Survivability of Soil and Permafrost Microbial Communities after Irradiation with Accelerated Electrons under Simulated Martian and Open Space Conditions

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Abstract: One of the prior current astrobiological tasks is revealing the limits of microbial resistance to extraterrestrial conditions. Much attention is paid to ionizing radiation, since it can prevent the preservation and spread of life outside the Earth. The aim of this research was to study the impact of accelerated electrons (~1 MeV) as component of space radiation on microbial communities in their natural habitat—the arid soil and ancient permafrost, and also on the pure bacterial cultures that were isolated from these ecotopes. The irradiation was carried out at low pressure (~0.01 Torr) and low temperature (−130°C) to simulate the conditions of Mars or outer space. High doses of 10 kGy and 100 kGy were used to assess the effect of dose accumulation in inactive and hypometabolic cells, depending on environmental conditions under long-term irradiation estimated on a geological time scale. It was shown that irradiation with accelerated electrons in the applied doses did not sterilize native samples from Earth extreme habitats. The data obtained suggests that viable Earth-like microorganisms can be preserved in the anabiotic state for at least 1.3 and 20 million years in the regolith of modern Mars in the shallow subsurface layer and at a 5 m depth, respectively. In addition, the results of the study indicate the possibility of maintaining terrestrial like life in the ice of Europa at a 10 cm depth for at least ~170 years or for at least 400 thousand years in open space within meteorites. It is established that bacteria in natural habitat has a much higher resistance to in situ irradiation with accelerated electrons when compared to their stability in pure isolated cultures. Thanks to the protective properties of the heterophase environment and the interaction between microbial populations even radiosensitive microorganisms as members of the native microbial communities are able to withstand very high doses of ionizing radiation.

Keywords: astrobiology; Mars; accelerated electrons; gamma radiation; microbial communities; radioresistance; native environment; soil; permafrost
1. Introduction

According to current knowledge, life on Earth exists within a limited range of the balanced impacts of physical and chemical factors. Significant changes in the amplitude of the impact can lead to the disturbances of biosystems up to the disappearance of life. Despite the long-standing scientific interest in the problem of living cell resistance to adverse impacts, and numerous biological studies of extreme habitats, the question of the limits of cell viability in native environment under stress conditions remains open, and in recent decades has acquired the scale of an astrobiological problem. The viability of microorganisms under the influence of extraterrestrial cosmic factors is one of the priority issues of modern astrobiological studies [1–3]. Much attention is paid to ionizing radiation being considered as one of the main factors that can prevent the preservation and spread of life outside the Earth [4–10]. Indeed, among physicochemical factors that change the natural habitat and, above all, organisms per se, ionizing radiation is undoubtedly of prior importance having a direct effect on the genetic material of a living cell.

The influence of ionizing radiation in relatively low doses on different organisms has been studied for a long time, and radiobiology generalizes and continues to accumulate experience of such studies [11–13]. Nevertheless, to plan and conduct astrobiological planetary studies, it is necessary to evaluate the effect of high doses of various components of cosmic radiation on microorganisms adapted to a native habitat [6–10]. In addition, the tasks of astrobiology require a differentiated approach to astrobiological models (Mars, ice satellites, Titan, etc.), taking into account hypothetical models of biospheres, as well as radiation fluxes and other data about the planet that characterize the environment.

It has been proved that the conditions under which irradiation occurs (temperature, pressure, etc.) substantially correct radiation effects [5–7,10]. Microorganisms in natural heterophase mineral media (soils, sediments) are well protected against the stressful effects of physical and chemical factors. A lot of data there is published already on the increased resistance to stress of microorganisms in biofilms or microbial communities in situ in native habitat in comparison with pure microbial cultures [6,14–18]. Therefore, in order to correctly assess the radioresistance of terrestrial microorganisms in the extraterrestrial regolith or other habitat, the closest possible reproduction of the relevant physicochemical factors is necessary, while preserving the natural environmental model, to which microbial communities are adapted.

Recent studies have shown that the resistance of microorganisms to gamma radiation is significantly underestimated [6–9,19]. Gamma radiation is a small part of cosmic radiation [5,10]. Different types of ionizing radiation differ in their biological effects [5]. For astrobiological modeling it is necessary to study in situ the stability of natural communities of microorganisms to various types of cosmic radiation in conditions that are as close as possible to extraterrestrial models. Elucidation of limiting doses and the identification of the most resistant in situ microorganisms or their communities will enable not only to correct the search tasks, but, despite the conventionality of extrapolating calculations to real conditions, it will allow for estimating the expediency of searching for life on a particular space object.

The aim of this research was to study the effect of accelerated electrons as a component of cosmic radiation at high doses of 10 kGy and 100 kGy on microbial communities in their natural habitat—arid soil and ancient permafrost, as well as on pure bacterial cultures that were isolated from these biotopes. Irradiation was carried out under low pressure (~0.01 Torr) and low temperature (~−130 °C) conditions, to simulate Martian and outer space environments. When considering the usage of different units of ionizing radiation dose in radiobiological literature we would clarify that gray (Gy) is defined as the absorption of one joule of radiation energy per kilogram of matter, and Gy is equal to 100 rad.

A study of the effect of elevated radiation doses, which significantly exceed the amplitude fluctuations of the radiation on the surface of Mars [20–23], is necessary for the extrapolation of biological effects on the geological time scale. Laboratory experiments do not allow for studying the long accumulation of doses of relatively low intensity in the microbial biomass of natural soil. The
main question of the present study is how long the biosphere of Mars could be maintained after the supposed catastrophic change in planetary conditions [24–29], the gradual loss of the atmosphere [30], and the formation of a modern climate.

2. Materials and Methods

2.1. Objects of Study

An arid soil sample (SN2) from the Negev desert and the ancient Arctic permafrost sample (M-1/91) were used as objects in the experiments, as well as pure cultures of bacteria *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60. The Arctic permafrost sample (M-1/91) was taken from the 1/91 well located in the Lower Kolyma Lowland (Yakutia) between the Bolshaya Chukochya and Malaya Konkovaja rivers in the sublatitudinal oler uplift [31]. The sample was taken at a depth of 34.0 m, from the ancient layers of the Oler suite, which did not melt for 1.8–2 million years. The temperature in the well was $-7 \ldots -12 \, ^\circ C$. Sampling and transportation methods and conditions were described earlier [32]. This sample was used earlier in our model experiments, with high doses of gamma irradiation at low temperature and low pressure [6,33].

The arid soil sample (SN2) was taken from the Negev desert in the Avdat (30°47’ N/34°46’ E) at a depth of 5–10 cm (horizon A) [34]. This area receives about 100 mm of precipitation during the whole year with months from June to October being the dry season with no precipitation at all. Soil-forming rocks are loess like loams. According to the WRB soil classification system, the soil is classified as Aridic Calcisols [35]. This sample was also used earlier in our model experiments with gamma irradiation at low temperature and low pressure [33].

