Ultramicrobacteria from Nitrate- and Radionuclide-Contaminated Groundwater

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Abstract: The goal of the present work was to investigate the physicochemical and radiochemical conditions and the microbial diversity in groundwater collected near the Lake Karachai (Russia), which was formerly used for the disposal of liquid radioactive waste, to isolate the dominant bacteria, and to determine their taxonomy and the physiological characteristics responsible for their adaptation to this environment. Groundwater samples contained high concentrations of acetate, oxalate, nitrate, and sulfate, as well as radionuclides. High-throughput sequencing and analysis of the clone libraries revealed lower microbial diversity in the most strongly contaminated groundwater and a predominance of bacteria of the genera Polynucleobacter, Pusillimonas, Candidatus Pelagibacter, and of the candidate phylum Parcubacteria; these groups include species with an ultra small cell size. Archaeal sequences in the libraries belonged to ammonium oxidizers of the phylum Thaumarchaeota and methanogens of the phylum Euryarchaeota. Pure cultures of obligate and facultative ultramicrobacteria belonging to the genera Chryseobacterium, Microbacterium, Salinibacterium, Pusillimonas, Roseomonas, and Janibacter were isolated from water samples. In genomes of Pusillimonas and Roseomonas strains the genes associated with nitrate reduction, resistance to heavy metals and metalloids were revealed. Several isolates are able to participate in the geochemical process of nitrate conversion to N₂ using acetate; this results in decreasing redox potential, which in turn may stimulate radionuclide reduction and decrease radionuclide migration in groundwater.

Keywords: nitrate- and radionuclide-contaminated groundwater; 16S rRNA gene; pyrosequencing; ultramicrobacteria; Chryseobacterium; Pusillimonas; Salinibacterium; Janibacter
1. Introduction

According to the modern concept, ultra-microbacteria are characterized by (1) ultra small sizes (the volume of less than 0.1 \( \mu \)m\(^3\) for most cells in the populations); (2) maintenance of ultra small cell size regardless of the growth conditions and culture development stage; and (3) small genome size (~3.2 to ~0.58 Mb) [1,2]. The majority of oligotrophic bacteria found in freshwater and marine environment are planktonic and have small cell volumes (<0.1 mm\(^3\)) [3]. Ultra-microbacteria are responsible for the major part of the biomass and metabolic activity in oceanic ecosystems, and express higher metabolic activity per unit of volume of seawater than larger bacterial cells [3]. Unlike copiotrophs that become small as a result of reductive cell division, oligotrophic ultra-microbacteria retain a small volume of cells even during the life cycle. Subsurface water and sediments not associated with petroleum, peat, or coal deposits are also a freshwater oligotrophic environment [4]. An inflow of organic and mineral contaminants to subsurface water can selectively increase the activity and abundance of microorganisms. The microbial populations of groundwater contaminated by radioactive waste are of special interest, since they may affect the valency and mobility of uranium and other variable-valency elements. While occurrence of bacteria with small genomes and ultra small cells in groundwater has been reported [5,6], culturable filterable bacteria isolated from subsurface horizons remain scarcely studied.

The first report of phylogenetic diversity of microorganisms with ultra small cells in deep environments was presented by Mioshi et al. [7] for anaerobic groundwater of the Tono uranium mine, Japan. The bacteria passing through the membranes with 0.22-\( \mu \)m pores belonged to the class Betaproteobacteria, while uncultured small-celled bacteria of the candidate groups OD1 and OP11 passed through the filters with 0.1-\( \mu \)m pores. The genomes of OD1 and OP11 bacteria were retrieved from fresh and brackish water samples and hydrothermal sites [8]. Based on single-cell genomic data, candidate Bacterial phyla OD1 and OP11 were assigned to the candidate phyla Parcubacteria and Microgenomates, respectively, of the superphylum ‘Patescibacteria’ [9]. Near-complete genome sequences of Parcubacteria (OD1) were obtained using metagenomic analysis of DNA from anoxic groundwater samples from an aquifer near Rifle, CO, USA [5,6]. Storage of uranium mill tailings at the Rifle alluvial aquifer site resulted in formation of a groundwater uranium plume [10]. Nelson and Stegen [11] reconstructed eight OD1 genomes from oxic groundwater samples from Hanford 300 Area at the Rifle. The organisms had small genomes (generally <1 Mb) missing the genes associated with biosynthetic capabilities and DNA repair, but containing the genes potentially encoding the attachment and adhesion proteins, which indicated that “the Parcubacteria were ecosymbionts or parasites of other organisms”. Anaerobic ultra Microbacterium ‘Dehalobium chlorocoercia’ DF-1 able to dechlorinate polychlorinated biphenyl using H\(_2\) and formate as electron donors was isolated from marine sediments of Charleston Harbor [12]. The genomes of Nanoarchaea with small size and limited metabolic capacity, which were probably involved in anaerobic carbon and hydrogen cycles, were reconstructed from an aquifer near Rifle, CO, USA also [13].

Sequences of the 16S rRNA genes both of the cultured members of the phyla Proteobacteria, Dehalococcoides, Actinobacteria, Acidobacteria, Firmicutes, Bacteroidetes, Planctomycetes and of uncultured bacteria of Candidate divisions OD1, OP3 and OP11 were revealed in clone libraries of microorganisms from groundwater of a disposal site for liquid radioactive wastes at the Siberian Chemical Complex (SCC, Tomsk-7, or Seversk, Russia) [14].

Advances in metagenomics, single-cell genome analysis, and metatranscriptomics promoted an increase in our knowledge of the genetics of uncultured microorganisms. However, investigation of pure cultures remains essential for correlation of the results of metagenomic analysis and the physiological and genomic characteristics of cultured microorganisms.

In the present work, the microorganisms of groundwater from the Lake Karachai region, one of the most contaminated basins in the world, are investigated. Since 1951, Lake Karachai (V-9 industrial reservoir) was used for storage of liquid intermediate-level nuclear wastes at the Production Association “Mayak” (PA “Mayak”), the first Russian plant producing plutonium, located in the north of the Chelyabinsk region [15]. PA “Mayak” activity resulted in a local accumulation of large quantities of wastes coming from radiochemical technology and in radioactive
contamination of some parts of the neighboring regions by long-lived radionuclides, mainly $^{90}$Sr and $^{239}$Pu [16]. The effluent discharged into the Lake Karachai consisted of radioactive, slightly alkaline (pH 7.9–9.3) brine with density between 1.006 and 1.095 g/cm$^3$; the dissolved salt varied from 16 to 145 g/L. The major components of the waste were nitrate, acetate, oxalate, sulfate, and sodium [17]. Waste effluents were enriched with Fe, Al, Zr, and rare earth elements. Long-term exploitation of V-9 reservoir resulted in large-scale groundwater contamination in the area, which was caused by infiltration of radioactive solutions through the permeable bottom of the reservoir in different directions. The radioactivity of $^{90}$Sr, $^{106}$Ru and $^{137}$Cs in the 50–100-m deep water horizon was 20–300 times lower than in the waters of the V-9 reservoir, and NO$_3^-$ concentration in the plume under the reservoir was 3.5–5 times higher than that in the water of the V-9 reservoir. In 2015, the Lake Karachai was filled with sand and covered with concrete and rock plates.

The high level of nitrate in the groundwater in the Karachai area prevents reductive immobilization of uranium. Denitrification intermediates (nitrite, nitric oxide, and nitrous oxide) are known to be capable of oxidizing and mobilizing U(IV) abiotically [18]. The removal of nitrate favored reduction of oxidized radionuclides and decreased their migration [18,19]. Geochemical evidence for denitrification in groundwater in the Karachai area is available [20], but the detection of microorganisms capable of nitrate reduction is a necessary stage for prediction of behavior of nitrate and radionuclides in contaminated subsurface waters.

The analysis of clone libraries of the 16S rRNA genes of microorganisms from groundwater of the Karachai area resulted in the retrieval of the sequences of uncultured bacteria of the candidate phylum OD1 (or Parcubacteria), Polynucleobacter sp. and ‘Candidatus Pelagibacter sp.’ with small-sized cells [21]. Over 50 strains of aerobic organotrophic and anaerobic denitrifying and iron-reducing bacteria belonging to 23 genera and 33 species were isolated from the samples. Further research on the composition of the microbial community and on the ecologically possible functions of the obtained pure cultures was essential.

The goal of the present work was to investigate the physicochemical and radiochemical conditions and the microbial community composition in nitrate- and radionuclide-contaminated groundwater collected near Lake Karachai (Russia), to isolate bacteria, including those with ultra-small cell size, and to determine the taxonomic position, physiological, and genomic characteristics responsible for their adaptation to this environment.

2. Materials and Methods

2.1. Sampling of Groundwater

Groundwater was sampled from the southern limb of the contamination plume in the PA “Mayak” industrial area. The samples were collected from observation wells (20–80 m deep), 2.0–3.2 km from Lake Karachai. Groundwater samples for microbiological analysis were collected into sterile bottles using an electrical pump and sealed hermetically. The samples for chemical and radiochemical studies were collected into 20-L bottles under nitrogen atmosphere to avoid contact with air. The samples for microbiological studies were collected into sterile 1-L bottles and hermetically sealed; samples for molecular studies were collected into 3-L bottles, each containing 0.75 L ethanol. The samples were transported to the laboratory in hermetic bottles for filtration and analysis.

