Protein patterns of black fungi under simulated Mars-like conditions

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Two species of microcolonial fungi – *Cryomyces antarcticus* and *Knutia perforans* - and a species of black yeasts – *Exophiala jeaneselmei* - were exposed to thermo-physical Mars-like conditions in the simulation chamber of the German Aerospace Center. In this study the alterations at the protein expression level from various fungi species under Mars-like conditions were analyzed for the first time using 2D gel electrophoresis. Despite of the expectations, the fungi did not express any additional proteins under Mars simulation that could be interpreted as stress induced HSPs. However, up-regulation of some proteins and significant decreasing of protein number were detected within the first 24 hours of the treatment. After 4 and 7 days of the experiment protein spot number was increased again and the protein patterns resemble the protein patterns of biomass from normal conditions. It indicates the recovery of the metabolic activity under Martian environmental conditions after one week of exposure.

Astrobiology is in the focus of many scientific investigations, trying to answer questions about the possible existence of life on other planets, survival in outer space and possible interplanetary transfers. Organisms which inhabit extreme environments – the so-called extremophiles - are generally considered to be the best model for exobiological studies1. Extremophiles can not only survive extreme thermo-physical conditions, but can also tolerate extreme saline, acidic, alkaline or other conditions which can induce DNA damage in less resistant organisms2–5.

Earth’s biosphere has evolved in more than 3 billion years while protected by the planet’s magnetosphere and the atmosphere from the hostile environment of outer space. The conditions of outer space like space vacuum, thermal extremes, cold, solar UV radiation, ionizing radiation and galactic radiation are real challenges for any form of life6. Fungi in general and especially melanized ones showed high resistance when were exposed to ionizing radiation7,8. Large quantities of highly melanized spores have been found in early Cretaceous period deposits when many species of animals and plants become extinct. This period matches with Earth’s crossing the ‘magnetic zero event’ when the planet lost its protection against cosmic radiation9.

Martian climate conditions are too cold and the atmosphere too thin for allowing bulk water to be stable. However, at environmental conditions similar to Mars-like conditions the metabolic activity has been detected in presence of one or two monolayers of water, which was proved by laboratory experiments10. A precondition for habitability on Mars would be the water availability and interaction between regolith and near-surface atmosphere11. Some investigations were performed on organisms which might have a certain potential to live on Mars1,4,12,13. Although organisms need water activities of \(a_w = 0.8–0.9\) for active metabolism, there are some special life forms which are also able to live in much drier conditions with a temporary saturation of atmospheric water vapor, for instance black microcolonial fungi (MCF) and some lichens. Such temporary saturation can occur on Mars1,4,13 as it is known by recent studies some extremophilic organisms are able to cope successfully with Mars-like conditions. For instance, it was reported that the vitality and photosynthetic activity of lichens is maintained under Mars-like conditions1,4,14.

Melanized fungi and lichens are colonizing extreme environments such as Arctic and Antarctic regions, including high altitude terrains. These specific regions are characterized by a combination of dry, cold, oligotrophic extremes along with huge fluxes of UV radiation16,17. Therefore microcolonial fungi - and particularly *C. antarcticus* – have been suggested as eukaryotic models for studies on the habitability of Mars14 and the biological exploration of Mars1,3,18. Both, Mars-simulation experiments and the exposition of the fungus to outer space have shown that *C. antarcticus* is able to survive under Mars and space conditions in a good manner1,19. Also the green alga *Stichococcus* sp. and the lichenized fungus *Acarospora* sp. were shown to be able to survive Mars and space conditions20.
Until now the cellular mechanisms underlying the resistance of MCF against harsh space and Mars conditions are not yet understood. At the moment there is limited information existing on a genomic sequences of MCF due to some obstacles occurring during DNA extraction and purification. Proteomics approach leads to understanding of the expression and regulation of the entire set of proteins which are directly related to biological functions. Therefore two dimensional gel electrophoresis (2D) for protein profiling is carried out in this study in order to get a more precise impression of cellular and metabolic activity of these fungi under simulated Mars-like conditions. Three model organisms were chosen for the experiments, that were carried out in the Mars-simulation chamber in the Mars Simulation Facility (MSF) of the German Aerospace Centre (DLR, Berlin): (1) Cryomyces antarcticus represents a group of extremophilic fungi occurring in Antarctic dry deserts; (2) Knufia perforans is a mesophilic but highly stress tolerant fungus found in hot and arid environments and (3) Exophiala jeanselmei represents a black yeast closely related to opportunistic pathogens in humans. A realistic simulation of the Martian climate was based on data provided by Mars-related missions.