The chemical characteristics of the samples are shown in Table 1. Ion concentrations in aqueous extracts from the samples were determined using Dionex ICS-1100 Ion Chromatography System (Dionex Corporation, Sunnyvale, CA, USA), according to manufacturer recommendations; total organic carbon content was determined using the method of oxidation in potassium dichromate [36].

| Attribute          | Sample SN2 | Sample M-1/91 |
|--------------------|------------|--------------|
| pH                 | 8.11       | 7.51         |
| NO$_2^-$, mg/kg    | Traces     | Traces       |
| NO$_3^-$, mg/kg    | 1.03       | 0.89         |
| NH$_4^+$, mg/kg    | 4.12       | 3.34         |
| Cl$^-$, mg/kg      | 58.15      | 49.84        |
| CO$_3^{2-}$, mg/kg | 330.41     | 129.29       |
| Na$^+$, mg/kg      | 512.7      | 87.57        |
| Mn$^{2+}$, mg/kg   | 1348.2     | 3.23         |
| Mg$^{2+}$, mg/kg   | 136.46     | 170.4        |
| K$^+$, mg/kg       | 877.5      | 40.47        |
| Fe$^{2+}$+Fe$^{3+}$, mg/kg | 4.35 | 29.55 |
| Total organic carbon, % | 1.27 | 0.32 |

Bacterial strains *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 were isolated previously from a sample of arid soil (sample SN), taken at the same location as the sample SN2. Sample SN and bacterial strains *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 were described in our previous papers [33,37–39]. It was shown that these strains are able to withstand gamma irradiation up to 10 kGy doses at low pressure (1 Torr) and low temperature ($-50 \, ^\circ C$) conditions, without a significant reduction in the number of living cells [38,39]. The strains are deposited to the Astrobiological Collection of bacteria of Lomonosov Moscow State University (the “Noah’s Ark” project, http://depo.msu.ru).
2.2. Preparation of Samples for Irradiation with Accelerated Electrons

Samples of arid soil SN2 (with ~40 g weight) and permafrost M-1/91 (with ~20 g weight) were incubated in a thermostat at +28 °C for 10 days before irradiation to activate microbial communities in situ and establish homeostasis in order to exclude the consequences of samples storage. Prior to incubation, the arid soil was moistened with sterile distilled water (up to 25% moisture by weight). Humidification of the permafrost sample M-1/91 was due to the melting of the ice contained in it. No nutrients were added. The samples were then incubated in tightly closed sterile polypropylene containers. After incubation, the samples were dried to an air-dry state at the same temperature for three days to prepare for the irradiation experiment under vacuum conditions.

The biomass of bacteria *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 in a stationary growth phase was suspended in sterile distilled water and the suspension was introduced into a sterile (heated at 600 °C for three hours) montmorillonite to immobilize the cells on the mineral matrix, simulating soil conditions. The samples were then thoroughly mixed with a sterile glass rod and dried for three days at +28 °C.

All of the objects under study were sieved through a sterile sieve with a hole diameter of 0.25 mm, then packed in sterile polyethylene bags (~20 mm in diameter). Each bag contained no more than 150 mg with the thickness of the sample layer being <1 mm. The bags were then hermetically sealed. To control possible contamination of the samples at all stages of the study, a control sample of sterile montmorillonite heated at 600 °C for three hours was prepared in the same way. Subsequently, this sample was transported, irradiated, stored, and analyzed together with the other samples.

2.3. Description of the Climatic Chamber

A climatic chamber was designed for the experiment (Figure 1). The chamber is a cylindrical tank with a “window” of aluminium foil (9) 100 μm thick on top. There are cylindrical cups that were installed inside the chamber (4) to reduce the pressure on the foil, and the samples were placed inside them and between them. The rubber ring (3) and a rigid metal ring tightens with 12 bolts (7) were used to seal the “window” of the foil. Under the bottom of the sample chamber there is a liquid nitrogen tank. To improve the thermal conductivity, rods (8) are welded to the bottom of the chamber. Taps (2) are used for pouring liquid nitrogen in. The chamber is pumped by a turbomolecular pump through a tap (6). Inside the chamber there is a thermocouple (5) for temperature control. The chamber together with the liquid nitrogen tank is thermally insulated with a syntepon bounded by an isolone cylinder (1). The chamber can maintain a pressure of ~0.01 Torr and a temperature of about −130 °C.

![Figure 1. Scheme of the climatic chamber: (a) Top view; (b) Side view. 1—Sintepon with isolon cylinder, 2—Tap for filling with liquid nitrogen, 3—Rubber ring, 4—Cylindrical cup, 5—Thermocouple, 6—Tap for air pumping out, 7—Bolt, 8—Rod, and 9—Foil.](image-url)
2.4. Irradiation of Samples with Accelerated Electrons

Samples in sealed polyethylene bags were placed in a pre-cooled to −100 °C irradiation chamber. The bags were then pierced quickly with a sterile steel needle to provide a balanced reduced pressure when exposed, and after that the chamber was immediately sealed and air was pumped out. The temperature during the irradiation was −130 °C with pressure being ~0.01 Torr. The irradiation was carried out using RTE-1B electron accelerator (USSR) with electron energy of ~1 MeV. Arid soil samples were irradiated with 10 kGy and 100 kGy doses, permafrost samples were irradiated with 100 kGy dose. The radiation intensity was 0.28 kGy/s and 2.8 kGy/s for 10 kGy and 100 kGy doses, respectively, and the duration of irradiation was ~36 s for both doses. The time from placing the samples in the chamber to unloading from the chamber (including the time of air pumping out, the onset of temperature equilibrium, the exit of personnel from the room, the starting and stopping of the accelerator, and the irradiation) was about 10 min. After irradiation the samples (in polyethylene bags) were unloaded into sterile polypropylene tubes, which were immediately placed in a cooled sterile thermos bottle. Samples were transported at a negative temperature (in thermos with cooling), and then stored at −18 °C.

In addition, control of low temperature and low pressure impact on the microorganisms was performed. The arid soil samples were placed in the chamber for 10 min at the same pressure and temperature as during irradiation. The absence of such control for permafrost, as well as permafrost irradiation with only one dose (100 kGy) was due to a small amount of available samples and the inaccessibility of similar ancient permafrost.

In the same way, pure bacterial cultures *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 immobilized in montmorillonite were irradiated with 10 kGy and 100 kGy doses. Control of low temperature and low pressure impact on the pure bacterial cultures was also performed.

2.5. Culturing of Microbial Communities

Determination of the number of culturable heterotrophic bacteria in the soil and permafrost samples was carried out by plating on glucose-peptone-yeast agar (GPY) (peptone—2 g/L, glucose—1 g/L, yeast extract—1 g/L, casein hydrolyzate—1 g/L, CaCO3—1 g/L, and agar-agar—20 g/L) [6]. Prior to inoculation, the microorganisms were desorbed from the mineral particles while using vortex Heidolph Multi Reax for 30 min at 2000 rpm. Suspensions of samples in different dilutions were plated in triplicate with simultaneous control of the nutrient medium sterility, sterility of the water used for dilutions preparing, and control of the presence of foreign air microflora. The plates were incubated at a temperature of +28 °C for two weeks.

2.6. Accounting for the Total Numbers of Prokaryotes

The total numbers of prokaryotic cells in the samples were determined using epifluorescence microscopy (EFM) with acridine orange dye. The cells were desorbed using ultrasound (22 kHz, 0.4 A, 2 min). The preparations for microscopy were made in six replicates and were fixed by heating, then stained, and viewed with a Biomed-6 PR LUM (Russia) microscope with ×700 magnification for 20 fields of view for each replication. Cells with a green fluorescence were counted. The water used for the dilution preparation was simultaneously examined as control. The numbers of prokaryotic cells in samples was calculated by the equation \[ N = \frac{S_1 \times a \times n}{V \times S_2 \times c}, \] where \( N \) is the number of cells per gram of soil; \( S_1 \) is the square of the preparation (µm²); \( a \) is number of cells in the field of view; \( n \) is the dilution index; \( V \) is the volume of the drop applied (mL); \( S_2 \) is the field of view of the microscope (µm²); and, \( c \) is the sample aliquot weight (g).
2.7. Multisubstrate Testing of Integral Metabolic Activity and Functional Diversity of Microbial Communities (MST)

Potential metabolic activity and the functional diversity of microbial communities have been investigated by the multisubstrate testing method [40–43]. Soil aliquots with 0.3 g weight were placed in 50 mL centrifuge cups, diluted with distilled water (1:100), and cells were desorbed using ultrasound (22 kHz, 0.4 A, 2 min). Mineral particles were then precipitated by centrifugation (2000 rpm, 2 min). A 2 mL of saturated solution of the substrate consumption indicator (dehydrogenase activity indicator triphenyltetrazolium bromide) was added to the 20 mL of supernatant, mixed, and inoculated 200 µL of the suspension into each well of a 96-well immunoassay plate containing a set of 47 test substrates [6,9] and mineral salts (the mineral base of the Czapek medium) [44]. The plates were incubated in a thermostat at +28 °C for 72 h. The water that was used for the dilution preparation was simultaneously examined as control. The optical density of the solutions in the wells was then photometrically measured at 510 nm wavelength, and the coefficients of functional biodiversity of the microbial community studied, which are characteristic features of its condition, were calculated using the Eco-Log software [45,46]. The analysis was performed in two replicates.

