NIAC Phase I Grant “Mars Ecopoiesis Testbed” NNX14AM97G Final Progress Report.

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Abstract/Executive Summary

Mars surface conditions where liquid water is absent were simulated for the purposes of laboratory research. A pressure-temperature (P-T) profile was maintained in which no combination of pressure or temperature corresponds to the liquid region of the water phase diagram. The triple point of pure water occurs at T = 0.1°C and P(H2O) = 6.01 mbar; therefore all temperatures and pressures must be kept below these values, respectively. A 35-day test was performed in a commercial planetary simulation system (Techshot, Inc., Greenville, IN) in which the minimum night-time temperature was -80°C, the maximum daytime temperature was +26°C, the simulated day-night light cycle in earth hours was 12-on and 12-off, and the total pressure of the pure CO2 atmosphere was maintained below 11 mbar. Any water present was allowed to equilibrate with the changing temperature and pressure. The gas phase was sampled into a CR1-A condensation-mirror low-pressure hygrometer, which uses liquid nitrogen (down to 77°K) to determine the dew point (Buck Technologies, Boulder, CO). Dew point was measured once every hour and recorded on a data logger, along with the varying temperature in the chamber, from which the partial pressure of water was calculated. The resulting calculated daily cycles were tracked on the water P-T diagram, and no points were found to fall within the liquid-phase region of the diagram. It is concluded that there was no liquid water present throughout the test except during the initial pump-down period when aqueous specimens were introduced on the first day (less than 1 hour). Mars regolith simulant was present during this test, and further investigation is needed to determine whether liquid water could have been present or absent in the regolith in the form of brine. Biological samples consisting of Cyanobacteria: *Anabena* sp., *Chroococcidiopsis* CCME171, *Plectonema boryanum*; Eubacteria: *Bacillus subtilis*, *Pseudomonas aeruginosa*, and Eukaryota: *Chlorella ellipsoidia* were maintained in the simulator under the above-described conditions. The exposed specimens were tested for intracellular esterase activity, chlorophyll content (where appropriate) and reproductive survival. All tests yielded low-level positive results in all cases. In parallel to these terrestrial studies a planned design study was undertaken for the proposed test bed. Design requirements include compact assembly for transport and installation on the planetary surface (multiple units per mission would be expected), protective internal package for the release of organisms, a means of atmosphere exchange, access to sunlight, a means of penetrating the planetary surface, and most importantly a means of acquiring regolith while meeting the requirements of planetary protection. In consultation with advisers a design was created, and a large-scale mock-up of this design was fabricated by additive manufacturing at Techshot, Inc. with moving parts that simulated the components of the design. The mock-up assembly has been demonstrated to interested parties. A means of detecting live metabolism will also be included in the test bed. Several options were reviewed, and it is concluded that, by the time the ecopoiesis test bed is ready for testing the optimum instrument will be the equivalent of a hand-held
mass spectrometer for metabolic gas analysis. This will maximize versatility and reveal much more information that could a detector of a single product (such as molecular oxygen), and the simple output signals will be compatible with telemetry. The objectives of this project, (1) Model and test the availability of liquid water in Techshot’s Mars simulator facility, (2) Identify current candidate pioneer organisms for testing, and initiate a selection program, (3) Create a mechanical and electronic design concept for Mars surface shallow penetrator with planetary protection, and (4) Identify electronic biological activity tests, were fulfilled by the completion of this Phase-1 research described in this final report.

Introduction

The cover of Astrobiology, September 2015, “Celebrating 15 Years” features, in bold color, photosynthetic cells reacting with extraterrestrial water to form hydrogen ions – the initial process in photosynthesis – within a microbial fuel cell. The molecular oxygen produced is reduced to water at the cathode generating an electric current proportional to the diurnal levels of insolation as a proposed means of detecting extraterrestrial life (Figueroedo et al., 2015). Active life cannot be sought in the absence of water and cannot be implanted in the absence of water. Mars orbital photographic evidence for the slow movement of perchlorate evaporites down slopes at Garni and Gale craters is currently taken as evidence that flowing brine may be responsible for recurring slope lineae on these steep slopes (Martin-Torres et al., 2015). The high salinity and low temperatures that correspond to this transient liquid condition (certain times of day) may not be hospitable toward terrestrial extremophiles. These recent findings further set the stage for searches for microbial life and ecopoiesis research.

What is a Mars Ecopoiesis Test Bed?

The term Ecopoiesis was introduced by Bob Haynes and Chris McKay, in collaboration with Carl Sagan in the 1970’s [Sagan, 1973], and conferences were held on this subject [Haynes, 1992; McKay, 1989, 1991, 2004]. McKay and Averner and others performed early calculations concerning the use of dark materials on the Mars polar caps, enhanced insolation and/or artificial greenhouse gases to initiate a runaway greenhouse effect that would result in conditions allowing liquid water to exist in abundance on the Red Planet [Averner, 1976; MacElroy, 1976; Haynes, 1992; Fogg, 1995; Gerstell, 2001; McInnes, 2006]. Our proposed concept emphasizes bio-ecopoiesis [McKay, 1989, 1997; Thomas, 1995], in which contained pioneer organisms will eventually be tested for bio-activity in a suitable location on the Martian surface (low latitude and low altitude, where pressure and temperatures combine to flirt with the possibility of metastable liquid water [Hecht, 2002; Heldmann, 2005; Carr, 1996; Jakosky, 1992; Levin, 2003]) using a robotic mechanism on a future rover.