Results
The results of the experiments showed (Table 1) that the number of expressed proteins has changed during the experiment in all fungi tested. After exposure to Mars-like conditions for 24 hours (day simulation) E. jeanselmei reduced the number of protein spots from 473 to 237 (Fig. 2). Protein spots representing proteins with high molecular weight (<120 kDa) and basic pI-ranges of <7 were reduced in the number. From 24 hours to 4 days the number of proteins increased to 359 spots by expressing large proteins with basic pH. After incubation for 7 days the number of protein spots reached nearly the initial spot number of 470. Within the protein profiles no significant changes of the patterns were observed; nearly all the spots were matched among the gels of different conditions, thus indicating there was no expression of new proteins after 24 hours (day simulation) or 4/7 days exposure. After 7 days the pattern and number of proteins resembles the cellular state at the starting point (the biomass from normal conditions) of the experiment, which indicated that no novel or stress-related proteins were expressed.

Also in K. perforans the number of proteins decreased after 24 hours before increasing during the following days of the experiment: after exposure for 24 hours (day simulation) the number of protein spots decreased significantly from 634 to 264 spots. The reduced proteins were of a large size (with the molecular weight of above 120 kDa and basic pI-ranges of 7 to 9). After 4 days of treatment 286 protein spots were detected and after 7 days the protein number increased to 328. The induction of new proteins was not detected in this fungus; however, many proteins were down-regulated and could not be visualized by silver-staining (Fig. 3). Protein extraction of E. jeanselmei and K. perforans for samples from night simulation was not successful and therefore the following 2D analysis could not be performed.

In C. antarcticus 406 spots were detected in the biomass from normal conditions before the start of the Mars-simulation treatment. After incubation for 24 hours at day and night simulation only 93 and 97 spots respectively were detected. There were no new protein spots appeared on the 2D gel patterns but over-expression of several proteins was observed (Fig. 4). Unfortunately, protein concentration of this fungus after 4 and 7 days of exposure to simulated Mars-like conditions was too low to perform further proteomic analyses.

After re-inoculation and incubation for 2 weeks at room temperature the increase of biomass (what is indicating growth of all the tested organisms) was observed. The biomass growth of all the tested strains manifested reproduction ability and survival for all fungal species after Mars-simulated treatments. The biomass, which used for the experiment was limited by the size of experimental chamber (Fig. 5). As it was mentioned in Materials and Methods section the extraction protocol was adapted to get sufficient protein amounts from those precious biomass samples. The analytical runs by 2DE was accomplished successfully for almost all the samples, however the amounts of the protein rather low to allow subsequent protein identification by proteomic approaches.

Discussion
The investigations of the microorganisms exposed to outer space and planetary simulated conditions support and enhance our understanding of basic biological mechanisms, such as biological effects caused by the radiation field in space and survival factors in the upper boundary of Earth’s biosphere. Moreover the likelihood of interplanetary transport of microorganisms via meteorites, the use of microorganisms in bioregenerative life support systems, the monitoring, characterization and control of spacecraft microflora and associated microbial crew health are under concern. Another aspect of space biological research is the investigation of the responses of microorganisms to simulated Mars-like conditions and their evaluation as potential forward contamination risks in the context of planetary protection.