2.8. Identification of Bacterial Isolates

Pure bacterial cultures isolated from arid soil and permafrost samples were identified by sequencing of the 16S rRNA genes. Polymerase chain reaction (PCR) was performed with primers 63F+1387R [47], 27F+Un1492R [48,49], and 27F+537R [48,50] (Table A1 in Appendix A). The PCR products that were obtained were purified and sequenced by the Research and Production Company “Synthol” (Moscow, Russia) using primers 1100R and 537R [50] (Table A1). Editing of the nucleotide sequences was carried out using the Chromas Lite 2.01 program (http://www.technelysium.com.au). For alignment, comparison, and identification of nucleotide sequences, the Clustal Omega program (http://www.ebi.ac.uk/Tools/msa/clustalo/) and the BLAST algorithm of the GenBank database (http://blast.ncbi.nlm.nih.gov/Blast.cgi) were applied. The isolated strains were deposited in the Astrobiological Collection of bacteria of Lomonosov Moscow State University (http://depo.msu.ru) under the strain numbers KBP.AS.297–KBP.AS.303, KBP.AS.317, KBP.AS.319–KBP.AS.326, KBP.AS.340–KBP.AS.347, and KBP.AS.531–KBP.AS.537. The obtained nucleotide sequences were deposited in GenBank under the accession numbers MH050938–MH050968.

2.9. Investigation of the Physiological Characteristics of Isolates

The physiological properties of the isolated strains were studied by inoculation into 96-well immunoassay plates with liquid GPY medium supplemented with triphenyltetrazolium chloride as an indicator of metabolic activity. Phosphate (pH 2–6) and carbonate (pH 7–12) buffer systems were used to determine the range of pH values that are suitable for growth [51]. Resistance to salts presence was analyzed using liquid GPY nutrient media with the addition of NaCl, KCl, MgSO₄, or NaHCO₃ at concentrations of 2, 5, 10, 15, or 20%. The temperature limits of growth were determined by incubating cultures in liquid GPY nutrient media at temperatures of +2, +4, +10, +25, +37, +45, or +50 °C. The cultures were analyzed after ten days of incubation. At +2 and +4 °C the incubation times were 30 and 60 days, respectively. The physiological properties of the few strains were not studied because they grew in pure culture very poorly and had low biomass.
2.10. Irradiation of Pure Bacterial Cultures with Gamma Radiation

Several bacterial strains (KBP.AS.301, KBP.AS.319, KBP.AS.323, KBP.AS.324, KBP.AS.341, KBP.AS.343, KBP.AS.347) isolated from control and irradiated samples, and also strains *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60, as well as *Deinococcus radiodurans* VKM B-1422T were irradiated with gamma radiation under conditions that were close to normal. This was done to compare radioresistance of isolated pure cultures with that one the same bacteria irradiated *in situ* in soil and permafrost samples as well as for comparison of their radioresistance with the data of other authors. Prior irradiation bacterial biomass in stationary growth phase was immobilized in sterile kaolinite, as described in Section 2.2.

The irradiation was carried out on a gamma facility “Issledovatel” (USSR) with a $^{60}$Co source at a radiation intensity of 6.7 kGy/h, normal atmospheric pressure and a temperature of +37 °C (the temperature was measured with a thermocouple) with a doses gradient of 4.7, 9.3, 14, 18.6, 28, 37.2, and 46.5 kGy. The doses were selected in accordance with known data on the radiation resistance of bacteria [52,53], and the increased survivability of dried cultures [54,55]. Culturing from irradiated and control samples was performed, as described in Section 2.5.

3. Results and Discussion

3.1. Bacteria Abundance in the Samples

Exposure to low temperature and low pressure did not affect the number of colony forming units (CFU) and the total number of prokaryotes.

Irradiation with accelerated electrons led to a decrease in the CFU number in the arid soil and permafrost by a one and more than three orders of magnitude, respectively (Figure 2). The total number of prokaryotic cells (EFM) in arid soil decreased 1.5 and 20 times after irradiation with doses of 10 kGy and 100 kGy, respectively. In the permafrost sample, the total number of cells remained at the control level after irradiation with 100 kGy. That is, in terms of the total cells abundance the permafrost microbial community showed a higher survivability than the arid soil’s one, which may be due to the accumulation of high radiation doses by microorganisms during natural cryopreservation in the permafrost [56] or by special protective properties of the organomineral substrate. It is shown that bacteria are able to increase radioresistance when exposed to radiation [57,58]. A more drastic decrease in the number of CFU in permafrost can be associated with both cell death and their passage to the unculturable state. The transition of prokaryotes to the unculturable or dormant state appears to be a common adaptation in the microbial communities of permafrost [59,60]. In particular, a recently published study in the Dry Valleys of Antarctica [60] revealed that microbial permafrost communities are enriched in genes that are involved with dormancy and sporulation. In such habitats, bacteria can have an increased readiness to transition to the unculturable state under the influence of various stress factors.

Culturable cells were not found in the control sample of montmorillonite, which indicates that there was no contamination during the experiment. Biological activity in this sample was also not detected by the EFM and MST methods.

3.2. Metabolic Activity of Microbial Communities

Microbial communities of control samples of arid soil and permafrost had high functional diversity and high potential metabolic activity and was able to consume 26 and 23 substrates, respectively. Exposure to low pressure and low temperature without irradiation did not affect the metabolic activity of the soil microbial community (Table 2). Irradiation with even a minimal dose (10 kGy) led to a sharp suppression of the potential metabolic activity: microbial communities of all the irradiated samples consumed only one substrate—peptone. Perhaps this was due to the fact that the peptone contains many different amino acids, while the remaining substrates were represented by pure substances.
Accordingly, more different types of microorganisms are able to consume peptone in comparison with other substrates.

![Figure 2](image-url) **Figure 2.** Impact of irradiation, low pressure, and low temperature on microbial abundance in (a) Arid soil samples; (b) Permafrost samples. Gray columns—Number of colony forming units, white columns—Total number of prokaryotes (EFM). TPC (Temperature and Pressure Control)—The sample, exposed to the low pressure and low temperature without irradiation. The error bars are within the standard deviation, $p < 0.05$.