2.2. Microbial Enumeration and Isolation

Numbers of anaerobic denitrifying, fermenting, iron- and sulfate-reducing, and methanogenic prokaryotes were estimated using the MPN technique with two replicates for each tenfold dilution of groundwater samples as described previously [14]. Aerobic organotrophs were enumerated in the TEG medium containing the following (g L$^{-1}$ distilled water): bacto-tryptone, 5.0; yeast extract, 2.5; glucose, 1.0; pH 7.0. Aerobic lithoautotrophic ammonium-oxidizing bacteria were revealed by inoculation of undiluted groundwater samples to the DSM medium 1583 (https://www.dsmz.de/microorganisms/medium/pdf/DSMZ_Medium1583.pdf). NO$_3^-$ production
was detected with the Griess reagent by development of pink color. Lithoautotrophic
nitrite-oxidizing bacteria were incubated in the medium containing per liter distilled water: 1 g
NaNO₂, 0.5 g KH₂PO₄, 0.5 g MgSO₄.7H₂O, 0.5 g NaCl; 0.4 g FeSO₄.7H₂O; 1.0 g Na₂CO₃; pH 8.0–9.2.
NO₃⁻ production was detected with diphenylamine by development of blue color. Inoculated media
were incubated at room temperature (18–24 °C) until the onset of growth, but no longer than 30
days. All the cultures were then examined under an Axio Imager.D1 epifluorescence microscope
(Carl Zeiss, Oberkochen, Germany), and specific microbial metabolites were analyzed.

Enrichment cultures of aerobic organotrophic and denitrifying bacteria obtained in our
experiments were used for isolation of pure cultures by successive transfer from liquid media to
respective solid media. Search for ultramicrobacteria in the samples was also carried out. For this
purpose, groundwater (20–40 mL) was filtered through 0.45- and 0.22-μm pore size membrane
filters (Whatman, Maidstone, UK). The filtrate was used to inoculate low-nutrient AMS1 medium
with methionine, glycine, pyruvate, vitamins and inorganic salts, as recommended for the growth of
‘Candidatus Pelagibacter ubique’ [22]. To exclude the development of rapidly growing bacteria, the
cultures were incubated for 7–14 days in a refrigerator at 6–8 °C, then at room temperature (18–22
°C), and then transferred onto solid media of relevant composition. Sequential transfers from liquid
to solid media resulted in isolation of pure cultures of filterable bacteria. Single colonies were used
for microbiological and molecular studies. Such characteristics of strains as temperature and pH
ranges and optima for growth, NaCl tolerance, catalase activity, anaerobic growth, nitrate reduction,
acid production, and growth with various substrates were tested as described by Smibert and Krieg
[23].

2.3. Microscopy

Cell morphology of the isolates was examined using a JEOL JEM-1011 (JEOL Ltd., Tokyo,
Japan) transmission electron microscope at accelerating voltage of 80 kV. The cells were contrasted
with 2% phosphotungstic acid. For ultrathin sectioning, cells were harvested by centrifugation and
then fixed in a solution of 1.5% (w/v) glutaraldehyde in 0.05 M cacodylate buffer (pH 7.2) at 4 °C for
1 h. The cells were washed three times in the same buffer and post-fixed in 1% OsO₄ in 0.05 M
cacodylate buffer (pH 7.2) for 3 h at 20 °C. The preparation was dehydrated in a series of increasing
ethanol concentrations, in acetone and embedded in Epon 812 epoxy resin. Ultrathin sections were
mounted on grids, post-stained with 3% (w/v) uranyl acetate in 70% (v/v) ethanol for 30 min, and
afterwards additionally stained with lead citrate according to Reynolds [24]. Thin sections were
examined in a JEOL JEM-100 transmission electron microscope at an acceleration voltage of 80 kV.

2.4. Identification of Pure Cultures

Pure bacterial cultures were identified by analysis of their 16S rRNA gene sequences. DNA
isolation, amplification, and sequencing of the 16S rRNA genes were carried out as described
previously [25]. The full-sized 16S rRNA gene was amplified using the 8f–1492r primers and was
used as a template for DNA sequencing with the primers 8f, 519r, 519f, and 1492r. Preliminary
analysis of the 16S rRNA gene sequences was carried out using the Ribosomal Database Project
(RDP, http://rdp.cme.msu.edu) data and software. The sequences were edited using BioEdit
(http://jwbrown.mbio.ncsu.edu/ BioEdit/bioedit.html) and aligned with those of closely related
Bacterial species using MAFFT 7.427 [26].

2.5. Analytical Methods

Molecular hydrogen, nitrogen, and methane produced by microorganisms were analyzed by
gas chromatography using approaches described previously [14]. Sulfide was determined with
dimethyl-p-phenylenediamine by a modified colorimetric method of Pachmayr [27]. The values of
pH, Eh, and total salinity of groundwater were measured using a HITON-S hydrogeochemical
sampler (LLC "Scientific and Production Center Pals, Samara, Russia) equipped with selective
electrodes; the sampler was submerged into the wells and the parameters at an examined depth
were recorded. Iron, magnesium, calcium, and sodium in groundwater were analyzed on an AASIN atomic absorption spectrophotometer (Carl Zeiss, Jena, Germany). Nitrate was analyzed on an Ecotest-01 device (NPP Econiks, Moscow, Russia) with an ELIT-21 (NPP Econiks, Moscow, Russia) ion-selective electrode. Nitrite was determined colorimetrically with sulfanilic acid and α-naphthylamine [28]. Other components of groundwater were analyzed as described earlier [29]. Radiochemical procedures for speciation of actinides in the environment were described previously [30].

2.6. Radiotracer Methods and Tolerance to Heavy Metals and Ionizing Radiation

The rates of sulfate reduction and methanogenesis in groundwater were determined by radiotracer methods using labeled Na$_{235}$SO$_4$, $^{14}$CH$_3$COONa, and NaH$^{14}$CO$_3$ according to the method employed by Nazina et al. [31]. Aliquots of formation water were incubated with labeled substrates for 24–48 h at 22 °C.

Stock solutions of 1.0 g L$^{-1}$ CdSO$_4$ 8H$_2$O, 2.0 g L$^{-1}$ K$_2$Cr$_2$O$_7$, 2.0 g L$^{-1}$ Zn(NO$_3$)$_2$ 6H$_2$O, 2.0 g L$^{-1}$ CuSO$_4$ 6H$_2$O, and 2.0 g L$^{-1}$ UO$_2$(NO$_3$)$_2$ 6H$_2$O were sterilized at 121 °C for 15 min, and aseptically added into the sterilized liquid TEG medium supplemented with 5% (w/v) NaCl to prepare selective media containing 10, 50, 100, 200, and 300 mg L$^{-1}$ Cd$^{2+}$; 25, 100, 200, 300, and 400 mg L$^{-1}$ Cr$^{6+}$; 10, 25, 50, 100, and 200 mg L$^{-1}$ Cu$^{2+}$; 10, 25, 50, 100, and 200 mg L$^{-1}$ Zn$^{2+}$; and 10, 100, 200, 300, and 400 mg L$^{-1}$ U$^{6+}$, respectively. Media containing series of concentrations of heavy metals were inoculated with isolated strains and incubated at 24 °C for 24 days. After incubation, the total cell numbers were determined by light microscopy. The numbers of living and dead cells were determined by staining with the LIVE/DEAD® BacLight™ Bacterial Viability Kit for microscopy & quantitative assays according to the recommendations (https://www.thermofisher.com/order/catalog/product/L7012) under a Microoptix MX 300 light microscope (West Medica Production und Handels-GmbH, Perchtoldsdorf, Austria) with an epifluorescence block. Inoculated media without metals were used as the controls.

Radioresistance of exponentially growing cultures of strains JR1/69-1-13, JR1/69-2-13, and JR1/69-3-13 was determined using an YELV-10-10-5-70 electron accelerator with beam energy 10 ± 0.4 MeV and beam current 500 ± 0.4 μA (IPCE RAS). For irradiation we used different radiation doses (0.1, 0.5, 1.0, 5.0, 10.0, 15.0 and 25.0 kGy) with dose rate 0.3 kGy/min and radiation doses 10.0, 15.0 and 25.0 kGy with dose rate 3 kGy/min. Radi ation doses were controlled using a Fricke ferrous sulfate dosimeter; for doses below 5 kGy, the radiation dose was controlled by Radiochromic film dosimeters SO PD(F)R 5/50 and SO PD(E) 1/10 FSUE “VNIIFTRI”.

The cell suspensions in tubes with 10 mL of TEG medium closed with rubber stoppers were exposed to radiation for predetermined times. At each time-point, three individual aliquots of each strain were removed from the radiation source. The effect of radiation on the microorganisms was assessed 60 min after irradiation according to the respiratory activity of living cells in the same test tubes (acute experiment), which was determined by staining with MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) with subsequent extraction of produced formazan by dimethyl sulfoxide (DMSO) and measuring the extract optical density at 500–600 nm, as was proposed previously [32,33]. The results were calculated as percentages of the respiratory activity in the control samples. The experiment was carried out in three replicates. Resistance of bacteria to heavy metals and ionizing radiation was determined by the standard approaches using semilethal (LD$_{50}$) and lethal doses (LD$_{100}$).

2.7. Construction and Analysis of the 16S rRNA Gene Clone Libraries

DNA Extraction: Groundwater samples from the observation wells 1/69 (from the depth 44 m), 2/68 (80 m), and 14/68 (80 m) (each 2.25 L) were fixed with ethanol (each with 0.75 L) immediately after sampling and were passed through 0.22-μm membrane filters. Prior to analysis, the filters with biomass were stored at −20 °C. The total DNA was isolated from the filters using DiatomTM DNAprep kit for DNA isolation (Shemyakin and Ovchinnikov Institute of Bioorganic Chemistry, Russian Academy of Sciences, Moscow, Russia) as described earlier [14]. For total DNA isolation,
the cell pellet was suspended with guanidine hydrochloride at 65 °C for 60 min. The sorbent (Diatomid/silica) was added to the cell lysate. After DNA sorption, the supernatant was removed and the sorbent was washed using 70% (v/v) ethanol. The total DNA preparation was dissolved in 100 μL H2O (MQ) and used for 16S rRNA gene amplification and microbial community analysis by cloning or/and high-throughput sequencing.