The fungal abilities to withstand harsh outer space conditions and tendency to contaminate spacecrafts are of high importance due to: (1) some of the fungi are potential human pathogens and this could possess certain danger for the well-being of the crew in long-distant space journeys and (2) fungi have strong enzymatic systems and secrete various metabolites which can cause degradation of structural materials. In their natural environment black microcolonial fungi from extreme environments are exposed to periods of temporary saturation with water vapor and temporary dryness. Although black microcolonial fungi, some lichens and cyanobacteria were proven to survive Martian and outer space conditions, it was never investigated on molecular level which cellular processes are responsible for this remarkable resistance.

The results of this study are the first approach to obtain a better understanding of survival mechanisms of MCF under Mars-like conditions. Although the experiments did not allow us to extract enough protein amounts for the identification of single protein spots (due to technical reasons allowing only very low amounts of the biomass to be placed inside the Mars simulation chamber), the results give some astonishing insights into ecology of MCF. The most important results of the study are:

The investigated fungi strains showed metabolic activity in close to anaerobic conditions during exposure to Mars-like conditions with a final exposure time of about 4 days in the first experiment and 7 days

| Table 1 | Number of protein spots detected in the 2D gels of the analyzed samples at each exposure condition. Opt-optimal laboratory conditions (normal conditions) at which fungi were grown for 30 days at temperature 20 °C for E. jeanselmei and K. perforans and 15 °C and for C. antarcticus. 24 hours (D/N) - samples were exposed to Mars-simulated conditions for 24 hours with day/night simulation respectively. 4/7 days- samples were exposed to Mars-simulated conditions for 4 and 7 days respectively. Blank cells (---) indicate that the protein concentration was too low for 2DE analysis. |
| Strain | Number of spots at each exposure condition | 24h-D | 24h-N | 4days | 7days |
| E. jeanselmei | 473 | 237 | - | 359 | 470 |
| K. perforans | 624 | 264 | - | 286 | 328 |
| C. antarcticus | 406 | 93 | 97 | - | - |

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in the second experiment. Although there was a significant decrease of the number of protein spots during the first 24 hours, the fungi seem to recover and gain the initial metabolic state after 7 days of the exposure. These results are in accordance with what was observed earlier by Zakharova and colleagues\textsuperscript{26}, when fungi were desiccated to the constant weight and a decrease of proteins was observed in \textit{C. antarcticus} whereas \textit{K. perforans} and \textit{E. jeaneselmei} reacted by expression of new proteins, possibly representing protective proteins. During the following increase of relative humidity fungi did never gain a fully rehydrated state but they gained the original protein pattern after one hour of rehydration. The results of the Mars simulation experiments confirm the hypothesis that the fungi are able to be metabolic active with minimum amount of water and that their proteins have special conformations working with minimum water content\textsuperscript{22,26}.

The fungi do not show the signs of stress reaction during the experiment. Previous studies have also shown that \textit{C. antarcticus} does not react to temperature stress or by desiccation through production of novel proteins\textsuperscript{26,27}. However, for \textit{E. jeaneselmei} and for \textit{K. perforans} it was shown that they express proteins which could be protein chaperons or the so-called heat shock proteins\textsuperscript{26,27}. In case of the Mars simulation none of the fungi showed such signals of stress-response although they seem to be active when exposed to Mars-simulated conditions.

The results of our study suggest that MCF seem to be not significantly stressed by Mars-like conditions, which are a combination of

![Temperature and relative humidity profile](image-url)
temperature extremes, pressure, dryness and radiation. The reason for that could be that the fungi have a basic set of stress resistant proteins and high doses of radiation enhance their growth capacity instead of damaging their cellular components. It was reported that the environments with high radiation, resulting from human activities, such as damaged reactor at Chernobyl and reactor cooling pool-water are habitats for black yeasts and MCF and that “melanizes fungal cells manifested increased growth after exposure to ionizing radiation”. Analyzing melanized fungal species which were found in Chernobyl reactor and in the reactor cooling pool water showed that high fluxes of radiation selected for highly radio-resistant types of microorganisms, showed increased catalase and nuclease activities.

