Exploiting a perchlorate-tolerant desert cyanobacterium to support bacterial growth for \textit{in situ} resource utilization on Mars

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\section*{Abstract}

The presence of perchlorate in the Martian soil may limit \textit{in-situ} resource utilization (ISRU) technologies to support human outposts. In order to exploit the desiccation, radiation-tolerant cyanobacterium \textit{Chroococcidiopsis} in Biological Life Support Systems based on ISRU, we investigated the perchlorate tolerance of \textit{Chroococcidiopsis} sp. CCME 029 and its derivative CCME 029 P-MRS. This strain was obtained from dried cells mixed with Martian regolith simulant and exposed to Mars-like conditions during the BIOMEX space experiment. After a 55-day exposure of up to 200 mM perchlorate ions, a tolerance threshold value of 100 mM perchlorate ions was identified for both \textit{Chroococcidiopsis} strains. After 40-day incubation, a Mars-relevant perchlorate concentration of 2.4 mM perchlorate ions, provided as a 60 and 40\% mixture of Mg- and Ca-perchlorate, had no negative effect on the growth rate of the two strains. A proof-of-concept experiment was conducted using \textit{Chroococcidiopsis} lysate in ISRU technologies to feed a heterotrophic bacterium, i.e. an \textit{Escherichia coli} strain capable of metabolizing sucrose. The sucrose content was fivefold increased in \textit{Chroococcidiopsis} cells through air-drying and the yielded lysate successfully supported the bacterial growth. This suggested that \textit{Chroococcidiopsis} is a suitable candidate for ISRU technologies to support heterotrophic BLSS components in a Mars-relevant perchlorate environment that would prove challenging to many other cyanobacteria, allowing a ‘live off the land’ approach on Mars.

\section*{Introduction}

Sustaining human outposts on Mars by providing life-support consumables from Earth is unrealistic mainly due to launch costs, travel times and failure risks. The development of Life Support Systems (LSS) is mandatory. On Earth, oxygenic photosynthesizers convert CO\textsubscript{2} to organic matter which serves as food, and H\textsubscript{2}O to O\textsubscript{2}, processes that should play a key role in Biological Life Support Systems (BLSS) off the planet (Binot \textit{et al.}, 1994; Wheeler \textit{et al.}, 2008). An important aspect of LSS is the utilization of local materials, the so-called \textit{in-situ} resource utilization (ISRU), although, to date, these technologies focused mainly on inorganic chemistry and building materials (Sridhar \textit{et al.}, 2000).

A link between local resources and BLSS components, otherwise unable to live ‘off the land’ might be provided by oxygenic phototrophs, such as cyanobacteria, thanks to the development of Bio-ISRU technologies (Verseux \textit{et al.}, 2016; Brown, 2018). Lithotrophic cyanobacteria have been proposed as ideal candidates for Bio-ISRU due to their capability of utilising rocks as a growth substrate (Olsson-Francis and Cockell, 2010).

The concept of using cyanobacteria to feed bacteria has been approached by genetically modifying the cyanobacterium \textit{Anabaena} sp. 7120 in order to secrete sucrose to grow \textit{Bacillus subtilis} (Moss and Ho, 2011; McCutcheon \textit{et al.}, 2015). This approach was further developed into the NASA PowerCell space experiment, on-board the DLR EuCROPIS satellite (Hauslage \textit{et al.}, 2018), aimed to germinate \textit{Bacillus subtilis} spores by using \textit{Anabaena} lysate under simulated Moon and Martian gravity (McCutcheon \textit{et al.}, 2015).

It has been suggested that cyanobacteria could be used in BLSS using Martian local resources with relatively low hardware requirements and attenuation of surface conditions, such as low pressure and high flux of ionizing and UV radiation (Lehto \textit{et al.}, 2006; Murukesan \textit{et al.}, 2016). However, the presence in the Martian soil of a highly oxidizing agent such as perchlorate in the range of 0.4–0.6 wt \% (Hecht \textit{et al.}, 2009) provides an additional limiting factor for Bio-ISRU.