The 16S rRNA gene amplification, cloning, and sequencing of cloned fragments of the 16S rRNA genes: Isolated DNA obtained from observation wells 1/69, 2/68 and 14/68 was amplified by PCR using the primers specific for bacteria (8-27f, and 1492r) [34,35] and *Archaea* (A8F, A109f, A517R, A800f, and A1041r) [36,37]. The amplicons were separated using 0.8% agarose gel, stained with ethidium bromide, and visualized under UV illumination. For PCR product purification, the fragments were excised from the gel, and the DNA was extracted using a V-gene DNA Gel Extraction Kit (V-gene Biotechnology Ltd. Hangzhou 310022, China) with subsequent precipitation of the DNA fragments by 70% ethanol and 0.3 M CH3COONa. Bacterial and archaeal 16S rRNA genes were purified and cloned into the pTZ57RT plasmid vector (Fermentas, Vilnius, Lithuania). The clones containing the DNA inserts were sequenced using the plasmid primers M13D and M13R. The sequencing was performed on a 3730 DNA Analyzer using BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, United States), in compliance with the recommendations of the manufacturers.

Phylogenetic analysis of the 16S rRNA gene sequences: The sequences were homology-searched using the NCBI BLAST software package (http://www.ncbi.nlm.nih.gov/blast) and SILVA (http://www.arb-silva.de). The sequences were edited using the BioEdit software package (http://jwbrown.mbio.ncsu.edu/BioEdit/bioedit.html). The sequence data were aligned with the sequences retrieved from the GenBank database using the MAFFT 7.427 [26], with the clones with at least 98% similarity grouped into operational taxonomic units (OTUs) or phylotypes. Detection of the chimeras and checking for the secondary structure anomalies was carried out using the CHIMERA-CHECK program from the Ribosomal Database Project.

Pyrosequencing of 16S rRNA amplicons and data analysis: The filtered cells from 2.25 L of groundwater taken from observation well 176/94 yielded 1656 ng of DNA. 10 ng of DNA was used for amplification of the 16S rRNA gene fragment comprising V3–V8 variable regions: U341F (5′-CCT ACG GGR BGC ASC AG) and E1406R (5′-GAC GGG CGG TGW GTR CA). The PCR fragments were sequenced with a Roche Genome Sequencer (GS) FLX, using the XL+ protocol. The 16S rRNA gene sequences determined by pyrosequencing were subjected to the standard filter for environmental pyrosequencing datasets and selected for tags that displayed perfect matches to the primers and contained no ambiguous nucleotides. The 16S rRNA reads starting from primer U341F were used for further analysis. The 16S rRNA gene data were analyzed using the RPD Classifier program package [38]. As a first step, the sequences were distributed between bacteria and archaea using the online RDP Naive Bayesian rRNA Classifier version 2.0 (http://rdp.cme.msu.edu/index.jsp). Then, *Bacterial* and *archaeal* data sets were analyzed separately.

Bacterial and archaeal 16S rRNA gene fragments were processed using a combination of QIIME [39] and USEARCH [40] pipelines. After demultiplexing and quality control, the sequences were de-replicated and chimeras and singletons were removed. The resulting datasets upon filtration consisted of 117685 16S rRNA reads assigned to bacteria and 104 reads assigned to *Archaea*. The sequences were de novo clustered into OTUs as having ≥98% sequence identity using QIIME. Taxonomic assignment of each OTU was accomplished using the online RDP Classifier (https://pyro.cme.msu.edu/classifier/form.spr).

Statistical Analysis. Alpha diversity metrics were calculated using the QIIME [39]. ClustVis was employed to generate the heat map of most abundant genera [41].

2.8. Genome Analysis

Genomes of *Pusillimonas* sp. JR1/69-2-13, *Pusillimonas* sp. JR1/69-3-13 and *Roseomonas aestuarii* JR1/69-1-13 were sequenced and annotated as described previously [42,43]. Identification of protein-coding sequences and primary annotation was performed using the NCBI Prokaryotic
Genome Automatic Annotation Pipeline (PGAAP) [44]. Additional gene prediction and functional annotation were performed in the Rapid Annotation using Subsystems Technology (RAST) server [45] and Integrated Microbial Genome-Expert Review pipeline [46], respectively. The analysis of metal resistance was performed with the AntiBacterial Biocide and Metal Resistance Genes Database (BacMet) [47].

2.9. Nucleotide Sequence Accession Numbers

The sequences of the 16S rRNA genes of microorganisms from libraries 1/69, 2/68, 14/68 and 176/94 were submitted to the GenBank/DDBJ/EMBL databases as BioProject: PRJNA548669 (http://www.ncbi.nlm.nih.gov/bioproject/548669). The draft genome sequences of strains *Pusillimonas* sp. JR1/69-2-13, *Pusillimonas* sp. JR1/69-3-13 and *R. aestuarii* JR1/69-1-13 are available at DDBJ/EMBL/GenBank under the accession nos. https://www.ncbi.nlm.nih.gov/nuccore/PDNV00000000, https://www.ncbi.nlm.nih.gov/nuccore/PDNW00000000, and https://www.ncbi.nlm.nih.gov/nuccore/NZ_PDOA00000000.1, respectively.

3. Results

3.1. Physicochemical, Radiochemical, and Microbiological Characteristics of Groundwater

Environmental conditions and an abundance of the major microbial physiological groups were studied in groundwater samples collected in the vicinity of Lake Karachai (industrial reservoir V-9), which was used as a repository of liquid radioactive waste at the PA “Mayak” area. The studied groundwater samples had high redox values (Eh from +110 to +420 mV), pH from 6.9 to 8.0, and elevated total salinity and radionuclide content (Table 1). Chloride, sulfate, nitrate, and acetate were the major anions. The concentrations of nitrate, acetate, and carbonate varied from 120 to 42,700 mg L\(^{-1}\), from 26 to 921 mg L\(^{-1}\), and from 59 to 1820 mg L\(^{-1}\), respectively. Uranium and americium concentrations were below the relevant intervention levels (IL) by Radiation safety standards (NRB-99/2009) of Russian Federation regulation, while strontium concentration exceeded IL of 5.0 Bq/dm\(^3\). Temperature of the studied groundwater in the Karachai area was 6–7 °C [48].

Groundwater samples were found to contain cultivated aerobic organotrophic (10–10\(^5\) cells mL\(^{-1}\)) and nitrifying prokaryotes, as well as anaerobic fermenting (10–10\(^5\) cells mL\(^{-1}\)), denitrifying (0–10\(^4\) cells mL\(^{-1}\)), and iron-reducing bacteria (0–10\(^4\) cells mL\(^{-1}\)) (Figure A1). The numbers of sulfate-reducing (<10 cells mL\(^{-1}\)) and methanogenic prokaryotes (0–10\(^5\) cells mL\(^{-1}\)) were very low, and sensitive radiotracer measurements with labeled Na\(^{235}\)SO\(_4\), 14CH\(_3\)COONa, and NaH\(^{14}\)CO\(_3\) revealed no sulfate reduction and methanogenesis in most of the studied samples (Table 1).

3.2. Phylogenetic Diversity of Archaea and Bacteria in Groundwater Samples

DNA extracted from groundwater samples collected at weakly contaminated (wells 1/69 and 176/94) and strongly contaminated zones (wells 2/68 and 14/68) was used for cloning of the 16S rRNA genes and pyrosequencing by means of the 454 platform (Table A1). Diversity indices for archaeal components in the library 176/94 obtained by pyrosequencing and in the clone libraries 1/69 and 2/68 were similar, while cloned sequences from the library 14/68 exhibited low diversity and domination of a small number of phylotypes (Table A2). Diversity indices of bacteria were significantly higher in the library 176/94 than in all clone libraries, especially the library 14/68, in which the dominance index was 0.97, indicating that this library was almost completely represented by a single phylotype.