The results achieved from our study led to the conclusion that black microcolonial fungi can survive in Mars environment. Further investigations are needed to characterize the present proteins and their functions. In other distantly related fungi strains some proteins are discovered, which might play a key role in stress-resistance to different environmental stresses. For instance the results obtained by Gocheva and colleagues demonstrate that the growth at low temperatures does clearly induce oxidative stress events. Antarctic strains (Penicillium spp.) demonstrated a marked rise in activities of protective enzymes such as superoxide dismutase and catalase at decreasing temperatures. Low-temperature resistance is partially associated with enhanced scavenging systems. Also it is known that protein carbonyls are biomarkers of protein oxidation followed by temperature stress. The protein AoSO prevents excessive cytoplasmic leakage upon hyphal injury by accumulating at the septal pore, moreover in response to various stresses (low and high temperature; extreme acidic and alkaline pH; nitrogen and carbon deplation). The protein Ip2431 was discovered being important under several stress-induced proteins in fungus Cryptococcus neoformans. Heterotrimeric Gα protein Pga1 plays a central role in the regulation of the whole growth-developmental program of Penicillium chrysogenum. This protein also plays important role in germination (mediating carbon source sensing); absence of Pga1 increases resistance to thermal, oxidative and osmotic stress. According to Deegenaars and colleagues, a 110 kDa proteins play a role in stress tolerance in psychrophilic yeast, similar to that of HSP 104 in mesophilic species. A described transcription factor PMSn2 in Beauveria bassiana and Metarhizium robertsi is important in tolerance to hyperosmolarity, oxidation, carbendazim, cell wall perturbing, high temperature and UV-B radiation. The results of our study, however, suggest that stress-related protein machinery of MCF is more complex compared to other fungi strains and there are a number of highly abundant proteins needed for the survival in Mars-simulated conditions.

We can conclude that an unknown metabolic pathway might be discovered, which enables the fungi to live in a quasi-anaerobic Mars-like environment.

Methods

The biological samples as model organisms. The organisms used for this study were: (1) Exophiala jeanselmei MA 2853, a rock inhabiting black yeast closely related to opportunistic pathogens in humans; (2) Knufia perforans MA 1299, a mesophilic but highly stress tolerant fungus found in hot and dry environments, like the Mediterranean; (3) Cryomyces antarcticus MA 5682, an extremophilic fungus from Antarctica. The strains were obtained from the ACBR culture collection (Austrian Center of Biological Resources and Applied Mycology, www.acbr-database.at).

Inoculi were prepared as cell suspensions and drop-inoculated onto sterilized cellophane membranes (Model 583 gel dryer Baking. Catalog #1650963, Bio-Rad), which were placed on the surface of 2% malt-extract agar (MEA, Applichem GmbH,
Darmstadt, Germany). Fungi were grown at normal conditions: 20 °C (293 K) for *E. jeanselmei* and *K. perforans* and 15 °C (288 K) and for *C. antarcticus* for 30 days. For samples from normal conditions the biomass was harvested by scratching the material from the plates using a scalpel, transferred into a sterile tube, then immediately frozen in liquid nitrogen and stored at −80 °C (193 K) for further analysis. Other samples were prepared as follows: round segments approximately three cm diameter from the cellophane membranes with well grown fungal colonies were cut out and transferred into empty three cm diameter dishes and then placed into the Mars simulation chamber.

**Mars-simulation.** The experiment was carried out in the Mars Simulation Facility (MSF) at the Institute of Planetary Research of the German Aerospace Center (DLR) Berlin. The MSF simulates Mars-like atmospheric conditions and is used to perform laboratory experiments with controlled time-profiles. The main part of the MSF is an “experimental chamber” (EC) located in a temperature test chamber. The experiments were performed in EC, which can be cooled, evacuated and filled with defined gas mixture and humidity in a controlled manner. The MSF, its performance and typical experiments are described by Lorek and Koncz.