**Table 2.** Consumption of different classes of substrates by microbial communities at Multisubstrate Testing (MST). TPC—The sample, exposed to the low pressure and low temperature without irradiation.

| Sample       | Pentoses | Hexoses | Oligoses | Alcohols | Amino Acids | Salts of Carboxylic Acids | Polymers | Amides, Amines, Nucleosides |
|--------------|----------|---------|----------|----------|-------------|---------------------------|----------|----------------------------|
| SN2, Control | 526      | 3950    | 5340     | 4910     | 6186        | 3894                      | 2654     | 124                        |
| SN2, TPC     | 544      | 3972    | 5294     | 4862     | 6268        | 3754                      | 2716     | 132                        |
| SN2, 10 kGy  | 0        | 0       | 0        | 0        | 0           | 0                         | 122      | 0                          |
| SN2, 100 kGy | 0        | 0       | 0        | 0        | 0           | 0                         | 114      | 0                          |
| M-1/91, Control | 2150   | 1420    | 1912     | 2562     | 5670        | 4318                      | 3642     | 22                         |
| M-1/91, 100 kGy | 0    | 0       | 0        | 0        | 0           | 0                         | 146      | 0                          |

### 3.3. Taxonomic Affiliation and Physiological Characteristics of Bacterial Isolates

The diversity of bacterial complexes cultured from soil samples after exposure at low temperature and pressure without irradiation and after irradiation with doses of 10 kGy and 100 kGy (~0.01 Torr, $-130 \, ^\circ\mathrm{C}$) decreased slightly after irradiation (Table 3, Table A2 in Appendix B). Based on the morphological features, 10, 8, and 9 strains were isolated from the mentioned variants of the arid soil, respectively (Table 3). However, the taxonomic composition of cultured communities differed significantly after different impacts. Thus, representatives of the genera *Arthrobacter, Bacillus, Cryobacterium, Microbacterium, Micrococcus, Phenylbacterium, Saccharothrix, Sphingourantiacus,* and *Streptomyces* were found in the sample exposed at low pressure and temperature without irradiation. After irradiation with 10 kGy, bacteria belonging to the genera *Arthrobacter, Massilia, Micrococcus, Planomicrobium, Rufibacter,* and *Spirosoma* were cultured from the same arid soil, and after irradiation with a dose of 100 kGy the genera *Arthrobacter, Microvirga, Pontibacter, Rufibacter,* and *Spirosoma* were revealed. At the same time, strains with 100% similarity of DNA sequences and identical morphology of colonies and cells were found among microbial isolates from different variants of SN2 sample. In particular, identical representatives of the genus *Arthrobacter* (strains KBP.AS.346, KBP.AS.300,
KBP.AS.323) were found in all three variants. *Arthrobacter* of another species (KBP.AS.531, KBP.AS.533), as well as *Micrococcus* (KBP.AS.344, KBP.AS.534), identical with each other were detected both after exposure at low pressure and temperature without irradiation, and after irradiation with 10 kGy. In the arid soil samples that were irradiated with different doses, identical representatives of the genus *Spirosoma* (KBP.AS.298, KBP.AS.297) were found. Among all of the isolates, four strains (KBP.AS.345, KBP.AS.321, KBP.AS.535, KBP.AS.326) showed a fairly low similarity of the 16S rRNA gene sequences to the GenBank database (about 98% and below), which allows for assuming that these strains belong to previously not described bacterial species. A detailed study of these strains is being conducted now by the authors of the present research. An interesting fact is also the finding of *Cryobacterium* in the desert soil (strain KBP.AS.341). Bacteria of this genus were previously isolated from cold habitats [61–63]; there is no information on *Cryobacterium* culturing from hot habitats in the scientific literature.

**Table 3.** Taxonomic affiliation of bacteria isolated from arid soil and permafrost. The strains, marked with identical superscript numbers have similar colony morphology and identical nucleotide sequences. The strains highlighted in bold have <98% nucleotide sequences similarity with GenBank and possibly are the bacterial species not described earlier. TPC—The sample, exposed to the low pressure and low temperature without irradiation.