The share of archaeal 16S rRNA gene sequences in the 176/94 library did not exceed 0.5%, and their OTUs (phyotypes) were represented mainly by the sequences of *Thaumarchaeota* belonging to autotrophic ammonium-oxidizing archaea of the genera *Nitrosopumilus* (30.7%) and *Nitrososphaera* (21.2%), uncultured *Pacearchaeota* (42.3%), and *Euryarchaeota* (5.8%), belonging to methanogens of the genera *Methanoregula* and *Methanothrix* (Figures 1a and A2a, Table A1). A similar archaeal composition was revealed in the clone library from well 1/69. In the libraries from highly
| Parameters                  | Standard Error | Observation Wells |
|----------------------------|----------------|-------------------|
|                            |                | 2/68 *            | 14/68 | 1/69 * | 1/69 | 176/94 |
| **Distance from the Karachai, m** |                | 2100             | 2000  | 3200   | 2500  |
| **Year of Analyses**       |                | 2010             | 2010  | 2013   | 2011  | 2013   | 2012   |
| **Depth of Sampling, m**   | 80             | 80               | 20–30 | 60     | 44    | 20–30  | 60     | 60     |
| **pH ±0.1**                | 8.0            | 7.6              | 7.7   | 7.2    | 7.9   | 7.3    | 6.9    | 7.8    |
| **Eh, mV ±1.0**            | 310            | 270              | 110   | 120    | 200   | 110    | 420    | 120    |
| **Na⁺ + K⁺, mg L⁻¹**:      | ±0.02          | 9071             | 1099  | 340    | 3230  | 145    | 370    | 2610   | 356    |
| **Ca²⁺ + Mg²⁺, mg L⁻¹**:   | ±0.09          | 6712             | 3102  | -      | -     | -      | 302    | -      | 2460   |
| **Fe₅Ο₆, mg L⁻¹**:         | ±0.004         | 12.1             | 0.86  | -      | -     | 3.3    | -      | -      | 0.75   |
| **Cl⁻, mg L⁻¹**:           | ±0.004         | -                | -     | -      | -     | 140    | -      | -      | 215    |
| **SO₄²⁻, mg L⁻¹**:         | ±0.02          | -                | -     | -      | -     | 110    | -      | -      | 80     |
| **NO₃⁻, mg L⁻¹**:          | ±0.09          | 42,700           | 13,600| 120    | 2900  | 120    | 200    | 2300   | 2090   |
| **CO₃²⁻/HCO₃⁻, mg L⁻¹**:   | ±0.23          | 1310             | 267   | 80     | 120   | 360    | 59     | 200    | 1820   |
| **Acetate, mg L⁻¹**:       | ±0.5           | 921              | 795   | -      | -     | 26     | -      | -      | -      |
| **Radioactivity, Bq L⁻¹**  | -              | -                | -     | -      | -     | -      | -      | -      | -      |
| **²³⁹,²⁴⁰Pu**               | 0.01           | 5.78             | -     | 0.03 ± 0.01 | 0.18 ± 0.04 | 0.14 | <0.02 | 0.12 ± 0.03 | 0.8 |
| **⁹⁰Sr**                    | ±0.001         | -                | -     | 220 ± 30 | 3300 ± 300 | 12.4 | 155 ± 20 | 2200 ± 250 | 300 |
| **¹³⁷Cs**                   | ±0.001         | -                | -     | <0.2    | 2.9 ± 0.4 | <0.2 | <0.2 | -      | -      |
|          | ±0.001 | 3.01 | <0.02 | 0.06 ± 0.01 | 0.8 | <0.02 | 0.20 ± 0.04 | 0.7 |
|----------|--------|------|-------|-------------|----|-------|-------------|----|
| $^{241}$Am | ±0.001 | 7.2  | <0.02 | 1.0 ± 0.3   | 0  | <0.02 | 3.7 ± 1.2   | <0.5 |
| $^{237}$Np | ±0.001 | 39   | -     | 3.1 ± 0.7 ** | 12.1 ± 2.0 ** | 2.6 | 0.3 ± 0.1 ** | 1.1 ± 0.2 ** | 45 |
| $^{238}$U, Bq L$^{-1}$ or mg L$^{-1}$** | ±0.001 | -    | -     | -            | -  | 0.01  | -            | -  |
| SR rate, μg S$^2$/(L day) | ±0.001 | -    | -     | -            | -  | 0     | -            | -  |
| MG rate, μg CH$4$/(L day) | ±0.001 | -    | -     | -            | -  | 0     | -            | -  |

* Data from the paper Nazina et al. [21]. ** Content $^{238}$U in the groundwater sample in mg L$^{-1}$. 


contaminated groundwater from wells 2/68 and 14/68, the sequences of ammonium-oxidizing \textit{Thaumarcheota} (\textit{Nitrosopumilus} and \textit{Nitrososphaera}) were numerically predominant. Thus, nitrate-contaminated groundwater was inhabited by archaea involved in the nitrogen cycle, as well as by uncultured \textit{Pacarcheota} (42.3\%) characterized by small genomes and involved in anaerobic carbon and hydrogen metabolism, probably associated with symbiotic and/or fermentation-based lifestyles [13].

Cloned sequences of \textit{Bacterial} 16S rRNA genes in the libraries 14/68, 2/68, and 1/69 belonged to 4, 60, and 16 phylotypes, respectively, while the more representative library 176/94 obtained by pyrosequencing was found to contain 927 phylotypes of the species level. \textit{Proteobacteria} sequences accounted for 35.2 to 97.1\% of total number of sequences in the libraries (Figures 1b and A2b, Table A1). Among the \textit{Proteobacteria} sequences, those of the Beta-, Alpha-, and Gammaproteobacteria were the most numerous.

Other phyla revealed in the libraries from contaminated groundwater were \textit{Candidatus} Parcubacteria (former OD1, 34.5\%), which include bacteria with ultra-small size of cells and reduced metabolic capacities [9], Bacteroidetes (10.0\%), Chloroflexi (10.7\%), and Firmicutes (3.5\%).

Members of the genus \textit{Pusillimonas} were responsible for 18.2 and 97.1\% of sequences in the libraries of highly contaminated groundwater from wells 2/68 and 14/68, respectively. Other numerous phylotypes belonged to bacteria affiliated with the candidate phylum Parcubacteria (34.5\%), uncultured Coxiellaceae (30.9\%), and genera Polynucleobacter (14.3\%), Frigidibacter, Levilinea, Curvibacter, Lysobacter, Gallionella, Sideroxydans, Simplicispira, Hydrocarboniphaga, Sulfuritalea, and Reyranella (Figure 1b, Table A1). Bacteria of the genera \textit{Pusillimonas}, Polynucleobacter, Microbacterium, Chryseobacterium, \textit{Candidatus Pelagibacter}, and the candidate phylum Parcubacteria include species with small or ultra-small size of the cells (Figure 1b, Table A1).

### 3.3. Isolation and Taxonomy of Bacteria from Groundwater

Since the 16S rRNA gene libraries contained the sequences of uncultured bacteria such as ‘\textit{Candidatus Pelagibacter}’, search for bacteria with small and ultra-small cells and their isolation were necessary. Groundwater samples were sequentially filtered through the filters with 0.45- and 0.22-μm pores. The latter filtrate was inoculated into the low nutrient medium for ‘\textit{Candidatus Pelagibacter}’ and using an approach recommended by [22] as “selective filtration enrichment procedure and long-term incubations at low temperatures”, we isolated six strains of small-celled bacteria. Based on analysis of their 16S rRNA gene sequences, filterable microorganisms were affiliated with phyla Actinobacteria and Bacteroidetes. Four strains were assigned to \textit{Janibacter limosus} (strain JR43/78-4-13, 99.7\% similarity with the 16S rRNA gene of the type strain of the species), \textit{Microbacterium oxydans} (JR3/2-13, 99.6\%), \textit{Paeniglutamicibacter sulfureus} (JR1/1-8b, 99.3\%), and \textit{Mycolicibacterium iranicum} (JR2/2-1, 100\%). The 16S rRNA gene sequences of strains J1-13Tb and JR2/1-6 had 98.2 and 98.5\% similarity with those of the type strains \textit{Chryseobacterium haifense} H38\textsuperscript{T} and \textit{Salinibacterium amurskyense} KMM 3673\textsuperscript{T}, respectively. These values were below 98.65\%, the threshold for differentiating two species [49]. Thus, strains J1-13Tb and JR2/1-6 probably belong to two new species. Attempts to isolate the target bacterium ‘\textit{Candidatus Pelagibacter}’ proved unsuccessful. Five strains were isolated on media for aerobic and denitrifying bacteria without the filtration stage. Three strains were assigned to known species \textit{Roseomonas aestuarii} (JR1/69-1-13, 99.6\%), \textit{Rhodococcus qingshengii} (JR3/2-4, 99.7\%), and \textit{Rhodococcus fascians} (J1-4, 100\%).
Figure 1. Heatmap analysis of the distribution of the dominant genera of Archaea (a) and bacteria (b) in groundwater samples. The double hierarchical dendrogram shows microbial distribution in these samples. The relative values for microbial genera are marked by colors from blue to brown designating the least abundant to most abundant. Abundance is expressed as the ratio of the number of targeted sequences to the total number of sequences from each groundwater sample.

The 16S rRNA gene sequence similarity of the strains JR1/69-2-13 and JR1/69-3-13 with closely related Pusillimonas species was within the range of 97.6–96.0, and the similarity between strains was 98.7%, showing their affiliation to two new species. Phenotypic characteristic and taxonomy of the isolated strains is given in Tables 2 and 3.

Table 2. Taxonomic position of microorganisms isolated from the groundwater at the Karachai area.

| Strain            | Closest Cultivated Microorganism, Accession No. of 16S rRNA Gene | Accession No.* of 16S rRNA Gene | 16S rRNA Similarity, % | Genome Size, Mb |
|-------------------|---------------------------------------------------------------|---------------------------------|------------------------|-----------------|
| JR1/69-1-13       | Roseomonas aestuarii JC17T, NR_116913                         | MG205609                        | 99.6                   | 5.2             |
| JR1/69-2-13       | Pusillimonas harenae B201, GQ232740                            | MG205613                        | 98.6                   | 4.3             |
| JR1/69-3-13       | Pusillimonas harenae B201, GQ232740                            | MG205614                        | 98.5                   | 4.5             |
| J1-13Tb           | Chryseobacterium haifense H38T, NR_044167                      | MG205639                        | 98.2                   | 2.8 **          |
| JR3/2-4           | Rhodococcus qingshengii JCM 15477T, DQ090961                  | MG205632                        | 99.7                   | 7.2 **          |
| J1-4              | Rhodococcus fascians ATCC 12974T, NR_119126                    | MG205633                        | 100                    | 5.8 **          |
| JR2/2-1           | Mycolicibacterium iranicum M05T, NR_117909                    | MG205634                        | 100                    | 6.3 **          |
| JR43/78-4-13      | Janibacter limosus DSM 11140T, NR_026362                       | MG205635                        | 99.7                   | 3.4 **          |
| JR2/1-6           | Salinibacterium amurskyense KMM 3673T, AF539697               | MG205636                        | 98.5                   | 2.8 **          |
| JR3/2-13          | Microbacterium oxydans DSM 20578T, NR_044931                   | MG205637                        | 99.6                   | 3.9 **          |
| JR1/1-8b          | Paeniglutamicibacter sulfureus DSM 20167T, NR_026237          | MG205638                        | 99.3                   | ND ***          |

* Accession no. for 16S rRNA genes of the isolated strains. ** Genome size for the type strain of the species obtained from NCBI database (https://www.ncbi.nlm.nih.gov/genome/). *** ND, no data.

The cells of the isolates were rod-shaped, ovoid, coccoïd, or pleomorphic (Figure 2). The cell size of most strains varied depending on the growth medium and cultivation stage. The formation of small filterable cells was probably the result of limited concentrations of available carbon sources in the groundwater.