The exploitation of desert strains of \textit{Chroococcidiopsis} to support human space exploration has been proposed taking advantage of their remarkable desiccation- and radiation-tolerance (Billi, 2019). In particular, strain CCME 029 isolated from the Negev Desert has been
extensively investigated and reported to survive years of desiccation (Billi, 2009), as well as high doses of ionizing (Billi et al., 2000; Verseux et al., 2017) and UV radiation (Baqué et al., 2013). In addition, strain CCMEE 029 was reported to cope with Mars-like conditions simulated in low Earth orbit, when exposed as dried cells mixed Martian regolith simulants (Billi et al., 2019a) or as dried biofilms (Billi et al., 2019b).

Although the strategies to withstand the above mentioned conditions have not been fully deciphered, for 

\[ \text{Chroococcidiopsis} \]
sp. CCMEE 029 the following mechanisms have been identified: (i) avoidance of protein oxidation during air-drying and ionizing radiation (Fagliarone et al., 2017); (ii) accumulation in response to desiccation of sucrose and trehalose that stabilize dried subcellular components, including ribosomal RNA and mRNAs codifying DNA repair proteins (Mosca et al., 2019; Fagliarone et al., 2020); and (iii) over-expression of DNA repair genes upon rehydration of dried cells exposed to a Mars-like UV flux (Mosca et al., 2019).

In the present work, we sought to address the hypothesis that as a consequence of its efficient antioxidant system, 

\[ \text{Chroococcidiopsis} \]
sp. CCMEE 029 might be perchlorate resistant and therefore a good candidate for Bio-ISRU on Mars. To test this hypothesis, we selected strain CCMEE 029 and its derive CCME 029 P-MRS obtained from dried cells mixed with Phyllisillicatic Martian Regolith Simulant (P-MRS) and exposed to Mars simulations during the BIOMEX (Biology and Mars EXperiment) space experiment performed during the EXPOSE-R2 space mission (Billi et al., 2019a). The perchlorate tolerance threshold of the two strains was identified by monitoring the growth rate during a 55-day exposure to 5, 50 and 100 mM Na-, Mg- and Ca-perchlorate. Once their perchlorate tolerance threshold was identified, the two strains were grown for 40 days in the presence of Mars-relevant perchlorate concentration, for example, 2.4 mM perchlorate ions supplied as a 60 and 40% mixture of Mg- and Ca-perchlorate, respectively, as reported by the NASA's Phoenix Mars Lander (Hassler et al., 2014). Then the biomass yielded by strain CCME 029 grown in 2.4 mM perchlorate ions was used to produce the lysate used to feed an \( \text{E. coli} \) strain capable of metabolizing sucrose.

**Material and methods**

**Organisms and culture conditions**

\[ \text{Chroococcidiopsis} \]
sp. CCMEE 029 was isolated by Roseli Ocampo-Friedmann from cryptoendolithic growth in the Negev Desert (Israel). The strain is part of the Culture Collection of Microorganisms from Extreme Environments (CCMEE), established by E. Imre and Roseli Ocampo-Friedmann, that is currently maintained at the Department of Biology, University of Rome Tor Vergata. \[ \text{Chroococcidiopsis} \]
sp. CCMEE 029 P-MRS was derived from dried cells of CCMEE 029 mixed with P-MRS and exposed to Mars simulations in low Earth orbit inside the EXPOSE-R2 facility; \( \text{CO}_2 \) atmosphere (780 Pa), \( 2.19 \times 10^2 \text{kJ m}^{-2} \) of \( \text{UV}_{200-400\text{nm}} \) radiation and 0.5 Gy of ionizing radiation (Billi et al., 2019a). Cyanobacterial strains were grown in BG11 medium (Rippka et al., 1979) by using 50 ml vented flasks, inside an incubator at 25°C, without shaking, under a photon flux density of 40 \( \mu \text{mol m}^{-2}\text{s}^{-1} \) provided by cool-white fluorescent lamp (4100 K) under continuous illumination.

\[ \text{Escherichia coli} \] strain W (ATCC 9637) was purchased from the American Type Culture Collection (Manassas, VA, USA) and grown in Luria-Bertani broth (Sambrook et al., 1989) at 37°C with orbital shaking.