For the experiment, two gases (CO₂ and air) were mixed and humidified. A gas volume flow of 10 liters per hour (approximately Standard Ambient Temperature 25 °C (298 K) and Pressure 101325 Pa) of the resulting gas mixture (CO₂ 95%/volume, air 5%/volume (N₂ 4%/volume and O₂ 1%/volume) was used to pass through the EC. Thus the humidity of the mixture inside the chamber, which was directly provided, corresponded to a partial water vapor pressure of about 3 Pa, which is the average closed to the 2.7 Pa vapor pressure in situ measured in polar regions on Mars. The pressure inside the experiment chamber was between 1000 Pa (at the starting time) and 700 Pa (main part until the end of the experiment). Complete Xenon-lamp radiation spectra between 200 nm and 2200 nm was used. The Xenon-lamp was switched on and off every day to stimulate the diurnal cycle of the sun. The day simulation was made by allowing light for 16 hours (for 8 hours light was off); night simulation for testing the effect of Mars-like atmospheric conditions without radiation was produced by covering the samples with aluminium foil. An applied

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**Figure 3 | 2D gel patterns obtained after exposure of Knufia perforans:** (A): sample from optimal conditions. (B): sample was exposed to Mars-simulated conditions for 24 hours with day simulation. (C): sample was exposed to Mars-simulated conditions for 4 days with day and night simulation. (D): sample was exposed to Mars-simulated conditions for 7 days with day and night simulation.

**Figure 4 | 2D gel patterns obtained after exposure of Cryomyces antarcticus.** (A): sample from optimal conditions. (B): sample was exposed to Mars-simulated conditions for 24 hours with day simulation. (C): sample was exposed to Mars-simulated conditions for 24 hours with night simulation.
Figure 5 | Experimental flow. (A): Sample preparation for loading into experimental chamber. (B): samples prepared for the exposure with day and night (the lid of the plate is covered with aluminium foil, enabling prevention of UV radiation exposure). (C): gas-mixing system. (D, E): samples after exposure to Martian simulated conditions. (F): Mini-Pam (photosynthetic activity), Xenon-Lamp housing and measurement equipment.

Table 2 | Mars-simulated conditions used for the experiments

| Parameter                           | Values                                                                 |
|-------------------------------------|------------------------------------------------------------------------|
| Gas mixing rates                    | CO₂ 95%/volume, air 5%/volume  
|                                     | [N₂ 4%/volume and O₂ 1%/volume]                                       |
| Gas humidity                        | 3 Pa                                                                   |
| Volume flow                         | 10 liters/hour*                                                        |
| Pressure inside the chamber          | 1000**–700*** Pa                                                       |
| Radiation spectra                   | 200 nm and 2200 nm                                                     |
| Simulation                           | day simulation for 24 hours: light on for 16 hours and light off for 8 hours; night simulation for 24 hours: light off |
| Temperature                         | was varying in a diurnal cycle between:  
|                                     | day simulation: 15 °C (288 K)  
|                                     | night simulation: −55 °C (218 K)                                       |
| Exposure time with corresponding dose| 24 hours(day and night simulation): 153 kJ/m²;  
|                                     | 4 days simulation: 598.8 kJ/m²;  
|                                     | 7 days simulation 1131 kJ/m²                                           |

*At approximately Standard Ambient Temperature and Pressure 101325 Pa and 25 °C.  
**At the starting time and ***main part until the end of the experiment.

0.9%) and proceeded as given in the basic protocol. After cell disruption the lysate was transferred into 15 ml polypropylene centrifuge tube and 3 ml of Tris-buffered phenol solution, pH 8.0 (Sigma–Aldrich, Steinheim, Germany) was added and mixed for 40 min at room temperature. The remaining steps were performed according to the basic protocol. Protein determination was carried out using Bradford Protein Assay.