The small filterable cells of strains JR2/1-6 and J1-13Tb were <0.1 μm³ in volume; the genomes of the closest type strains, S. amurskyense KMM 3673T and C. haifense H38T, had a small size characteristic for ultra-microbacteria (both 2.8 Mb) (Tables 2 and 3). However, larger cells were also present in the cultures. Cells of strain JR3/2-13 possessed a volume of <0.1 μm³ and retained this volume irrespective of growth conditions, but genome size of the closest type strain Microbacterium oxydans DSM 20578T was 3.9 Mb. This value exceeded the 3.5-Mb limit accepted for ultramicrobacteria. Thus, these filterable bacteria are facultative ultramicrobacteria or ultramicroforms. The remaining three strains of filterable Actinobacteria, JR2/2-1, JR43/78-4-13, JR1/1-8b, with a normally large cell size possibly decreased cell size under pronouncedly starvation conditions of the groundwater similar to what was shown for the majority of planktonic bacteria [1].
### Table 3. Phenotypic characteristic and taxonomy of the strains isolated from groundwater samples.

| Characteristic | JR3/2-13 | J1-13Tb | JR43/78-4-13 | JR2/1-6 | JR2/2-1 | JR3/2-4 | J1-4 | JR1/8b | JR1/69-2-13 | JR1/69-3-13 | JR1/69-1-13 |
|----------------|---------|---------|-------------|---------|---------|---------|------|-------|-------------|-------------|-------------|
| **Cells morphology** | Rods | Rods, cocci | Cocci, rods | Rods | Rods | Pleomorphic | Pleomorphic | Cocci, rods | Rods | Rods | Cocciobacilli |
| **Cell size, μm** | 0.1–0.2 × 0.4–0.9 | 0.4–0.9 × 1.3–2.5; 0.3 | 0.3 × 1.3–2.0 | 0.8–1.5 × 1.6–2.4 | 0.9–1.2 × 1.2–4.7; 0.9–2.5 | 1.1–1.2 × 1.5–3.1; 0.9–1.5 | Cocci 0.7–1.6 | 0.8–1.2 × 1.5–2.1 | 0.7–1.3 × 0.9–2.1 | 1.2–1.6 × 1.5–2.4 |
| **Motility** | + | – | – | – | – | – | – | – | + | + | + |
| **Substrates for aerobic growth** | | | | | | | | | | | |
| Acetate | + | W | + | + | + | + | + | + | + | – |
| Oxalate | – | + | – | – | – | – | – | – | – | – |
| Glucose | + | + | – | + | + | + | + | + | + | + |
| Sucrose | + | + | – | + | + | + | + | – | W | W | + |
| Ethanol | W | – | + | + | + | W | – | – | – | – |
| Alanine | W | + | – | – | – | – | – | + | + | – |
| Lactate | + | + | – | + | + | W | + | W | W | + |
| Acetate + Fe³⁺ → Fe²⁺ | – | – | – | – | – | – | – | – | – | + |
| Acetate + NO₃⁻ → NO₂⁻ | – | – | + | – | – | – | W | + | + | + |
| Acetate + NO₂⁻ → N₂ | – | – | – | – | – | – | – | + | + | – |
| **Temperature, range/optimum, °C** | 5–42/30 | 5–40/30–32 | 4–50/28 | 5–37/25–28 | 13–42/35 | 21–42/33 | 13–35/28 | 28–38/28 | 5–35/30 | 5–35/30 | 13–42/30 |
| **NaCl, range/optimum (%) w/v** | 0–7/0.5–3 | 0–2/0–1.5 | 0–4 | 0–10/1–2 | 0–5(7)/0–2 | 0–5(7)/0–2 | 0–7(9)/0–3 | 0–8/0.5–2 | 0–3/0.5–1 | 0–5 0.5–3 | 0–3/0.5–1 |

+, positive growth; –, negative growth; W, weak reaction.
Figure 2. Morphology of the isolated strains as shown by transmission electron microscopy of ultrathin sections (a–c) and negatively stained whole cells (d–j): (a) *Salinibacterium* sp. JR2/1-6; (b) *Pusillimonas* sp. JR1/69-2-13; (c) *Microbacterium oxydans* JR3/2-13; (d) *Mycobacterium iranicum* JR-2/2-1; (e) *Chryseobacterium* sp. J1-13Tb; (f) *Rhodococcus qingshengii* JR 3/2-4; (g) *Pusillimonas* sp. JR1/69-2-13; (h) *Pusillimonas* sp. JR1/69-3-13; (i) *Roseomonas aestuarii* JR1/69-1-13; (j) *Paeniglutamicibacter sulfureus* JR1/1-8b. The cells were grown in TEG medium at 20–22 °C for 2–3 days. Scale bars in all panels correspond to 0.5 μm.

Glöckner and co-workers [50] used group-specific oligonucleotide probes to demonstrate that the cells of most of the hybridized freshwater actinobacteria are very small in situ.

The filterable bacterium *Chryseobacterium* sp. J1–13Tb had unique morphology. When grown on agarized media, strain J1-13Tb formed beige slimy colonies 2–4 mm in diameter. The cells were spherical, ovoid, bean-shaped, short rods (coccobacilli), or irregularly shaped rods (Figure 2e). The cells divided either by binary fission or formed small spherical buds laterally or at the poles. The size of spherical and kidney-shaped forms varied from 0.1 to 0.3 μm; undivided and pleomorphic cells were 0.4–0.9 μm in diameter and 1.3–2.5 μm long. Since the volume of small coccoid cells 0.3 μm in diameter was 0.014 μm³, they may be described as nanoforms, which explains their occurrence among filterable bacteria.

All isolates were aerobic organotrophs capable of using acetate, which was present in groundwater, as well as sugars, alcohols, and organic acids (Table 3). Five strains grew anaerobically with acetate, reducing nitrate to nitrite or to dinitrogen. Most strains were psychrotolerant and grew at temperatures of ~10 °C, which was an indication of their adaptation to the environment.

3.4. Tolerance of the Isolates to Heavy Metals and Ionizing Radiation

In view of wide occurrence of members of the genus *Pusillimonas* in radionuclide-contaminated groundwater, their resistance to heavy metals (Zn²⁺; Cr⁶⁺; Cu²⁺; Cd²⁺; and U⁶⁺) and γ-radiation was studied. *Roseomonas aestuarii* strain JR1/69-1-13 was used for comparison. Both *Pusillimonas* strains, JR1/69-2-13 and JR1/69-3-13, exhibited high resistance to cadmium, chromium, and uranium.
(exceeding 400 mg U\(^{6+}\) L\(^{-1}\)) (Table 4). The lethal chromium concentration (LD\(_{100}\)) for \textit{Pusillimonas} sp. JR1/69-3-13 was rather high (400 mg L\(^{-1}\)). \textit{Roseomonas aestuarii} JR1/69-1-13 was less resistant to the tested heavy metals (including uranium) than \textit{Pusillimonas} strains. All three strains were highly resistant ionizing radiation (Figure A3).

### Table 4. Tolerance of isolated strains to heavy metals and ionizing radiation.

| Strain                  | LD\(_{50}\), LD\(_{100}\) Radiation Dose (kGy) | Maximum Concentration (mg L\(^{-1}\)) | Zn\(^{2+}\) | Cr\(^{6+}\) | Cu\(^{2+}\) | Cd\(^{2+}\) | U\(^{6+}\) |
|-------------------------|--------------------------------------------|--------------------------------------|------------|-----------|------------|-----------|------|
| \textit{R. aestuarii} JR1/69-1-13 | LD\(_{50}\) 5.0                           |                                      | 10         | 25        | 50         | 50        | 10   |
|                         | LD\(_{100}\) 15.0                          |                                      | 100        | 100       | >200       | 200       | 100  |
| \textit{Pusillimonas} sp. JR1/69-2-13 | LD\(_{50}\) 1.0                           |                                      | 50         | 25        | 100        | 100       | 250  |
|                         | LD\(_{100}\) 10.0                          |                                      | 200        | 200       | >200       | 300       | >400 |
| \textit{Pusillimonas} sp. JR1/69-3-13 | LD\(_{50}\) 1.0                           |                                      | 50         | 100       | 100        | 100       | 250  |
|                         | LD\(_{100}\) 10.0                          |                                      | 200        | 400       | >200       | 300       | >400 |

#### 3.5. Metabolic Potential of \textit{Pusillimonas} and \textit{Roseomonas} Strains, as Assessed by Physiology and Genomes Analyses

Members of the genus \textit{Pusillimonas}, which were numerically predominant among the 16S rRNA gene sequences in the libraries from most strongly contaminated water of wells 14/68 and 2/68, were isolated in pure cultures for in-depth investigation. Two \textit{Pusillimonas} strains, JR1/69-2-13 (=VKM B-3222 = KCTC 62614) and JR1/69-3-13 (=VKM B-3223 = KCTC 62615), grew within the range of 5–35 °C with the optimum at 30–35 °C, and thus probably were psychrotolerant bacteria. They grew at pH 7.0, in the presence of 0%–3% and 0%–5% (w/v) NaCl with optimums at 1% and 2% NaCl, respectively. Carbon substrates utilized for aerobic growth by both strains included D-glucose, D-fructose, D-maltose, sucrose, D-xylene, acetate, \(n\)-valerate, fumarate, DL-lactate, L-malate, pyruvate, L-phenylalanine, and L-proline. Strain JR1/69-2-13 also utilized L-arabinose, D-mannose, L-rhamnose, D-ribose, butyrate, L-alanine, and gluconate; strain JR1/69-3-13 utilized D-galactose and succinate. Under anaerobic conditions in the medium with acetate strain JR1/69-3-13 reduced nitrate to N\(_2\), while strain JR1/69-2-13 was incapable of reducing nitrate, but reduced nitrite to N\(_2\).