| Sample         | Strain—Genbank Accession Number | The Most Closely Related Sequences in Genbank—Similarity, % | Taxonomic Affiliation          |
|----------------|---------------------------------|-------------------------------------------------------------|--------------------------------|
| SN2, TPC       | KBP.AS.531 MH050938             | JX049673 *Arthrobacter* sp.—100                             | *Arthrobacter* sp. 1           |
|                |                                 | NR_026198 *Arthrobacter agilis*—99.5                        |                                 |
|                |                                 | NR_148833 *Arthrobacter echinis*—99.3                       |                                 |
|                |                                 | JQ419069 *Saccharothrix* sp.—99.8                           |                                 |
| SN2, TPC       | KBP.AS.532 MH050939             | NR_109447 *Saccharothrix echatensis*—99.5                   | *Saccharothrix* sp.            |
|                |                                 | NR_109103 *Saccharothrix hoggyrensis*—99.5                  |                                 |
| SN2, TPC       | KBP.AS.340 MH050940             | KT191026 *Phenylobacterium panocos*—99.5                    | *Phenylobacterium* sp.         |
|                |                                 | NR_117783 *Phenylobacterium muchangponense*—99.5            |                                 |
|                |                                 | DQ177485 *Microbacteriaceae bacterium*—99.6                 |                                 |
|                |                                 | KB857373 *Cryobacterium* sp.—99.4                           |                                 |
|                |                                 | FR691394 *Cryobacterium* sp.—99.4                           |                                 |
|                |                                 | NR_108605 *Cryobacterium arcticum*—99.4                     |                                 |
|                |                                 | JX205201 *Cryobacterium psychrotolerans*—99.2                |                                 |
| SN2, TPC       | KBP.AS.342 MH050942             | KGY280588 *Streptomyces fulvisimus*—99.3                    | *Streptomyces* sp.             |
|                |                                 | AJ781354 *Streptomyces mediolan*—99.3                       |                                 |
| SN2, TPC       | KBP.AS.343 MH050943             | NR_119202 *Microbacterium saperdae*—100                     | *Microbacterium* sp.           |
|                |                                 | NR_025405 *Microbacterium phyllosphaeraceae*—100             |                                 |
|                |                                 | GQ232449 *Bacterium RK91*—100                               |                                 |
| SN2, TPC       | KBP.AS.344 MH050944             | JN378531 *Micrococcus luteus*—99.9                          | *Micrococcus* sp. 2            |
|                |                                 | NR_134088 *Micrococcus aloeverae*—99.7                      |                                 |
|                |                                 | NR_116578 *Micrococcus yunnanensis*—99.6                    |                                 |
|                |                                 | DQ906916 *Uncultured bacterium clone*—99.4                   |                                 |
| SN2, TPC       | KBP.AS.345 MH050945             | NR_147725 *Sphingouarantiacus polygranulatus*—96.8         | *Sphingouarantiacus* sp.       |
|                |                                 | NR_148321 *Sphingomonas chloroacetidivorans*—95.4           |                                 |
|                |                                 | KT321369 *Sphingouarantiacus capsulatus*—95.3                |                                 |
|                |                                 | JX03970 *Arthrobacter oxidans*—100                           |                                 |
| SN2, TPC       | KBP.AS.346 MH050946             | NR_108849 *Arthrobacter siccitolerans*—99.7                 | *Arthrobacter* sp. 3           |
|                |                                 | NR_026192 *Arthrobacter polychromogenes*—99.6                |                                 |
| SN2, TPC       | KBP.AS.347 MH050947             | NR_117474 *Bacillus frigoritolerans*—100                     | *Bacillus* sp.                |
|                |                                 | NR_114919 *Bacillus simplex*—100                            |                                 |
| SN2, TPC       | KBP.AS.298 MH050948             | NR_113978 *Spirosoma rigui*—99.8                           | *Spirosoma* sp. 4             |
| SN2, 10 kGy    | KBP.AS.299 MH050949             | AB637009 *Uncultured bacterium clone*—100                   | *Rufibacter* immobiles         |
|                |                                 | HG316123 *Rufibacter immobilis*—99                          |                                 |
|                |                                 | CP012645 *Rufibacter tibetensis*—97.8                       |                                 |
| Sample          | Strain—Genbank Accession Number | The Most Closely Related Sequences in Genbank—Similarity, % | Taxonomic Affiliation       |
|-----------------|----------------------------------|------------------------------------------------------------|-----------------------------|
| SN2, 10 kGy     | KBP.AS.300 MH050950              | NR_042469 Arthrobacter phenanthrenivorans—100              | Arthrobacter sp. 3          |
|                 |                                  | NR_117535 Arthrobacter cryotolerans—100                    |                             |
|                 |                                  | NR_041546 Arthrobacter humidula—100                        |                             |
|                 |                                  | KY386623 Arthrobacter sp.—100                              |                             |
| SN2, 10 kGy     | KBP.AS.533 MH050951              | HQ860629 Uncultured bacterium clone—100                    | Planomicrobium sp. 5        |
|                 |                                  | GQ140340 Planomicrobium okeanokoites—99.9                  |                             |
|                 |                                  | NR_116601 Planomicrobium flavidum—99.7                     |                             |
| SN2, 10 kGy     | KBP.AS.301 MH050952              | SN2, 10 kGy KBP.AS.301 MH050952                            |                             |
|                 |                                  | NR_113978 Spirosoma rigui—99.8                            |                             |
| SN2, 10 kGy     | KBP.AS.302 MH050953              | EF507901 Spirosoma aquatica—99.8                          |                             |
| SN2, 10 kGy     | KBP.AS.303 MH050954              | NR_117040 Massilia consociata—99.0                         | Massilia sp.                |
|                 |                                  | NR_042502 Massilia aurea—98.4                             |                             |
| SN2, 10 kGy     | KBP.AS.334 MH050955              | NR_134088 Micrococcus aloveareae—99.8                      | Micrococcus sp. 2           |
|                 |                                  | NR_116578 Micrococcus yunnanensis—99.8                     |                             |
| SN2, 100 kGy    | KBP.AS.297 MH050956              | JX504809 Microvira vignae—98.7                            | Microvira sp.               |
|                 |                                  | NR_114298 Microvira aerilata—98.7                         |                             |
| SN2, 100 kGy    | KBP.AS.321 MH050957              | JF417863 Uncultured bacterium clone—97.9                   | Rufibacter sp.              |
|                 |                                  | HJ316124 Rufibacter immobile—96.8                          |                             |
| SN2, 100 kGy    | KBP.AS.322 MH050958              | JX504809 Microvira vignae—98.7                            | Microvira sp.               |
|                 |                                  | KC534446 Arthrobacter sp.—100                              |                             |
| SN2, 100 kGy    | KBP.AS.323 MH050959              | JX504809 Microvira oxydans—100                             | Arthrobacter sp. 3          |
|                 |                                  | XR_108949 Arthrobacter siccitolerans—99.7                   |                             |
|                 |                                  | XR_247636 Microvira sol—99.7                               |                             |
| SN2, 100 kGy    | KBP.AS.324 MH050960              | JF295810 Uncultured bacterium clone—99.9                   | Microvira soli             |
|                 |                                  | HJ316124 Rufibacter immobile—96.8                          |                             |
| SN2, 100 kGy    | KBP.AS.325 MH050961              | JR_042252 Arthrobacter parietis—100                        | Arthrobacter sp.            |
|                 |                                  | KPI25973 Arthrobacter subterraneus—100                     |                             |
|                 |                                  | FR691450 Pontibacter sp.—99.9                              | Pontibacter sp.             |
| SN2, 100 kGy    | KBP.AS.335 MH050962              | NR_148858 Pontibacter amylobiyticus—98.1                   | Microvira sp.               |
|                 |                                  | NR_116853 Pontibacter salisaro—97.3                        |                             |
| SN2, 100 kGy    | KBP.AS.336 MH050963              | KX247636 Microvira soli—99.4                               | Microvira sp.               |
|                 |                                  | XR_044563 Microvira guangxiensis—99.1                      |                             |
| SN2, 100 kGy    | KBP.AS.326 MH050964              | JJN037908 Uncultured Bacteroidetes bacterium—98.2           | Pontibacter sp.             |
|                 |                                  | KK355015 Pontibacter sp.—99.0                              |                             |
| M-1/91, 100 kGy | KBP.AS.317 MH050965              | NR_116722 Brevundimonas naejangsanensis—99.9               | Brevundimonas sp.           |
|                 |                                  | NR_113602 Brevundimonas diminuta—99.7                      |                             |
| M-1/91, 100 kGy | KBP.AS.319 MH050966              | NR_114986 Microbacterium maritipicum—100                    | Microbacterium sp. 7        |
|                 |                                  | KT889483 Microbacterium oxydans—100                         |                             |
| M-1/91, 100 kGy | KBP.AS.320 MH050967              | NR_114986 Microbacterium maritipicum—100                    | Microbacterium sp.          |
|                 |                                  | KT889483 Microbacterium oxydans—100                         |                             |
| M-1/91, 100 kGy | KBP.AS.337 MH050968              | NR_121739 Stenotrophomonas rhizophila—99.2                  | Stenotrophomonas rhizophila |
|                 |                                  | LJ906480 Stenotrophomonas maltophilia—97.9                 |                             |

Diversity of bacteria that were cultured from the irradiated permafrost sample was significantly lower than in arid soil. Four bacterial strains belonging to three genera were identified: *Brevundimonas*, *Microbacterium*, and *Stenotrophomonas*.

The isolation of bacteria of the genus *Arthrobacter* after irradiation is in complete agreement with the previously obtained data on their high radioresistance *in situ* to gamma radiation under the...
model conditions of Mars [6], as well as with data on their high radioresistance in pure culture [64] and with data on their high ultraviolet resistance [65]. There is also data on the high radioresistance of the Spirosoma [66], Pontibacter, Rufibacter, Microvirga [64], Micrococcus [67], Planomicrobium [64,68], Massilia [69], Brevundimonas [10], and Stenotrophomonas [70]. We have not been able to find information about the high radioresistance of Microbacterium. A number of studies have shown that bacteria of this genus have a high resistance to ultraviolet radiation and have multiple stress tolerance [70–73]. Thus, all the heterotrophic aerobic bacteria that were isolated from the samples irradiated with accelerated electrons belong to genera for which high radioresistance or high resistance to ultraviolet radiation is known.

Most of the isolated strains, including arid soil isolates, showed facultative-psychrophilic (psychotropic) properties, that is, they could grow in the temperature range extended to the low-temperature (Table 4). This can be both a manifestation of adaptation to storage and exposure of samples at low temperatures, and the increased resistance of psychrophiles to oxidative stress [74], and, consequently, to radiation. According to the data on resistance to pH, neutrophiles with wide pH-ranges of growth (4–12), alkalophiles (8–12), and acid-tolerant alkalophiles (4–8, 5–9) were found among isolates. Recall that the samples had a slightly alkaline pH (Table 1). The variety of pH-ranges of growth probably reflects the heterogeneity of the soil environment. However, it is noteworthy that most of the isolated strains are resistant to the acidity of the medium and grow in a wide pH range.