The Bradford protein Assay was performed to determine the concentration of protein in fungal extracts. Reactions were carried out in microtiter plates according to the manufacturer instructions. A standard curve was established using serial dilutions from 0.8 µg ml⁻¹ to 100 µg ml⁻¹ of bovine serum albumin (BSA). The resulting optical density (OD) at 595 nm was analyzed with a plate reader (Magellan; Tecnac Austria, Graz, Austria). All experiments were carried out in triplicate. For each gel 20 µg of proteins were applied. IEF separation was performed using 13 cm strips pH 3–10NL. For each condition 2D gels were made in triplicate. Visualization of protein spots was made by a high sensitive mass spectrometry compatible silver staining. The gels were fixed in 50% (v/v) methanol and 5% (v/v) acetic acid for 20 min, followed by washing in 50% (v/v) methanol for 10 min and then left in MilliQ water (Millipore, MA, USA) overnight at 4 °C (277 K) with gentle agitation. Thereafter, the 2D gels were sensitized using a 0.02% (w/v) sodium thiosulphate solution for 1 min followed by incubation in 0.04% (v/v) formalin and 2% (w/v) sodium carbonate solution until the desirable intensity of staining was achieved. All the washing solutions were prepared in MilliQ water (Millipore, MA, USA).

After staining the gels were scanned in TIFF 16 bit format; for each sample (normal condition, 24 hours(day/night simulation) and exposure for 47 days); the 3 gels were matched by warping (Image Master 2D Platinum version 5.0, Amersham Biosciences, Swiss Institute of Bioinformatics, Geneva, Switzerland) and the sum of all spots which were present at least in two gels was taken into account for the protein pattern analysis.