Genomes of \textit{Pusillimonas} sp. JR1/69-2-13 and \textit{Pusillimonas} sp. JR1/69-3-13 were sequenced and annotated with the goal of elucidation of their taxonomic position and identification of the genetic determinants providing for their occurrence in the environment contaminated with radioactive waste [42].

The genome of the strain JR1/69-2-13 consists of 4,307,924 bp and comprises 33 scaffolds, with an overall G + C content of 57.2%. The genome of the strain JR1/69-3-13 consists of 4,541,669 bp and comprises 61 scaffolds, with an overall G + C content of 57.9%. The genomes of the strains JR1/69-2-13 and JR1/69-3-13 were predicted to include 4108 (98.5%) and 4260 (98.6%) protein-coding genes, respectively. Among the protein-coding genes, 85.0% and 85.3%, respectively, were assigned to putative functions, and the remaining genes were annotated as hypothetical proteins. The distribution of genes into COG functional categories is summarized in Figure A4.

The genes involved in central carbon metabolism in both strains included a set of genes encoding glycolysis/gluconeogenesis, the Entner-Doudoroff pathway, the pentose phosphate pathway, and the tricarboxylic acid (TCA) cycle. In JR1/69-2-13 genome, the Calvin-Benson cycle of CO\(_2\) fixation was revealed, including the large and small subunits of I Rubisco, Rubisco activase, fructose-1,6-biphosphatase and phosphoribulokinase (PLC53897–PLC53901), which were absent in the JR1/69-3-13 genome.
The genomes of both strains encoded the enzymes necessary for assimilatory sulfate reduction, including sulfate adenylyltransferase subunit 1 (PLC54533, PLC49270), sulfate adenylyltransferase subunit 2 (PLC54704, PLC49269), phosphoadenosine phosphosulfate reductase (PLC54534, PLC49268), sulfite reductase alpha subunit (PLC55466, PLC51459), and sulfite reductase beta subunit (PLC55268, PLC51319). Interestingly, JR1/69-2-13 genome harbors the complete set of sulfur-oxidizing genes including soxA (PLC53471), soxB (PLC53691), soxC (PLC53692), soxD (PLC53474), soxX (PLC53470), soxY (PLC53473), and soxZ (PLC53472) for thiosulfate oxidation. Sox multienzyme complex was not revealed in the JR1/69-3-13 genome.

The denitrification gene cluster included the genes predicted to encode the enzymes for sequential reduction of nitrate to dinitrogen gas, consistent with the complete denitrification phenotype of the strain JR1/69-3-13 (Table A3). In the JR1/69-2-13 genome, the genes encoding nitrate reductase were absent, but the genes encoding nitrite reductase (Nir), nitric oxide reductase (Nor), and nitrous oxide reductase (Nos) were revealed.

Strain JR1/69-2-13 reduced selenite under anoxic conditions, and the genes responsible for selenite transport into the cell and selenite reduction were revealed in the genome. Numerous genes responsible for heavy metal tolerance and detoxification were identified in the genome of JR1/69-2-13, including those encoding copper resistance protein CopB, copper chaperone, copper-translocating P-type ATPase, cobalt-zinc-cadmium resistance protein CzcA, mercuric reductase, DNA gyrase subunit A and B involved in fluoroquinolone resistance, various polyols ABC transporter and DedA protein involved in the uptake of selenate and selenite, an arsenate reductase and mercuric reductase. In the genome of strain JR1/69-2-13, the genes for cobalt-zinc-cadmium resistance protein CzcA, chromate resistance protein ChrB, and the arsenate reductase coding gene were identified. Detection of an arsenate reductase coding gene suggests that the reduction capabilities of both strains are broader than expected.

Our previous study [42] demonstrated that average nucleotide identity (ANI) values of JR1/69-2-13 and JR1/69-3-13 genomes with genomes of *Pusillimonas* strains were within the range of 77.6%–84.5%. These ANI values were below the species border, proposed to be 95%–96% [51]. In silico DNA-DNA hybridization (dDDH) values against reference genomes of *Pusillimonas* and *Candidimonas* strains were within the range of 19.3–28.8, and also were below the 70% threshold to differentiate bacterial species [52].

Thus, analysis of genomes of the strains JR1/69-2-13 and JR1/69-3-13 testifies to their belonging to two novel species within the *Pusillimonas* genus with proposed names ‘*Pusillimonas nitritireducens*’ and ‘*Pusillimonas subterraneus*’, respectively [42]. The genome sequences of both strains provide an insight into the genomic basis of their resistance to heavy metals and metalloid and ability to survive in nitrate-contaminated groundwater.

**Roseomonas aestuarii** JR1/69-1-13 (=VKM B-3221) was another strain isolated from the 1/69 groundwater sample. Strain JR1/69-1-13 was a Gram-negative, aerobic chemooorganoheterotrophic bacterium, motile in the early exponential phase, and forming orange pigmented colonies. The strain was able to grow at temperatures ranging from 13 to 42 °C with an optimum at 35 °C, at pH 7.0–7.5, in the presence of 0%–3% (w/v) NaCl with an optimum at 0.5%–1% (w/v) NaCl. The strain utilized sugars, amino acids, aromatic compounds, and organic acids. Under anaerobic conditions in the medium with glucose, strain JR1/69-1-13 reduced nitrate to nitrite. The genome of the strain consisted of 5,195,849 bp with an overall G + C content of 71.5% and contained 4849 genes, of which 4650 were coding DNA sequences, 52 coded tRNAs, and 4 belonged to ncRNAs [43]. Numerous genes responsible for utilization of various organic substrates (sugars, amino acids, organic acids, and chlorinated aromatic compounds), reduction of nitrate to nitrite, and heavy metal tolerance and detoxification were identified in the genome of *R. aestuarii* JR1/69-1-13.

4. Discussion

Use of uranium for fuel production and weapons manufacturing has resulted in widespread environmental contamination in the world [15,53]. Microorganisms are known to affect the migration of both radioactive and non-radioactive waste components [54–56]. Investigation of
radionuclide-contaminated environments is therefore important both for determination of the composition and functional activity of microbial communities, and for development of biotechnologies for remediation of contaminated environments.

The present work reports the results of the investigation of the physicochemical and radiochemical conditions and microbial composition in nitrate- and radionuclide-contaminated groundwater samples collected near Lake Karachai (Russia), which has been used for storage of intermediate-level liquid radioactive wastes of PA “Mayak” in the Chelyabinsk region [15]. We also aimed at the isolation and identification of bacteria, including those with ultra-small cell size, and determination of the physiological and genomic characteristics responsible for their adaptation to this environment.

Groundwater samples were collected from the depth of 20–80 m using the observation (monitoring) wells located at weakly contaminated and strongly contaminated zones. The studied groundwater samples had a high redox potential (Eh from +110 to +420 mV), their pH varied from 6.9 to 8.0. They had higher total mineralization and higher concentrations of nitrate (up to 13.6–42.7 g L\(^{-1}\)), acetate (up to 921 mg L\(^{-1}\)), and radionuclides (Table 1) than the uncontaminated natural water [48]. Many systems with these contaminants are highly acidic because of the nitric acid contamination. At the Lake Karachai measures were taken to establish the hydrochemical regime promoting radionuclide retention. Acidic solutions were always neutralized. For this purpose, solutions with high concentrations of metal hydroxides (mainly those of aluminum, iron, and manganese) with pH > 8 were dumped into the lake until 1985. Soda ash was also added. According to the data of many years, pH of the water varied from 7.2 to 9.8 with the average value of 8.7 [57].

Cultured aerobic organotrophic and nitrifying bacteria, as well as anaerobic denitrifying and iron-reducing prokaryotes were revealed in the groundwater samples. The numbers of sulfate-reducing bacteria and methanogens, as well as the rates of sulfate reduction and methanogenesis, were low or undetected, probably due to the high redox values and high nitrate concentration (Table 1, Figure A1).

Pyrosequencing of the 16S rRNA genes revealed that the share of archaeal sequences in the 176/94 library did not exceed 0.5%, and they belonged to ammonium-oxidizing archaea of the genera *Nitrosopumilus* and *Nitrososphaera* (*Thaumarchaeota*), uncultured *Pacearchaeota*, and methanogens of the genera *Methanoregula* and *Methanothrix* (*Euryarchaeota*) (Figures 1 and A2a, Table A1). The sequences of *Nitrosopumilus* and *Nitrososphaera* were predominant also in the three clone libraries obtained, which showed the presence of the archaea involved in the nitrogen cycle in contaminated groundwater. Ammonium-oxidizing members of the phylum *Thaumarchaeota* were also revealed in a subsurface radioactive thermal spring, the Franz-Josef-Quelle in Bad Gastein, Austria [58]. Uncultured *Pacearchaeota* characterized by small cells and small genomes were found in an aquifer amended with acetate at the Colorado River, near Rifle (CO, USA). Castelle and co-workers [13] performed metagenomic analyses of *Pacearchaeota* found in this aquifer amended with acetate and showed that these archaea were characterized by small cells and small genomes, and “involved in anaerobic carbon and hydrogen metabolism, probably associated with symbiotic and/or fermentation-based lifestyles.”

Our results correlate with the data of metagenomic studies of microbial communities from contaminated groundwater (FW106 site) sampled at the Oak Ridge Integrated Field Research Center (OR-IFRC) [59]. At the FW106 site, groundwater was also characterized by circumneutral pH, high concentrations of nitric acid, uranium and other radionuclides, heavy metals, and organic solvents. Microbial diversity in contaminated groundwater decreased significantly, with predominance of *Rhodanobacter* populations adapted to the combination of stresses at the site, in comparison with more diverse pristine groundwater community having a higher functional and metabolic potential. Moreover, denitrifying bacteria of the genus *Rhodanobacter* dominated also in groundwater with low pH at the FW106 site [55].