**Monitoring of growth rates**

Cyanobacterial growth was monitored by determining the optical density at 730 nm (OD\text{730}) of triplicates, each one consisting of 100 \( \mu \)l aliquot. Bacterial growth was determined by measuring the optical density at 600 nm (OD\text{600}) of triplicates, each one consisting of 100 \( \mu \)l aliquot. Two calibration curves, one for OD\text{730} versus cyanobacterial concentration and one for OD\text{600} versus bacterial concentration, were obtained by determining cell concentrations (cells ml\textsuperscript{-1}) with a Burker’s chamber.

**Determination of cyanobacterial perchlorate tolerance**

One-ml aliquots of the two cyanobacterial cultures in the early stationary phase (about \( 1 \times 10^7 \) cells) were centrifuged and pellets inoculated into 50 ml vented flasks containing 10 ml of liquid BG-11 with 5, 50 and 100 mM Na-, Mg- and Ca-perchlorate. Controls were obtained by inoculating 1 ml aliquots in liquid BG-11. During the 55 days of growth, the culture volume was kept constant by adding BG-11 or BG-11 with 5, 50 and 100 mM Na-, Mg- and Ca-perchlorate. The experiment was performed in triplicate.

**Cyanobacterial growth in Mars-relevant perchlorate**

One-ml aliquots of the two cyanobacterial cultures in the early stationary phase (about \( 1 \times 10^7 \) cells) were centrifuged, and the pellets inoculated into 50 ml vented flasks containing 10 ml of BG-11 with 2.4 mM Cl\text{O}_4\text{\textsuperscript{-}} provided as a 40% Mg-perchlorate and 60% Ca-perchlorate mixture. Each experiment was performed in triplicate. During the 40 days of growth, the culture volume was maintained constant by adding BG-11 or BG-11 with 2.4 mM Cl\text{O}_4\text{\textsuperscript{-}}. The experiment was performed in triplicate.

**Cyanobacterial lysis**

Six-ml aliquots (about \( 6 \times 10^8 \) cells ml\textsuperscript{-1}) obtained from 40-day-old cultures in BG-11 and BG-11 containing 2.4 mM Cl\text{O}_4\text{\textsuperscript{-}} (as reported above) were centrifuged at 7000 \( \times \)g, at 20°C for 15 min. Pellets were washed twice in dd-H\text{2}O, air-dried overnight under a laminar-flow hood, and weighted prior to and after drying. Air-dried cells (30 mg) were resuspended in 3 ml dd-H\text{2}O and split into three aliquots. Pellets from each 1 ml aliquot were lysed as follows: (i) mortaring for 2 min, (ii) resuspension in 500 \( \mu \)l dd-H\text{2}O and immersion in liquid N\text{2}, for 1 min, (iii) thawing at 37°C for 10 min; (iv) centrifugation at 6000 \( \times \)g, for 10 min at 20°C to collect the soluble supernatant used as lysate medium.

**Escherichia coli growth with cyanobacterial lysate-based medium**

One-ml aliquots of overnight \( \text{E. coli} \) were diluted to about \( 1 \times 10^6 \) cells ml\textsuperscript{-1}, washed with 1X PBS, and resuspended in 1 ml lysate-based medium. As a positive control, \( 1 \times 10^6 \) cells ml\textsuperscript{-1} were washed with PBS and inoculated in 1 ml of M9 minimal medium (Sambrook et al., 1989) supplemented with 0.5% glucose. As negative controls, about \( 1 \times 10^6 \) cells ml\textsuperscript{-1} were washed with PBS and inoculated into 1 ml of sterile dd-H\text{2}O and 1X PBS. Each sample was incubated in 1.5 ml Eppendorf tubes at 38°C
overnight, under continuous shaking of 180 rpm. The experiment was performed in triplicate. 

Perchlorate measurements

Perchlorate concentrations were determined by EPA Method 331.0 Rev. 1.0 – Liquid Chromatography/Electrospray Ionization/Mass Spectrometry (2012). Briefly, samples were diluted 1:20 with ultrapure H₂O and 5 μl injected for analysis on a Waters ACQUITY I-Class UPLC/Xevo G2-XS QTof system. The quantification of perchlorate anion was based on Tof MRM negative acquisition mode of the following masses, under −25 V collision energy and by using a calibration curve that covered 0.01–10 mM concentration interval.