1. Onofri, S. et al. Resistance of Antarctic black fungi and cryptendolithic communities to simulated space and Mars conditions. Stud. Mycol. 61, 99–109 (2008).
2. Billi, D. Subcellular integrities in Chroococcidiopsis sp. CCME029 survivors after prolonged desiccation revealed by molecular probes and genome stability assays. Extremophiles. 13, 49–57 (2009).
3. Cavicchioli, R. Extremophiles and the search for extraterrestrial life. Astrobiology. 2, 281–292 (2002).
4. de Vera, J.-P. et al. Survival potential and photosynthetic activity of lichens under Mars-like conditions: a laboratory study. Astrobiology. 10, 215–227 (2010).
5. de Vera, J.-P. Lichens as survivors in space and on Mars. Fungal Ecol. 5, 472–479 (2012).
6. Hornbeck, G., Klaus, D. M. & Mancinelli, R. L. Space Microbiology. Microbiol. Mol. Biol. Rev. 74, 121–156 (2010).
7. Dighton, J., Tugay, T. & Zhdanova, N. Fungi and ionizing radiation from radionuclides. EEMS Microbiol. Lett. 281, 109–120 (2008).
8. Dadachova, E. & Casadevall, A. Ionizing radiation: how fungi cope, adapt, and exploit with the help of melanin. Curr. Opin. Microbiol. 11, 525–531 (2008).
9. Hulot, G. & Gallet, Y. Do superchrons occur without any palaeomagnetic warning? Earth and Planetary Science Letters. 210, 191–201 (2003).
10. Rivkina, E. M., Friedmann, E. I., McKay, C. P. & Gilichinsky, D. A. Metabolic activity of permafrost bacteria below the freezing point. Appl. Environ. Microbiol. 66, 3230–3233 (2000).
26. Zakharova, K. et al. 19. Onofri, S.
25. Sto¨ffler, D. et al
28. Mironenko, N. V., Alekhina, I. A., Zhdanova, N. N. & Bulat, S. A. Intraspecific
27. Tesei, D. et al
23. Novikova, N. D. Review of the knowledge of microbial contamination of the
15. Mo¨hlmann, D. Adsorption water-related potential chemical and biological
et al
13. McKay, C. P. Icarus.
11. Stoker, C. R. Fungal Genet. Biol.
10. Scalzi, G. Icarus.
9. Onofri, S. et al Survival of rock-colonizing organisms after 1.5 years in outer
8. Moeller, R., Schuerger, A. C., Reitz, G. & Nicholson, W. L. Protective role of spore
7. McKay, C. P. Biochem.
6. Robinson, C. H. Cold adaptation in Arctic and Antarctic fungi. New Phytologist.
5. Möhlmann, D. Adsorption water-related potential chemical and biological
4. de Vera, J.-P. et al Adaptation of an Antarctic lichen to Martian niche conditions can
3. McKay, C. P. Microcolonial fungi on rocks: a life in constant drought?
2. Moeller, R., Schuerger, A. C., Reitz, G. & Nicholson, W. L. Protective role of spore
1. Stoker, C. R. et al. Habitability of the Phoenix landing site. J. Geophys. Res. 115: E09E20, 24 pp. DOI: 10.1029/2008JE003421 (2010).
12. Morozova, D., Möhlmann, D. & Wagner, D. Survival of methanogenic archaea from Siberian permafrost under simulated Martian thermal conditions. Orig. Life Ecol. Biosph. 37, 189–200 (2007).
14. de Vera, J.-P. et al. Adaptation of an Antarctic lichen to Martian niche conditions can occur within 34 days. PSS. (in press) (2013).
15. Möhlmann, D. Adsorption water-related potential chemical and biological processes in the upper Martian surface. Astrobiology. 5, 770–777 (2005).
16. Robinson, C. H. Cold adaptation in Arctic and Antarctic fungi. New Phytologist. 151, 341–353 (2001).
17. Sterflinger, K., de Hoog, G. S. & Haase, G. Phylogeny and ecology of meristematic ascomycetes. Stud. Mycol. 43, 5–22 (1999).
18. Onofri, S., Selbmann, L. & Pagano, S. Antarctic microfungi as models for exobiology. Planetary Space Sci. 52, 229–237 (2004).
19. Onofri, S. et al Survival of rock-colonizing organisms after 1.5 years in outer space. Astrobiology. 12, 508–516 (2012).
20. Scalzi, G. et al. LIFE Experiment: isolation of Cryptoendolithic organisms from Antarctic colonized sandstone exposed to space and simulated Mars conditions on the international space station. Orig. Life Evol. Biosph. 42, 253–262 (2012).
21. Marzbazan, G., Tesei, D. & Sterflinger, K. A review beyond borders: proteomics of microcolonial black fungi and black yeast. Nat. Sci. 5, 640–645 (2013).
22. Moeller, R., Schuerger, A. C., Reitz, G. & Nicholson, W. L. Protective role of spore structural components in determining Bacillus sublitis spore resistance to simulated Mars surface conditions. Appl. Environ. Microbiol. 87, 8849–8853 (2012).
23. Novikova, N. D. Review of the knowledge of microbial contamination of the Russian manned spacecraft. Microb. Ecol. 47, 127–132 (2004).