Microbial community in ethanol-amended sediments from Area 1 in Oak Ridge with acidic groundwater (pH ranging from 3.0 to 6.8) and high concentrations of nitrate, U(VI), Tc(VII), and sulfate was enriched with members of the genus *Castellaniella* (class Betaproteobacteria), which are
capable of nitrate reduction [60]. In another experiment at Area 1, addition of ethanol or glucose and pH neutralization of initially acidic groundwater with low concentrations of nitrate resulted in an increase of metal-reducing bacteria of the genera *Anaeromyxobacter* and *Geobacter* of the *Deltaproteobacteria* in acidic subsurface sediments [54]. These studies show that nitrate and radionuclides pollution decreases microbial diversity and stimulates growth of the community components best adapted to the conditions of the contaminated habitat [54,60].

The diversity of bacteria was highest in weakly contaminated groundwater from wells 1/69 and 176/94 (Figures 1b and A2b, Table A1). Among the sequences revealed in the libraries obtained from groundwater, those of *Candidatus* Parcubacteria (former OD1, 34.5% sequences in the 1/69 library), which include bacteria with ultra-small size of cells and reduced metabolic capacities [9], were revealed, as well as the sequences of *Pusillimonas*, *Polynucleobacter*, *Microbacterium*, *Chryseobacterium*, *Elusimicrobiurn*, and ‘*Candidatus Pelagibacter*,’ which also include bacteria with small or ultra-small size of the cells (Figure 1b, Table A1). In the samples of highly contaminated groundwater from wells 2/68 and 14/68, members of the genus *Pusillimonas* predominated (18.2% and 97.1% sequences in the libraries, respectively).

Aerobic chemoorganotrophic ultramicrobacteria of the genus *Polynucleobacter*, which utilize a narrow range of substrates (acetate, pyruvate, malate, etc.) under aerobic conditions or grow as facultative anaerobes, are known to be common in freshwater environments [61]. Chemoorganotrophic marine bacteria of the SAR11 clade ‘*Candidatus Pelagibacter ubique*’ are the Earth’s most abundant organisms [22]. Detection of these freshwater and marine bacteria in the studied samples is hardly occasional, considering the broad ranges of salinity and composition in the groundwater affected by Lake Karachai contaminated water.

Bacteria belonging to *Dechloromonas*, *Acinetobacter*, uncultured *Coxiellaceae*, *Candidatus* Parcubacteria, and undefined *Comamonadaceae* retrieved in the libraries from groundwater of the Karachai area were also revealed in uranium-contaminated groundwater amended with acetate sampled from an aquifer at Rifle, CO, USA [62]. Genome sequences of *Candidatus* Parcubacteria (OD1) were reconstructed from groundwater sampled in Rifle [5,6,11]. Sequences of *Candidatus* Parcubacteria (OD1) were also revealed in groundwater samples in the Karachai area. Thus, similar physicochemical and radiochemical conditions resulted in development of similar microbial communities.

Presence of the sequences of small-celled bacteria in the 16S rRNA gene libraries stimulated their isolation and studying. Using a selective filtration enrichment procedure and long-term incubations at low temperatures as recommended by Carini et al. [22], we isolated the strains *Janibacter limosus* JR43/78-4-13, *Microbacterium oxydans* JR3/2-13, *Paeniglutamicibacter sulfureus* JR1/1-8b, *Mycolicibacterium iranicum* JR2/2-1, *Chryseobacterium* sp. J1-13Tb, and *Salinibacterium* sp. JR2/1-6. Aerobic and denitrifying bacteria *Roseomonas aestuariii* JR1/69-1-13, *Rhodococcus qingshengii* JR3/2-4, *Rhodococcus fascians* J1-4, and *Pusillimonas* sp. strains JR1/69-2-13 and JR1/69-3-13 were isolated without the filtration stage. Although the strains with ultra-small cell size (below 0.1 μm³) were isolated (Table 3, Figure 2) cell size of most strains varied depending on the growth medium and cultivation stage. These filterable bacteria were facultative ultra-microbacteria or ultra-microforms. Formation of small filterable cells was probably the result of limited concentrations of available carbon and energy sources in the groundwater. Reduction in cell size caused by nutrient and energy limitations was shown for microorganisms from other oligotrophic habitats [63].

All isolates were aerobic organotrophs utilizing sugars, alcohols, and organic acids, as well as acetate, which was present in groundwater (Table 3). Five strains grew anaerobically with acetate, reducing nitrate to nitrite or to dinitrogen. Numerous stress factors in the studied groundwater (high concentrations of nitrate, radionuclides, and metals, as well as radioactivity) resulted in decreased microbial diversity and predominance of the *Pusillimonas* population adapted to the conditions of the habitat. *Pusillimonas* strains were resistant to heavy metals (Zn²⁺, Cr⁶⁺, Cu²⁺, Cd²⁺, and U⁶⁺) and γ-radiation (Table 4, Figure A3). The resistance capability of *Pusillimonas* and *Roseomonas* strains to the four heavy metals tested was much higher than that of *E. coli*. The
maximum tolerance concentration to a single heavy metal for \( E. coli \) was for \( \text{Cd}^{2+} (56 \text{ mg L}^{-1}) \), \( \text{Pb}^{2+} (41 \text{ mg L}^{-1}) \), \( \text{Cu}^{2+} (64 \text{ mg L}^{-1}) \), and \( \text{Zn}^{2+} (65 \text{ mg L}^{-1}) \) [64]. The maximum tolerance concentrations reported for \( \text{Pseudomonas} \), \( \text{Cupriavidus} \), \( \text{Bacillus} \), and \( \text{Acinetobacter} \) strains isolated from heavy-metal-polluted soil to \( \text{Cd}^{2+} \), \( \text{Pb}^{2+} \), \( \text{Zn}^{2+} \), and \( \text{Cu}^{2+} \) were 200, 1600, 600, 265 \text{ mg L}^{-1} \), respectively [65]. Genome sequencing of denitrifying bacteria of the genus \( \text{Pusillimonas} \) made it possible to determine their taxonomic position and genetic determinants providing for their occurrence in the environment contaminated with radionuclides and nitrates (Tables 2 and A3, Figure A2b).

It should be noted that occurrence of the metal resistance genes is typical of microorganisms from radionuclide-contaminated environments. In plasmidome analysis of groundwater samples from the Oak Ridge Field Research Center (Oak Ridge, TN, USA), which were also characterized by low cell counts but diverse microbial communities, several hundred circular plasmids were identified bearing the genes involved in metal, antibiotic, and phage resistance [66].

5. Conclusions

Groundwater microorganisms from the Lake Karachai region used for the disposal of liquid radioactive waste were investigated. Inflow of radioactive waste resulted in decreased microbial diversity in highly contaminated sites compared to the background groundwater samples. High-throughput sequencing and analysis of the clone libraries revealed lower microbial diversity in the most strongly contaminated groundwater and predominance of bacteria of the genera \( \text{Pusillimonas} \), \( \text{Polynucleobacter} \), \( \text{Candidatus Pelagibacter} \), and of the candidate phylum \( \text{Parcubacteria} \) in the most contaminated groundwater samples; these groups include species with ultra-small cell size. Pure cultures of aerobic organotrophic and denitrifying bacteria isolated from the groundwater samples and identified as members of the genera \( \text{Janibacter} \), \( \text{Microbacterium} \), \( \text{Paeniglutamicibacter} \), \( \text{Mycolicibacterium} \), \( \text{Chryseobacterium} \), \( \text{Salinibacterium} \), \( \text{Roseomonas} \), \( \text{Rhodococcus} \), and \( \text{Pusillimonas} \) were facultative ultra-microbacteria or ultra-microforms, possibly formed because of nutrient and energy limitations. The studied bacteria of the genus \( \text{Pusillimonas} \), which predominated in the groundwater were resistant to the environment (heavy metals and radioactivity) and capable of reducing nitrate; these findings were supported by their genomes analysis. Bacteria inhabiting groundwater of the Lake Karachai area, including the isolated small-celled strains, are able to grow aerobically or/and anaerobically. Degradation of acetate and organic components of waste and denitrification with formation of \( \text{NO}_2^- \), \( \text{N}_2\text{O} \), and \( \text{N}_2 \) are probably the most important microbial geochemical processes in the groundwater studied. A high abundance of denitrifying bacteria and decreasing nitrate concentrations in the groundwater of Lake Karachai area confirm the possible geochemical activity of these bacteria. Removal of nitrates from groundwater in the course of denitrification may result in decreased redox potential, which could stimulate other anaerobic processes, enhancing radionuclide reduction, and may decrease the rate of radionuclide migration.

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## Appendix A

Table A1. Diversity of the phylotypes of the bacteria and *Archaea* in 16S rRNA gene libraries of the microorganisms from groundwater.