Results

Cyanobacterial tolerance towards increasing perchlorate concentrations

The perchlorate tolerance of *Chroococcidiopsis* strains CCME 029 and CCME 029 P-MRS was investigated by exposure for 55 days in liquid BG-11 medium containing increasing concentration of Na-, Mg- and Ca-perchlorate up to 100 mM (Fig. 1). The presence of 5 mM of Na-, Mg- and Ca-perchlorate (5, 10 and 10 mM perchlorate ions, respectively) did not inhibit the growth of the two strains. While the presence of 50 mM perchlorate resulted in a reduced growth rate of both strains, that was more evident in Mg- and Ca-perchlorate (100 mM perchlorate ions), than in Na-perchlorate (50 mM perchlorate ions). The growth of both strains was reduced by the 100 mM concentration of Na-perchlorate, while no growth occurred in 100 mM Mg- and Ca-perchlorate (200 mM perchlorate ions) (Fig. 1).

Effects of increasing perchlorate concentrations on cyanobacterial morphology

The exposure to increasing concentration of Na-, Mg- and Ca-perchlorate did not show any severe morphological changes. For example, after 55 days of exposure in 100 mM Na-perchlorate, *Chroococcidiopsis* sp. CCME 029 (Fig. 2a) and CCME 029 P-MRS (Fig. 2b) occurred as single cells and as four-celled aggregates as typical of strain CCME 029 when grown in BG-11 medium (Fig. 2c).

Effects of Mars-relevant perchlorate concentration on cyanobacterial growth

The ability of *Chroococcidiopsis* strains CCME 029 and CCME 029 P-MRS to grow in the presence of Mars-relevant perchlorate concentrations was investigated by incubation in liquid BG-11 containing 2.4 mM ClO₄⁻ provided as a 40% Mg-perchlorate and 60% Ca-perchlorate mixture (Fig. 3). The combination of the two salts did not affect the growth rate. After 40-day incubation, strains CCME 029 and CME 029 P-MRS showed a slight increase in cell densities compared to control cells grown in BG-11 medium (Fig. 3).

Perchlorate content during cyanobacterial growth

The concentration of perchlorate ions in *Chroococcidiopsis* strains CCME 029 and CCME 029 P-MRS was measured at the beginning and after 40-day incubation in BG-11 medium with 2.4 mM perchlorate ions. In the overall, no significant variation occurred between the perchlorate concentrations in the BG-11 medium recovered after the cyanobacterial growth and that in control BG-11 medium with 2.4 mM perchlorate ions.

Escherichia coli growth with cyanobacterial lysate-based medium

A cyanobacterial lysate-based medium was derived from the soluble fraction of lysed cells of *Chroococcidiopsis* sp. CCME 029 grown for 40 days in BG-11 with 2.4 mM perchlorate ions (Fig. 4). When about 1 × 10⁶ *E. coli* cells were inoculated in 1 ml of the lysate-based medium obtained from the lysis of about 30 mg of air-dried cyanobacteria, the cell density resulted increased to about 2 × 10⁸ cells ml⁻¹ after overnight incubation. No significant difference occurred in *E. coli* cell density when using the lysate obtained from strain CCME 029 grown in

![Fig. 1. Cell densities of cultures of *Chroococcidiopsis* sp. CCME 029 and CCME 029 P-MRS grown in BG-11 containing 5, 50 and 100 mM Mg-, Ca- and Na-perchlorate. Control cultures were grown in BG-11. Data are shown as mean ± standard deviation in three independent trials.](https://doi.org/10.1017/S1473550420000300) Published online by Cambridge University Press
BG-11 in the absence of 2.4 mM perchlorate ions. *Escherichia coli* reached an average density of $4 \times 10^8$ cells ml$^{-1}$ after incubation in M9 minimal medium (containing salts, nitrogen and 0.5% glucose) and about $1 \times 10^9$ cells ml$^{-1}$ in LB medium. No growth occurred when *E. coli* was inoculated in dd-H$_2$O or PBS (Fig. 4).

**Discussion**

The aim of this work was to investigate whether the desert cyanobacterium *Chroococcidiopsis* could be used in Bio-ISRU technologies to support the growth of BLSS components, namely bacteria, that need fixed carbon and that might be perchlorate sensitive (Rothschild, 2016; Verseux et al., 2016).