24. de la Torre, R. et al. Likelihood of interplanetary transfer of rock-inhabiting microbial communities: results from the space experiment Lithopanspermia. Icarus. 208, 735–748 (2010).
25. Stoffler, D. et al. Experimental evidence for the potential impact emission of viable microorganisms from Mars and Mars-like planets. Icarus. 186, 585–588 (2007).
26. Zakharova, K. et al. Microcolonial fungi on rocks: a life in constant drought? Mycopathologia 175, 537–547 (2013).
27. Tesei, D. et al. Alteration of protein patterns in black rock inhabiting fungi as a response to different temperatures. Fungal Biol. 116, 932–940 (2012).
28. Mironenko, N. V., Alekhina, I. A., Zhdanova, N. N. & Bulat, S. A. Intraspacific variation in Gamma-radiation resistance and genomic structure in the filamentous fungus Alternaria alternate: a case study of strains inhabiting Chernobyl reactor No.4. Ecotoxicol. Environ. Safety. 45, 177–187 (2000).
29. Robertson, K. L. et al. Adaptation of the Black yeast Wangiella dermatitidis to ionizing Radiation: Molecular and Cellular Mechanisms. PLoS ONE 7 (2012).
30. Gocheva, Y. G. et al. Cell response of Antarctic and temperate strains of Penicillium spp. to different growth temperature. Mycol. Res. 110, 1347–1354 (2006).
31. Li, Q., Harvey, L. M. & McNeil, B. The effects of elevated process temperature on the protein carbonyls in the filamentous fungus, Aspergillus niger B1–D. Process Biochem. 43, 877–881 (2008).
32. Maruyama, J., Escaño, C. S. & Kitamoto, K. AoSO protein accumulates at the septal pore in response to various stresses in the filamentous fungus Aspergillus oryzae. Biochem. Biophys. Res. Commun. 391, 868–873 (2010).
33. Kroll, K., Pahtz, V. & Kniemeyer, O. Elucidating the fungal stress response by proteomics. J. proteomics (in press) (2013).
34. Garcia Rico, R. O., Martin, J. F. & Fuero, F. Heterotrimetric Gx protein Pg1 from Penicillium chrysogenum triggers germination in response to carbon source and affects negatively resistance to different stress conditions. Fungal Genet. Biol. 48, 641–649 (2011).
35. Deegenaars, M. L. & Watson, K. Stress proteins and stress tolerance in an Antarctic, psychrophilic yeast, Candida psychrophila. FEMS Microbiol. Lett. 151, 191–196 (1997).
36. Liu, Q., Ying, S.-H., Li, J.-G., Tian, C. G. & Feng, M.-G. Insight into the transcriptional regulation of Msn2 required for condiation, multi-stress responses and virulence of two entomopathogenic fungi. Fungal Genet. Biol. 54, 42–51 (2013).
37. Lorek, A. & Koncz, A. [Simulation and measurement of extraterrestrial conditions for experiments on habitability with respect to mars]. Habitability of Other Planets and Satellites [de Vera, J.-P. & Seckbach, J. (ed.)] [145–162] (Springer Netherlands, Dordrecht, 2013).
38. Zent, A. P. et al. Initial results from the thermal and electrical conductivity probe (TECP) on Phoenix. J. Geophys. Res. 115: DOI: 10.1029/2009JE003420 (2010).
39. Isola, D. et al. Establishment of sample preparation and 2-DE procedure for protein expression profiling of black fungi. Fungal Biol. 115, 971–977 (2011).
40. Bradford, M. M. A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein–dye binding. Anal. Biochem. 72, 248–254 (1976).
41. Shevchenko, A., Wilm, M., Vorm, O. & Mann, M. Mass spectrometric sequencing of proteins from silver stained polyacrylamide gels. Anal. Chem. 68, 850–858 (1996).

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Author contributions
K.S. conceived the project, she is the project-leader of the FWF project grant; she established the collaboration with J.-P.dV. K.S., J.-P.dV. and K.Z. made an experimental design. J.-P.dV. provided access to the facility of German Aerospace Center (DLR); J.-P.dV. and A.L. supervised the laboratory work at DLR; provided the data for ‘Mars simulation’ subunit in ‘Materials and Methods’ chapter; proof-read the manuscript. K.Z. performed the laboratory work (exposure of the samples to Mars-simulated conditions) at DLR; laboratory work and computer analysis (proteomics) at University of Natural Resources and Life Sciences, Vienna (BOKU); wrote the manuscript. G.M. and K.S. supervised the practical work in the laboratory of BOKU; proof-read the manuscript.

Additional information
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