### Table 4. Physiological characteristics of bacteria isolated from arid soil and permafrost. TPC—The sample, exposed to the low pressure and low temperature without irradiation.

| Sample          | Strain          | Taxonomic Affiliation | Temperature Limits of Growth, °C | pH-range of Growth | Maximum Salt Concentrations at Which Growth Is Possible, % | NaCl | KCl | MgSO₄ | NaHCO₃ |
|-----------------|-----------------|-----------------------|----------------------------------|--------------------|-----------------------------------------------------------|------|-----|-------|-------|
| SN2, TPC        | KBP.AS.531      | Arthrobacter sp.       | 2–37                            | 5–9                | 2 2 15 0                                                   |      |     |       |       |
| SN2, TPC        | KBP.AS.532      | Saccharothrix sp.      | 10–37                           | 4–12               | 2 5 20 0                                                   |      |     |       |       |
| SN2, TPC        | KBP.AS.340      | Phenylbacterium sp.    | 10–37                           | 6–12               | 0 0 5 0                                                   |      |     |       |       |
| SN2, TPC        | KBP.AS.341      | Cryobacterium sp.      | 2–37                            | 7–11               | 20 15 20 0                                                 |      |     |       |       |
| SN2, TPC        | KBP.AS.342      | Streptomyces sp.       | 10–45                           | 4–12               | 10 10 20 5                                                 |      |     |       |       |
| SN2, TPC        | KBP.AS.343      | Microbacterium sp.     | 10–37                           | 4–11               | 10 15 20 0                                                 |      |     |       |       |
| SN2, TPC        | KBP.AS.344      | Micrococcus sp.        | 10–37                           | 5–8                | 0 5 0 0                                                   |      |     |       |       |
| SN2, TPC        | KBP.AS.346      | Arthrobacter sp.       | 2–50                            | 6–10               | 0 0 2 0                                                   |      |     |       |       |
| SN2, TPC        | KBP.AS.347      | Bacillus sp.           | 2–37                            | 5–8                | 0 5 0 0                                                   |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.298      | Spirosoma sp.          | 10–37                           | 5–12               | 20 15 20 0                                                 |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.299      | Rufibacter immobilis   | 2–37                            | 6–8                | 0 0 2 0                                                   |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.300      | Arthrobacter sp.       | 2–37                            | 6–8                | 0 0 2 0                                                   |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.533      | Arthrobacter sp.       | 10–37                           | 4–9                | 10 15 15 0                                                 |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.301      | Planomicrobium sp.     | 2–37                            | 5–12               | 0 0 15 0                                                  |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.302      | Planomicrobium sp.     | 2–37                            | 6–12               | 0 0 15 0                                                  |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.303      | Massilia sp.           | 10–37                           | 5–8                | 0 0 0 0                                                   |      |     |       |       |
| SN2, 10 kGy     | KBP.AS.534      | Micrococcus sp.        | 10–37                           | 5–12               | 20 15 20 0                                                 |      |     |       |       |
| SN2, 100 kGy    | KBP.AS.297      | Spirosoma sp.          | 10–37                           | 7–12               | 0 0 0 2                                                   |      |     |       |       |
| SN2, 100 kGy    | KBP.AS.323      | Arthrobacter sp.       | 4–37                            | 8–12               | 5 0 0 2                                                   |      |     |       |       |
| SN2, 100 kGy    | KBP.AS.324      | Microvirga sp.         | 4–37                            | 7–12               | 10 15 20 0                                                 |      |     |       |       |
| SN2, 100 kGy    | KBP.AS.325      | Arthrobacter sp.       | 2–50                            | 5–12               | 0 5 15 0                                                  |      |     |       |       |
| M-1/91, 100 kGy | KBP.AS.317      | Brevundimonas sp.      | 10–37                           | 5–12               | 20 15 20 0                                                 |      |     |       |       |
| M-1/91, 100 kGy | KBP.AS.319      | Planomicrobium sp.     | 10–37                           | 5–8                | 2 2 2 0                                                   |      |     |       |       |
| M-1/91, 100 kGy | KBP.AS.320      | Microbacterium sp.     | 4–50                            | 7–12               | 0 0 2 0                                                   |      |     |       |       |
| M-1/91, 100 kGy | KBP.AS.537      | Stenotrophomonas rhizophila | 2–37                           | 6–11               | 0 0 5 2                                                   |      |     |       |       |

Some strains were sensitive to the presence of salts in the medium, others were halotolerant and grew with a content of up to 20% NaCl or MgSO₄ and up to 15% KCl in the medium (Table 4). Bacterial strains practically did not grow on media containing NaHCO₃, even in small concentrations. Correlations between resistance to the studied physical and chemical factors (T, pH, salts) and radioresistance were not observed.
3.4. Irradiation of Pure Bacterial Cultures with Accelerated Electrons under Simulated Extraterrestrial Conditions

To assess the protective role of the natural environment and interpopulation relations in situ to protect microorganisms from radiation, irradiation of strains *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 with accelerated electrons under low pressure and low temperature conditions was carried out. Pure bacterial cultures were immobilized in a mineral matrix, according to the protocol described in Section 2.2.

Low temperature and low-pressure conditions had practically no impact on the number of CFU (Figure 3). After irradiation with a dose of 10 kGy, the number of CFUs of *A. polychromogenes* SN_T61 and *K. rosea* SN_T60 decreased by three and four orders of magnitude, respectively. After irradiation with a dose of 100 kGy, no cultured cells were detected. Pure cultures showed significantly less resistance to radiation than microbial communities in situ. Such a result can be caused by features of a heterogeneous soil environment, as well as by interactions of microorganisms within the community.

The soil is an extremely heterogeneous environment [75–77]. The physicochemical conditions in this environment can drastically change on a micro scale. That is, the soil is a combination of microzones with various conditions—different concentrations of water, oxygen, organic substances, salts, minerals, etc. As mentioned above, the physicochemical conditions at irradiation can affect the radiation damage of microorganisms. Probably, in the soil with its high heterogeneity, there should exist niches most favorable for the survival of bacteria upon irradiation. Also soils contain various organics, including pigments and microbial’s growth autoregulators. It is shown that these substances play an important role in the stress tolerance of microorganisms [18,78–81]. There is also information on the higher radioresistance of microorganisms in soils with high concentrations of humus, but these data are contradictory [82].

It is shown that bacteria within biofilms have increased resistance to stress in comparison with planktonic cells [16,17]. The protective role of intra- and interpopulation interactions can also be manifested through the production of some metabolites, as mentioned above. Moreover,
microorganisms possess the quorum-sensing, which also contributes to resistance to adverse effects, in particular, to oxidative stress [18,83].

These factors, as well as possibly some unrevealed factors, may correct the radioresistance of microorganisms in situ. However, it should be noted that the role of these factors in the radioresistance of natural microbial communities is currently poorly understood.

3.5. Survivability of Pure Bacterial Cultures under Gamma Irradiation

Another series of bacterial strains were irradiated with gamma radiation under normal conditions. Some of the strains were isolated from the studied arid soil and permafrost samples after irradiation with high radiation doses and after exposure to the simulated extraterrestrial conditions (Table 5). There were seven such strains, and *Arthrobacter polyochromogenes* SN_T61 and *Kocuria rosea* SN_T60 isolated earlier from arid soil (see Section 2.1) were irradiated along with them, and *Deinococcus radiodurans* VKM B-1422T strain was simultaneously irradiated for comparison. Viable cells of *D. radiodurans* VKM B-1422T were detected after irradiation with doses up to 28 kGy inclusive. Of the remaining nine strains, only two (*Bacillus* sp. KBP.AS.347 and *Planomicrobium* sp. KBP.AS.301) survived the irradiation with a minimal dose of 4.7 kGy, but these two strains did not withstand higher doses (Figure 4). It should be noted that resistance to 4.7 kGy dose allows for these strains to be considered radioresistant [11]. However, most strains did not show high radioresistance. Taking into account that some strains were isolated from the samples after irradiation with various high doses of accelerated electrons, it becomes evident that as the part of natural microbial communities, even radiosensitive microorganisms are able to withstand exposure to high radiation doses. This agrees with the data of Section 3.4 and also confirms the assumption that the natural organomineral heterophase medium has a protective function for the microorganisms inhabiting it, as well as, possibly, interrelationships in the community itself. Survival of soil microorganisms after exposure to high radiation doses (15 kGy) in the absence of the same high radioresistance in isolates was observed by other researchers [84]. In such studies, isolated strains that are offspring of irradiated in situ in soil microorganisms may carry residual genetic disturbances. However, as already mentioned, it has been experimentally shown that repeated irradiation of bacterial cultures leads to adaptation and enhancement, rather than a decrease in their radioresistance [57,58]. Thus, we demonstrated a much higher resistance of bacteria in situ in the composition of microbial communities in the natural habitat when compared to their resistance in pure culture.