The incubation of *Chroococcidiopsis* sp. CCME 029 and CCME 029 P-MRS in increasing perchlorate concentrations identified a perchlorate threshold value at 100 mM perchlorate ions. No remarkable significant difference in the perchlorate tolerance occurred between CCME 029 and its space-derivate CCME 029 P-MRS, suggesting that the exposure to Mars simulations in low Earth orbit followed by rehydration on the ground after retrieval did not select any enhanced oxidative-stress resistance. Indeed on-going comparative genomic analysis between the genome sequence of strain CCME 029 and its space-derivate CCME 029 P-MRS will verify the eventual presence of differences between the two strains.

The identified perchlorate tolerance provided a first prerequisite for the use of *Chroococcidiopsis* in Bio-ISRU technologies. In contrast, at the Phoenix landing site, a concentration of about 2.4 mM perchlorate ions (0.5 wt %) was measured, whereas the Curiosity rover reported perchlorate concentrations up to 1% at the Gale Crater (Glavin et al., 2013).

Long-term (55 days) exposure of the two *Chroococcidiopsis* strains to increasing perchlorate concentrations revealed that growth rate was: (i) not affected in 5 mM Na-, Mg- and Ca-perchlorate (5, 10 and 100 mM perchlorate ions); (ii) slightly reduced in 50 mM Na-perchlorate; (iii) further impaired in 100 mM Na-perchlorate; (iv) reduced in 50 mM Mg- and Ca-perchlorate (100 mM perchlorate ions); and (v) completely inhibited in 100 mM Mg- and Ca-perchlorate (200 mM perchlorate ions). No growth occurred in 300 mM Na-perchlorate (not shown).
The identified 100 mM perchlorate ion threshold was comparable to that of a halophilic bacterium isolated from Big Soda Lake, growing in 2% Na-perchlorate, that is, 160 mM perchlorate ions (Matsubara et al., 2017). However, Chroococcidiopsis perchlorate tolerance was lower than that of several halophiles. For example, halophilic Archaea of the family Halobacteriaceae and Halomonas elongata showed a weak growth in 600 mM Na-perchlorate (Oren et al., 2014), the growth of the archaeon Halorubrum lacusprofundi was still measurable in 800 mM perchlorate ions (Laye and DasSarma, 2018), while Hydrogenothermus marinus did not replicate in 400 mM Na-perchlorate (Beblo-Vranesvic et al., 2017). So far the highest perchlorate tolerance was reported for the yeast Debaryomyces hansenii growing in 2.4 M Na-perchlorate (Heinz et al., 2020), a value twice that of the bacterium Planococcus halocryophilus (Heinz et al., 2019).

Short-term (15 min) exposure in Na-perchlorate highlighted $D_{10}$ values of 2.7, 1.3, 5 and >5 M, for Deinococcus radiodurans, E. coli, B. subtilis spores and H. marinus, respectively (Beblo-Vranesvic et al., 2017). In the present work, Chroococcidiopsis strains were not exposed to short-term incubation in higher perchlorate concentrations because the main goal was to identify the tolerance threshold in long-term cultivation.

No significant morphological changes were observed in Chroococcidiopsis exposed up to 200 mM perchlorate ions. On the contrary, halophilic archaea occurred as swollen cells in perchlorate ion concentrations $>$200 mM (Oren et al., 2014). H. marinus turned into long cell chains in Na-perchlorate concentrations $>$100 mM (Beblo-Vranesvic et al., 2017), while P. halocryophilus formed large cell clusters in 1.1 M Na-perchlorate (Heinz et al., 2019).

The two Chroococcidiopsis strains better tolerated Na- and Mg-perchlorate than Ca-perchlorate. Indeed Ca-perchlorate is a ch cryptocurrency, causing macromolecules destabilization and growth inhibition stronger than Mg-perchlorate and Na-perchlorate, the latter being a weak ch cryptocurrency agent (Craty et al., 2013; Nagler and Moeller, 2015). Moreover, since Chroococcidiopsis sp. CCME 029 has been reported to bleach in 680 mM NaCl (Hershkovitz et al., 1991), the observed slightly reduced growth in 100 mM Na-perchlorate might be ascribed to the oxidizing action of perchlorate.