| Super-Kingdom | Phylum, Class | Genus               | **Share of the Number of Sequences in the Library * from Well, %** |
|---------------|---------------|---------------------|---------------------------------------------------------------|
|               |               |                     | 176/94  2/68  1/69  14/68                                     |
|               |               |                     | **from Well, %**                                             |
| *Archaea*     |               |                     | **from Well, %**                                             |
| Crenarchaeota |               |                     | **from Well, %**                                             |
| Euryarchaeota | Methanobacterium | 4.8                 | 0  2.4  0  0                                                |
|               | Methanococcus  | 14/68               | 0  8.7  0  0                                               |
|               | Methanothrix   | 1.0                 | 0  0  0  0                                                |
|               | Pacearchaeota  | 42.3                | 19.4  8.9  0  0                                           |
|               |               |                     | **from Well, %**                                             |
|               |               |                     | **from Well, %**                                             |
| Thaumarchaeota | Nitrosopumilus | 30.7                | 63.4  66.9  81.0                                            |
|               | Nitrososphaera | 21.2                | 6.5  8.9  19.0                                             |
|               |               |                     | **from Well, %**                                             |
|               |               |                     | **from Well, %**                                             |
| Woesearchaeota | Incertae_Sedis_AR13 | 42.3                | 19.4  8.9  0  0                                           |
|               | Incertae_Sedis_AR15 | 0                   | 0  2.4  0  0                                             |
|               | Incertae_Sedis_AR18 | 0                   | 1.1  1.6  0  0                                           |
| Proteobacteria |               |                     | **from Well, %**                                             |
|               |               |                     | **from Well, %**                                             |
| Alphaproteobacteria | Novospingobium | 1.1                 | 0  0  0  0                                               |
|               | Reyranella    | 0                   | 5.1  0  0  0                                               |
|               | Brevundimonas | 0                   | 3.0  0  0  0                                               |
|               | Parvibaculum  | 0                   | 0  2.4  0  0                                               |
|               | Candidatus_Pelagibacter | 0           | 0  4.8  0  0                                           |
| Bacteria      | Marinicuda    | 0                   | 0  2.4  0  0                                               |
|               | Frigidibacter | 0                   | 11.1  0  0  0                                               |
|               | Inquilinus    | 0                   | 2.0  0  0  0                                               |
|               | Pseudaminobacter | 0           | 2.0  0  0  0                                           |
|               | Gallionella   | 6.9                 | 0  0  0  0                                               |
|               | Curvibacter   | 9.2                 | 0  0  0  0                                               |
|               | Rhodobacter   | 7.9                 | 0  0  0  0                                               |
| Taxonomic Group          | Percentage | 230 | 55  | 0  |
|--------------------------|------------|-----|-----|----|
| Sulfuritalea             | 5.6        | 0   | 0   | 0  |
| Limnohabitans            | 2.0        | 0   | 0   | 0  |
| Dechloromonas            | 1.6        | 0   | 0   | 0  |
| Polynucleobacter         | 1.5        | 0   | 14.3| 0  |
| Sideroxydans             | 6.4        | 0   | 0   | 0  |
| Sphingiscirra            | 0          | 6.1 | 0   | 0  |
| Pusillimonas             | 0          | 18.2| 0   | 97.1|
| Candidimonas             | 0          | 4.0 | 0   | 0  |
| Georgfuchsia             | 2.1        | 0   | 0   | 0  |
| Azospira                 | 1.8        | 0   | 0   | 0  |
|                          | 34.0       | 23.0| 5.5 | 0  |
| Uncultured Coxiellaceae  | 30.9       | 0   | 0   | 0  |
| Gammaproteobacteria      |            |     |     |    |
| Pseudomonas              | 0          | 2.0 | 0   | 0  |
| Nitrosococcus            | 0          | 0   | 2.4 | 0  |
| Lysobacter               | 0          | 9.1 | 0   | 0  |
| Hydrocarbinophaga        | 0          | 6.1 | 0   | 0  |
| Nitrospirae              |            |     |     |    |
| Nitrospira               | 0          | 0   | 8.3 | 0  |
| Bacteroidetes            |            |     |     |    |
| 2.1                      | 10.0       | 0   | 0   | 0  |
| Paracubacteria            |            |     |     |    |
| Incertae_sedis           | 0          | 0   | 34.5| 0  |
| Chloroflexi              |            |     |     |    |
| Levilinea                | 0          | 0   | 10.7| 0  |
| Fimicutes                |            |     |     |    |
| 1.4                      | 0          | 3.5 | 0   | 0  |
| Firmicutes               |            |     |     |    |
| Thermodesulfobium        | 0          | 0   | 2.4 | 0  |
| Acetobacterium           | 1.3        | 0   | 0   | 0  |
| Gemmatimonadetes         |            |     |     |    |
| Gemmatimonas             | 1.0        | 0   | 0   | 0  |

* The taxonomic groups comprising ≥1% of the total number of Archaea and Bacteria sequences are listed.
Table A2. Diversity indices of the 16S rRNA gene sequences of *Archaea* (A) and bacteria (B) in the libraries obtained by pyrosequencing and/or cloning of 16S rRNA genes generated from groundwater samples.

| Parameter                                      | Libraries from Observation Wells |
|------------------------------------------------|----------------------------------|
|                                                | 176/94-A | 176/94-B | 1/69-A | 1/69-B | 2/68-A | 2/68-B | 14/68-A | 14/68-B |
|                                                |          |          |        |        |        |        |         |         |
| Year of Sampling                               | 2012     | 2011     | 2010   | 2010   |        |        |         |         |
| Depth of Sampling, m                           | 60       | 44       | 80     | 80     |        |        |         |         |
| Library                                        | 176/94-A | 176/94-B | 1/69-A | 1/69-B | 2/68-A | 2/68-B | 14/68-A | 14/68-B |
| Number of sequences in the library             | 104      | 17,685   | 104    | 57     | 93     | 99     | 84      | 103     |
| Number of OTUs                                 | 15       | 927      | 14     | 16     | 21     | 60     | 5       | 4       |
| Shannon-Weaver diversity index (H)             | 2.21     | 5.08     | 1.27   | 2.58   | 2.54   | 3.65   | 0.85    | 0.16    |
| Simpson’s diversity index (1/S)                | 6.11     | 35.50    | 1.97   | 11.0   | 9.0    | 20.80  | 1.70    | 1.10    |
| Evenness                                       | 0.82     | 0.74     | 0.92   | 0.92   | 0.83   | 0.89   | 0.53    | 0.12    |
| Berger-Parker Dominance index (D)              | 0.32     | 0.13     | 0.70   | 0.21   | 0.23   | 0.15   | 0.75    | 0.97    |

* Data from paper Nazina et al. [21].

Table A3. Genes for denitrification found in the *Pusillimonas* sp. JR1/69-2-13 and *Pusillimonas* sp. JR1/69-3-13 genomes based on the annotation from IMG.

| Gene   | Locus Tag       | GenBank Identifier (pBLAST Best Hit) | % Identity |
|--------|-----------------|-------------------------------------|------------|
|        | JR1/69-2-13     | JR1/69-3-13                         | JR1/69-2-13 | JR1/69-3-13 |
| **narG**        | CR159_07195     | WP_073102996.1 Bordetella petrii     | -          | 91         |
| **narH**        | CR159_07190     | WP_008166145.1 Achromobacter arsenitoxydans | -          | 91         |
| **narI**        | CR159_07185     | WP_084658162.1 Chitinimonas taiwanensis | -          | 72         |
| **narJ**        | CR159_07180     | WP_050873229.1 Comamonas testosterone | -          | 81         |
| **narX**        | CR159_07165     | WP_088588207.1 Achromobacter marplatensis | -          | 64         |
| **narL**        | CR159_07160     | WP_076411578.1 Achromobacter xylosoxidans | -          | 77         |
| **narK1**       | CR159_07205     | WP_072427369.1 Chitinimonas taiwanensis | -          | 80         |
| **narK2**       | CR159_07200     | WP_028353905.1 Bordetella petrii     | -          | 80         |
| **nirK**        | CR155_06925     | CR159_13410                          | WP_084135858.1 Candidimonas bauzanensis | 88         | 87         |
| **nirK**        | CR155_06925     | CR159_19240                          | WP_073101649.1 Candidimonas bauzanensis | 87         | 88         |
| **norC**        | CR155_05890     | CR159_19250                          | WP_073101466.1 Candidimonas bauzanensis | 82         | 84         |
| Gene | Accession | Accession | Description | Count 1 | Count 2 |
|------|-----------|-----------|-------------|---------|---------|
| norB | CR155_05895 | CR159_19255 | WP_084135643.1 *Candidimonas bauzanensis* | 91      | 90      |
| norQ | CR155_05900 | CR159_19256 | WP_084135672.1 *Candidimonas bauzanensis* | 85      | 86      |
| norD | CR155_05905 | CR159_19257 | WP_073101467.1 *Candidimonas bauzanensis* | 74      | 76      |
| norZ | CR155_19235 | CR159_02315 | WP_073106495.1 *Candidimonas bauzanensis* | 91      | 90      |
| norA | CR155_19230 | CR159_02310 | WP_073106498.1 *Candidimonas bauzanensis* | 83      | 84      |
| norR | CR155_19225 | CR159_02305 | WP_073106501.1 *Candidimonas bauzanensis* | 89      | 89      |
| nosR | CR155_08950 | CR159_05710 | WP_073103310.1 *Candidimonas bauzanensis* | 80      | 81      |
| nosZ | CR155_08945 | CR159_05705 | WP_088147579.1 *Achromobacter denitrificans* | 89      | 90      |
| nosD | CR155_08940 | CR159_05700 | WP_073103312.1 *Candidimonas bauzanensis* | 79      | 80      |
| nosF | CR155_08935 | CR159_05695 | WP_073103313.1 *Candidimonas bauzanensis* | 84      | 83      |
| nosY | CR155_08930 | CR159_05690 | WP_073103314.1 *Candidimonas bauzanensis* | 89      | 88      |
| nosL | CR155_08925 | CR159_05685 | WP_084135995.1 *Candidimonas bauzanensis* | 72      | 66      |
| nosX | CR155_08920 | CR159_05680 | WP_073103316.1 *Candidimonas bauzanensis* | 66      | 68      |
Figure A1. Numbers of aerobic and anaerobic microorganisms in groundwater samples at the Karachai area (well 209, 2011 year of sampling; well 176/94, 2012; wells 1/69 and 14/68, 2013).

Figure A2. Taxonomic classification of archaeal (a) and bacterial (b) 16S rRNA gene fragments in the libraries from groundwater samples at the phylum level (at the class level for Proteobacteria) using the RDP classifier.
Figure A3. Resistance of the isolates to γ-radiation: 1, *R. aestuarii* JR1/69-1-13; 2, *Pusillimonas* sp. JR1/69-2-13; and 3, *Pusillimonas* sp. JR1/69-3-13. Error bars represent the standard deviation on the graph. The graph is representative of results obtained on three different replications.

Figure A4. Number of genes assigned to COGs categories in the genomes of *Pusillimonas* sp. JR1/69-2-13 and *Pusillimonas* sp. JR1/69-3-13.
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