| Sample          | Strain Number | Taxonomic Affiliation               |
|-----------------|---------------|-------------------------------------|
| SN2, TPC        | KBP.AS.341    | Cryobacterium sp.                   |
| SN2, TPC        | KBP.AS.343    | Microbacterium sp.                  |
| SN2, TPC        | KBP.AS.347    | Bacillus sp.                        |
| SN2, 10 kGy     | KBP.AS.301    | Planomicrobium sp.                  |
| SN2, 100 kGy    | KBP.AS.323    | Arthrobacter sp.                    |
| SN2, 100 kGy    | KBP.AS.324    | Microvirga sp.                      |
| M-1/91, 100 kGy | KBP.AS.319    | Microbacterium sp.                  |
| SN$^1$          | SN_T60        | Kocuria rosea                       |
| SN$^1$          | SN_T61        | *Arthrobacter polyochromogenes*     |
| The strain obtained from the VKM collection | VKM B-1422T | *Deinococcus radiodurans*          |

$^1$ The strain was isolated earlier in the course of other studies, see Section 2.1.

When comparing the radioresistance of *A. polyochromogenes* SN_T61 and *K. rosea* SN_T60 at irradiation under normal conditions and at low temperature and low pressure demonstrates a sharp decrease in radiation damage in the model conditions of Mars and outer space. The strains survived 10 kGy under model conditions, while retaining a high number of cells, but it did not survive 4.7 kGy...
under normal conditions. This is probably due to a decrease in the activity of free radicals with a decrease in temperature [10], as well as a decrease in the amount of water and oxygen, which are the main sources of free radicals [5,11], with a decrease in pressure. In addition, some differences may be due to the different types of radiation that we used.

The high radioresistance of *Planomicrobium* in pure culture was previously shown only twice [64,68]. Our data indicate that high resistance to radiation might be characteristic to the entire genus *Planomicrobium* (or for many of its species).

It should be emphasized how much the combination of Mars physical factors and the natural environment corrects radioresistance—the strains that survived *in situ* as a part of microbial community after 100 kGy irradiation (moreover, even those that retained high abundance after 100 kGy irradiation) did not survive irradiation with 4.7 kGy dose in pure culture under normal conditions. Thus, the microorganisms irradiated under simulated Martian conditions in soil *in situ* showed more than 20 times increased survivability when compared to pure bacterial cultures irradiated under normal conditions.

### 3.6. Implications for Habitability Assessment

The radiation resistance of microbial communities in our experiment was significantly higher than it is generally assumed [53,82,84,85]. This may be due to the reduction of radiation damage under low pressure and low temperature conditions [5,10], and was discussed in detail in already published papers [6,9].

The intensity of ionizing radiation on the surface of modern Mars is about 0.076 Gy/year [23]. 100 kGy radiation dose can be accumulated in the surface layer of regolith for 1.3 million years. This suggests that microorganisms in the modern cold conditions of Mars (in the absence of the possibility of actively repairing radiation damages) are able to remain viable for the specified period. At 5 m depth, the radiation intensity is about 5 mGy/year [20], and 100 kGy dose can be accumulated after 20 million years. These estimates of the possible duration of viable conservation of the hypothetical
deep Martian biosphere in the cryopreserved state coincide with those we indicated earlier [6]. It can also be assumed that viable microorganisms can be preserved in the ice of Europa for at least ~170 years at 10 cm depth, taking into account the data on the radiation intensity of ~600 Gy/year at this depth [5]. In this case, it should be noted that in our experiment the samples were dried, while the presence of water can significantly affect (advance) radiation damage [5,11]. The effect of ionizing radiation is also a limiting factor for the realization of some mechanisms of panspermia [5,86,87]. During a long journey of meteorites in outer space, hypothetical microorganisms that are located within them must accumulate high doses of radiation. It is estimated that most of the Martian meteorites reach Earth after thousands and millions of years of flight in open space [86]. Based on the radiation intensity inside meteorites, which is not more than 0.25 Gy/year depending on the size and material of the space body [87], the results of our study allow for hypothesizing the survival of microorganisms for at least 400 thousand years in open space (i.e., accumulation of 100 kGy dose for this period). These data can be used to estimate the probability of interplanetary life transport.

We note that the irradiation intensity that was used in the experiment is by the several orders of magnitude higher than in extraterrestrial conditions [20,23,86,87]. It is shown that in general the effectiveness of sparsely ionizing radiation (including accelerated electrons) dose is decreases with irradiation intensity decreasing due to radiation damage repairing by cells [5]. In low-temperature conditions of open space and Europa, a cells’ repairing seems unlikely [88]. So, differences in radiation intensity should not affect the survivability of microorganisms. But considering Mars, the possibility of cells’ repairing should be taken into account. There are a lot of studies testifies possibility of it. Microbial metabolism is revealed at temperatures at least down to $-33 \, ^\circ C$ [88,89]. Moreover, it is shown that econiches with liquid water and positive temperatures can occur on the current Mars [88,90–93]. Terrestrial microorganisms can grow under combined low pressure, temperature, and anoxic atmosphere conditions [94], it can withstand the impact of Martian salts and strong oxidizers [95–98], and can survive in subzero brines [99,100]. If even slow repairing of cells’ damages occurs, it can be of great effect during geological time. Thus, it can be concluded that, on Mars, the extension of viable cells’ in situ cryopreservation time should occur and be not fully compatible with our model experiment.

The age of Martian permafrost is evaluated at approximately more than 3 Ga [24]. At the current intensity of ionizing radiation during 3 Ga, the putative microorganisms in the cryopreserved state should accumulate ~230 MGy and ~15 MGy doses in the shallow layer of the regolith and at 5 m depth, respectively. This calculation suggests that survivability after irradiation with 100 kGy dose is not enough for viable cryopreservation of the putative ancient biosphere of early Mars till present. But, this dose was not sterilizing in our study—the true sterilizing doses may be significantly higher. Definition of the microbial communities’ radioresistance limit is in the scope of the further research. Nevertheless, survival of putative ancient biosphere over 3 Ga in the cryopreserved state without cell replication or metabolism (i.e., without radiation damages repairing) seems unlikely. However, there are some possibilities for cells’ repairing and viable cryopreservation time extension in conditions of the current Mars, as mentioned above. Moreover, there is evidence of the presence of liquid water and of atmosphere with higher temperature and pressure in the recent past of the Mars—hundreds of thousands and few millions of years ago [101–104]. Our data indicate that microorganisms should survive in the permafrost sediments of such ages, even at shallow depths.

Data on high radioresistance of microorganisms (including our earlier papers [6–9,37]) also indicates that putative contaminants that were carried with past and future space missions can survive on different space objects for a long time. Moreover, as discussed above, there are possibilities for microorganisms’ growth and replication under current Martian conditions. Contaminants from shallow regolith layers might be widely spread on the planet by the dust storms [10], and this can significantly obstruct the search for the indigenous life on the Mars. Due to this, it is necessary to advance methods and protocols for reducing the microbial load on space missions. In particular, new antibacterial surfaces [105] and sterilization methods [106,107] are being developed. Since the results of the study
presented testify in favor of the possibility of extraterrestrial life exists, they should be considered at sample return missions planning meaning appropriate protocols development and implementation in order to prevent Earth contamination.

