archaea and bacteria with surprising microdiversity show shifts in dominance over 1,000-year time scales in hydrothermal chimneys. Proc Natl Acad Sci USA. 2010;200905369.
25. Daee FL, Okland I, Dahle H, Jorgensen SL, Thorseth IH, Pedersen RB. Microbial life associated with low-temperature alteration of ultramafic rocks in the Leka ophiolite complex. Geobiology. 2013; 11 (4):318–39. Epub 2013/04/05. https://doi.org/10.1111/gbi.12035 PMID: 23551703.

26. Gravuer K, Eskelinen A. Nutrient and rainfall additions shift phylogenetically estimated traits of soil microbial communities. Front Microbiol. 2017; 8(1271). https://doi.org/10.3389/fmicb.2017.01271 PMID: 28744266.

27. Mei N, Zergane N, Postec A, Erauso G, Ollier A, Payri C, et al. Fermentative hydrogen production by a new alkaliphilic Clostridium sp. (strain PROH2) isolated from a shallow submarine hydrothermal chimney in Prony Bay, New Caledonia. Int J Hydrogen Energ. 2014; 39(34):19465–73.

28. Oline DK. Phylogenetic comparisons of bacterial communities from serpentine and nonserpentine soils. Appl Environ Microbiol. 2006; 72(11):6965–71. Epub 2006/09/05. https://doi.org/10.1128/AEM.00690-06 PMID: 16959096; PubMed Central PMCID: PMC1636195.

29. Ronholm J, Goordial J, Sapers HM, Izawa MRM, Applin DM, Pontefract A, et al. Characterization of microbial communities hosted in quartzofeldspathic and serpentinite lithologies in Jeffrey Mine, Canada. Astrobiology. 2018; 18(8):1008–22. Epub 2018/07/11. https://doi.org/10.1089/ast.2017.1885 PMID: 29989429.

30. Suzuki S, Ishii S, Wu A, Cheung A, Tenney A, Wanger G, et al. Microbial diversity in The Cedars, an ultrabasic, ultrareducing, and low salinity serpentinitizing ecosystem. Proc Natl Acad Sci USA. 2013; 110 (38):15336–41. Epub 2013/09/05. https://doi.org/10.1073/pnas.1302426110 PMID: 24003156; PubMed Central PMCID: PMC3780913.

31. Tiago I, Verissimo A. Microbial and functional diversity of a subterranean high pH groundwater associated to serpentinitization. Environ Microbiol. 2013; 15(6):1687–706. https://doi.org/10.1111/1462-2920.12034 PMID: 23731249.

32. Woychees e KM, Meyer-Dom bard DAR, Cardace D, Argayosa AM, Arcilla CA. Out of the dark: transi- tional subsurface-to-surface microbial diversity in a terrestrial serpentinitizing seep (Manleluag, Pangas i, the Philippines). Front Microbiol. 2015; 6:44. https://doi.org/10.3389/fmicb.2015.00044 PMID: 25745416.

33. Ettenauer J, Pišlar G, Sterflinger K, Gonzalez-Muñoz MT, Jroundi F. Molecular monitoring of the microbial dynamics occurring on historical limestone buildings and after the in situ application of different bio-consolidation treatments. Sci Total Environ. 2011; 409(24):5337–52. Epub 09/22. https://doi.org/10.1016/j.scitotenv.2011.08.063 PMID: 21944202.

34. Frank YA, Kadnikov VV, Gavrilot SN, Banks D, Gerasimchuk AL, Podosokorskaya OA, et al. Stable and variable parts of microbial community in Siberian deep subsurface thermal aquifer system revealed in a long-term monitoring study. Front Microbiol. 2016; 7(2101). https://doi.org/10.3389/fmicb.2016.02101 PMID: 28082967.

35. Miettinen H, Kietavainen R, Sohliberg E, Numminen M, Ahonen L, Itavaara M. Microbiome composition and geochemical characteristics of deep subsurface high-pressure environment, Pyhäsalmi mine Finland. Front Microbiol. 2015; 6:1203. Epub 2015/11/19. https://doi.org/10.3389/fmicb.2015.01203 PMID: 26579109; PubMed Central PMCID: PMC4626562.

36. Hänchen M, Prigojibe V, Baciocchi R, Gonzal ez-Muñoz MT, Jroundi F. Molecular monitoring of the microbial communities hosted in quartzofeldspathic and serpentinite lithologies in Jeffrey Mine, Canada. Astrobiology. 2018; 18(8):1008–22. Epub 2018/07/11. https://doi.org/10.1089/ast.2017.1885 PMID: 29989429.

37. McCutcheon J, Southam G. Advanced biofilm staining techniques for TEM and SEM in geomicrobiol- ogy: Implications for visualizing EPS architecture, mineral nucleation, and microfossil generation. Chem Geol. 2018; 498. https://doi.org/10.1016/j.chemgeo.2018.09.016.

38. Yao M, Lian B, Teng H, Tian Y, Yang X. Serpentine dissolution in the presence of bacteria Bacillus mucilaginosus. J Geomicrobiol. 2012; 30. https://doi.org/10.1080/1490451.2011.653087.