Since 2.4 M Na-perchlorate ions were reported to occur on Mars as a 60 and 40% mixture of Mg- and Ca-perchlorate, respectively (Hassler et al., 2014), the two Chroococcidiopsis strains were grown in Mars-relevant perchlorate ions as reported for the Phoenix landing site. The growth of the two Chroococcidiopsis strains was not impaired by 40-day incubations in 2.4 mM perchlorate ions, rather it increased slightly compared to control. This might be due to a lack of a synergistic negative effect of Mg- and Ca-perchlorate as reported for bacteria isolated from Big Soda Lake (Matsubara et al., 2017), but also to a positive effect of the increased cation concentration. Indeed Anabaena sp. PCC 7120 was reported to grow in 1 mM CaCl$_2$ (Singh et al., 2016). Here Chroococcidiopsis cells were grown in concentrations of 0.24 mM CaCl$_2$ and 0.30 mM MgSO$_4$ supplied by the BG-11 medium (Rippka et al., 1979) while 0.5 mM Mg$^{2+}$ and 0.75 mM Ca$^{2+}$ were present in 2.4 M perchlorate ions supplied as a 60 and 40% mixture of Mg- and Ca-perchlorate, respectively.

After 40-day growth in 2.4 mM perchlorate ions, no variation in the perchlorate content was observed; therefore, the perchlorate concentration in the lysate was not determined. Indeed cyanobacteria are non-perchlorate-reducing prokaryotes, although the presence of chloride dismutase-like proteins has been reported in a few nitrogen-fixing species (Nerenberg, 2013; Schaffner et al., 2015). Hence, the perchlorate tolerance of the two Chroococcidiopsis strains was likely a consequence of their efficient antioxidant system (Fagliarone et al., 2017).

The observed Chroococcidiopsis capability to tolerate Mars-relevant perchlorate ions was further exploited to obtain a lysate-based medium to feed an E. coli strain, capable of utilizing sucrose as a carbon source (Lee and Chang, 1993). Therefore, after 40-day growth in 2.4 M perchlorate, Chroococcidiopsis cells were air-dried to induce a fivefold increase in their sucrose content from, up to 5 mg/g dry weight (not shown), as previously reported (Fagliarone et al., 2020). The lysate-based medium obtained from 10 mg of dried cyanobacteria supported an increase of the E. coli cell density from $1 \times 10^8$ to $2 \times 10^9$ cells after overnight incubation. A comparable increase of E. coli cell density was obtained using the lysate obtained from Chroococcidiopsis grown in standard BG-11 medium. This suggested that washing the cyanobacterial cells before lysis removed perchlorate and avoided a negative effect on E. coli growth. However, since a doubled cell density (about $4 \times 10^8$ cells) was supported by overnight growth in M9 minimal medium, an increased E. coli cell density is expected to be supported by lysing an increased amount of dried cyanobacteria.

However, results provided a first proof-of-concept of feeding a bacterium with a lysate-based medium obtained from Chroococcidiopsis grown in Mars-relevant perchlorate concentration. The use of this cyanobacterium as a pioneer for Bio-ISRU on Mars is further supported by its capability of lithotrophic growth on Mars regolith analogues, namely anorthosite and basal (supplemented with NaNO$_3$), resulting in elemental release and biomass production (Olsson-Francis and Cockett, 2010).

Moreover, the feeding of E. coli with a cyanobacterial lysate is relevant since this bacterium is a typical chassis for genetic manipulation and metabolic engineering of bacteria was proposed to provide consumables to human outposts on Mars (Rothschild, 2016). Recently, the use of cyanobacterial lysate to support B. subtilis engineered to synthetize aromatic polymers for space application has been reported (Aversch and Rothschild, 2019) and tested in NASA’s PowerCell payload aboard the DLR EuCROPIS Satellite (McCutcheon et al., 2015).

In conclusion, Chroococcidiopsis might enable Bio-ISRU on Mars by combining its capability of converting CO$_2$ to organic compounds with its perchlorate resistance, and its use in Bio-ISRU might be further reinforced by synthetic biology (Montague et al., 2012; Snyder et al., 2019). Indeed, the development of Chroococcidiopsis CCME 029 as a chassis for synthetic biology will endow it with the ability to synthetize compounds not naturally produced and this approach will take advantage of the availability of a genetic system for its synthetic manipulation (Billi et al., 2001) and maintenance of plasmid DNA after prolonged air-dried storage (Billi, 2012).

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