Irradiation with accelerated electrons led to more damage of microbial communities than irradiation with the same gamma radiation doses under similar conditions [6,7,33,38,39]. This was expressed in a decrease in the total number of prokaryotes as determined by the EFM method, as well as in the sharp suppression of potential metabolic activity. This result is also confirmed with pure cultures: it was previously shown that strains of *Arthrobacter polychromogenes* SN_T61 and *Kocuria rosea* SN_T60 are able to withstand gamma irradiation under low pressure (1 Torr) and low temperature (−50 °C) conditions with doses of at least 10 kGy without a significant reduction number of viable cells [38,39]. Irradiation with accelerated electrons with 10 kGy dose led to a decrease in CFU abundance by 3–4 orders of magnitude. It is necessary to study the effects of the entire spectrum of ionizing radiation, as well as to study the effect of higher radiation doses for a more accurate prediction of the duration of conservation of microorganisms and biomarkers at various objects of the Solar system and beyond.

The modern concept of the Mars evolution assumes a scenario of the planet’s development according to the terrestrial type with the possible emergence of the biosphere [24–28]. The impact of radiation could contribute to the origin of life, being, possibly, a key factor [108]. The discovery of ancient ecosystems in palaearcheal sediments [109–112] suggests that life appeared very quickly after the formation of our planet, and it quickly spread on its surface. However, this was not the Earth familiar to us. The radiation of the young Sun was significantly higher than the current one [113–115]. In particular, X-ray and EUV emission was six and up to 1000 times stronger than those of the present Sun during 3.5 Ga and 4.5 Ga, respectively. It is suggested, that the early Earth’s surface was subjected to the high levels of cosmic ionizing and ultraviolet radiation, although there is uncertainty on the density of the early Earth’s atmosphere [116], which could protect the planet surface against radiation. Moreover, the background from the decay of radioisotopes on the ground was about five times higher [117]. From the standpoint of modern scientific logic, life on such a planet is impossible. But, it arose and quickly evolved, improving the mechanisms of adaptation in a changing environment. The experience of the early evolution of life in the conditions of the young Earth (or outside it, in accordance with the hypothesis of pansperma) should be fixed in the genetic material of descendants of the primary biosphere [118]. This gives grounds for assumptions that the high stability of the currently observed effects of adaptation of biosystems to cosmo-geophysical impacts are atavisms of the geological epoch of the much more active early Sun [119,120]. Similarly, the biosphere could arise and evolved on Mars, developing the processes of adapting life to the very harsh conditions of the planet at the initial stages of planet’s development. At the same time, adaptive-genetic acquisitions of cells in the natural habitat could be accumulated and they could ensure a sustained life-support in a new round of unfavorable conditions with the loss of a significant part of the atmosphere by Mars. For a logically justified estimate of the duration of maintenance of a putative life in a Mars soil, the Earth model is the only object. Therefore, further research is needed on the conditions and limits of life that is embedded of the planetary body, using the example of study natural terrestrial biotopes.

4. Conclusions

We performed the study of viability of natural microbial communities of extreme habitats after irradiation with accelerated electrons (10 kGy, 100 kGy) under conditions of low pressure (~0.01 Torr) and low temperature (~−130 °C), and also the viability of pure bacterial cultures under the same conditions. Radioresistance of a number of bacteria strains under normal conditions has also been evaluated. It is shown that irradiation with electrons with 100 kGy dose in the model conditions of Mars and open space did not sterilize the samples of arid soils and permafrost. The data obtained suggest that viable microorganisms can be viably cryopreserved for at least 1.3 and 20 million years in the Mars regolith in the shallow surface layer and at 5 m depth, respectively; for at least ~170 years in the ice of Europa at 10 cm depth; for at least 400 thousand years in the open space inside the meteorites.
We found that bacteria have a much higher radioresistance as a part of microbial communities in situ in the natural habitat when compared to their stability in pure culture. Due to the protective function of the natural environment and possibly due to the intra- and interpopulation interactions of organisms, even radiosensitive microorganisms being part of natural microbial communities are able to withstand the effects of high doses of ionizing radiation.

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Appendix A

Table A1. The primers used for 16S rRNA gene amplification and sequencing.

| Strain—GenBank Accession Number | Primers Used for the Amplification | Primers Used for the Sequencing |
|---------------------------------|-----------------------------------|---------------------------------|
| KBP.AS.531—MH050938             | 63F+1387R                         | 1100R                           |
| KBP.AS.532—MH050939             | 27F+537R                          | 537R                            |
| KBP.AS.340—MH050940             | 63F+1387R                         | 1100R                           |
| KBP.AS.341—MH050941             | 63F+1387R                         | 1100R                           |
| KBP.AS.342—MH050942             | 63F+1387R                         | 1100R                           |
| KBP.AS.343—MH050943             | 63F+1387R                         | 1100R                           |
| KBP.AS.344—MH050944             | 63F+1387R                         | 1100R                           |
| KBP.AS.345—MH050945             | 63F+1387R                         | 1100R; 537R                     |
| KBP.AS.346—MH050946             | 63F+1387R                         | 1100R                           |
| KBP.AS.347—MH050947             | 63F+1387R                         | 1100R                           |
| KBP.AS.298—MH050948             | 63F+1387R                         | 1100R                           |
| KBP.AS.299—MH050949             | 63F+1387R                         | 1100R                           |
| KBP.AS.300—MH050950             | 63F+1387R                         | 1100R                           |
| KBP.AS.533—MH050951             | 63F+1387R                         | 1100R                           |
| KBP.AS.301—MH050952             | 63F+1387R                         | 1100R                           |
| KBP.AS.302—MH050953             | 63F+1387R                         | 1100R                           |
| KBP.AS.303—MH050954             | 63F+1387R                         | 1100R                           |
| KBP.AS.534—MH050955             | 27F+Un1492R                       | 1100R                           |
| KBP.AS.297—MH050956             | 63F+1387R                         | 1100R                           |
| KBP.AS.532—MH050957             | 63F+1387R                         | 1100R; 537R                     |
| KBP.AS.532—MH050958             | 63F+1387R                         | 1100R                           |
| KBP.AS.323—MH050959             | 63F+1387R                         | 1100R                           |
| KBP.AS.324—MH050960             | 63F+1387R                         | 1100R                           |
| KBP.AS.325—MH050961             | 63F+1387R                         | 1100R                           |
| KBP.AS.535—MH050962             | 63F+1387R                         | 1100R                           |
| KBP.AS.536—MH050963             | 63F+1387R                         | 1100R                           |
| KBP.AS.326—MH050964             | 63F+1387R                         | 1100R; 537R                     |
| KBP.AS.317—MH050965             | 63F+1387R                         | 1100R                           |
| KBP.AS.319—MH050966             | 63F+1387R                         | 1100R                           |
| KBP.AS.320—MH050967             | 63F+1387R                         | 1100R                           |
| KBP.AS.537—MH050968             | 63F+1387R                         | 1100R                           |
Appendix B

Table A2. The bacterial genera cultured from the arid soil samples. The table summarizes some data of the Table 3 and allows to compare bacterial diversity in a simpler view.

| SN2, TPC | SN2, 10 kGy | SN2, 100 kGy |
|----------|-------------|--------------|
| Arthrobacter | Arthrobacter | Arthrobacter |
| Saccharothrix | | |
| Phenyllobacterium | | |
| Cryobacterium | Micrococcus | |
| Streptomyces | Micrococcus | |
| Microbacterium | Spirosoma | Spirosoma |
| Micrococcus | Rufibacter | Rufibacter |
| Sphingouarantiacus | Planomicrobium | |
| Bacillus | Massilia | Microvirga |
| | | Pontibacter |

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