The proposed concept is illustrated diagrammatically in Figure 1.
Figure 1. Rendering of the Mars surface shallow penetrator showing, from top to bottom: transparent dome for the admission of light for photosynthesis, porous ceramic ring that allows free exchange between the internal vapor phase and Mars atmosphere while providing planetary protection, sensor of biological activity (TBD) connected to datalink to a Mars orbiter for relaying signals, specimen containers that robotically release one set of contents onto the surface and one set of contents below the surface, threads that cause the penetrator to penetrate the regolith when rotated, the containment cylinder which constitutes the entire shell of the penetrator and which contains the sample of regolith to be tested, and at the very bottom the planetary protection seal that will be formed in situ after the penetrator is at full depth and before the release of specimens.

Relevance to NASA Astrobiology Roadmap

Goal 6.2:

Objective 6.2—Adaptation and evolution of life beyond Earth.
Explore the adaptation, survival and evolution of microbial and other organisms under environmental conditions that simulate conditions in space or on other potentially habitable planets. Identify survival strategies to evaluate the potential for interplanetary transfer of viable organisms and to establish requirements for effective planetary protection. Identify and validate roles that microorganisms might play in life support and resource acquisition during human missions envisioned by US Space Policy. Develop tools to track the function and adaptation of microbes and other organisms to extraterrestrial environments during mankind’s exploration efforts.

Example investigations

Document the effects of the space environment upon microbial ecosystems.
Examine the survival, genomic alteration, and adaptation of microbial ecosystems in a wide range of simulated martian environments. Interpret the significance of these experiments regarding the potential for the forward biological
contamination of Mars and for utilizing microorganisms to support the needs of human exploration. Examine the effects of the space environment upon the biosynthesis and utilization of biomolecules that play key roles in biogeochemical processes and also upon the viability of microbes that might be transferred between planets by natural processes (e.g., impact ejection). Develop automated assay tools to monitor the adaptation of organisms in lunar and martian environments, especially those areas most likely to be visited by human explorers over the next century.

Team Members and Their Contributions

A core scientific and engineering team will be assembled in Phase I. This will be accomplished by the PI, the Project Scientist, The Project Engineer, The Project Astrobiologist, and the Project Adviser. These Phase-I activists are:

PI: Dr. Eugene Boland, Chief Scientist Techshot, Inc. (resume in Section 5)
Project Scientist: Dr. Paul Todd, Chief Scientist Emeritus, Techshot, Inc. (resume in section 5)
Project Engineer: Mr. Michael (Andy) Kurk, Techshot, Inc. (resume in section 5)
Project Astrobiologist: Prof. David J. Thomas, Professor, Lyon College
Project Adviser: Dr. Lawrence Kuznetz, Principal, Spinoff, NASA, Retired

The participating engineers will be drawn from Techshot’s space-hardware-experienced staff, and participating scientists will be drawn from Drs. Thomas’ and Todd’s international circle of colleagues.

Objectives of the Phase I Project

The Phase I objectives, exactly as briefly stated in the proposal, were as follows:

Objective 1. Model and test the availability of liquid water in Techshot’s Mars simulator facility. A low-temperature hygrometer will be acquired, and conditions surrounding 7-10 mbar will be sampled by interrupting experiments at specific times in Martian sols and measuring regolith moisture and by monitoring the formation of mineral evaporites.

Objective 2. Identify current candidate pioneer organisms for testing, and initiate a selection program.

Objective 3. Create a mechanical and electronic design concept for Mars surface shallow penetrator with planetary protection.

Objective 4. Identify electronic biological activity tests (O₂ sensor, for example); initiate testing in a laboratory simulator.

Progress on Phase I Objectives
Objective 1. Model and test the availability of liquid water in Techshot’s Mars simulator facility. A low-temperature hygrometer will be acquired, and conditions surrounding 7-10 mbar will be sampled by interrupting experiments at specific times in Martian sols and measuring regolith moisture and by monitoring the formation of mineral evaporites.

Rationale

Ecopoiesis will require water. That means maximizing the chances of liquid-phase water being transiently present in the test bed with the most likely sites being found at Mars’ lowest altitudes and latitudes [Kuznetz, 2006]. A preliminary identification of these “landing” sites, already considered for certain past and future robots, is given briefly in Table 1. The tidal pressure swings of ±0.5 mbar need to be considered. These sites are also thought to contain evaporites, possibly including nitrates (all of which are water soluble) to provide nitrogen and magnesium salts [Tosca, 2006]. Recent results from the Curiosity Rover in Gale Crater are encouraging with regard to the availability of minerals to support autotrophic life [Navarro-González, 2013]. The big question of course has to do with the thermodynamics and transport processes of water in real and simulated Martian environments. Even at 11 mbar, the vapor pressure of water is well below the 6.1-mbar triple point, where, at increased temperature ice will normally sublume. However, speculative calculations modeling the diffusion of water vapor from ice surfaces during sublimation indicate a local (within a few mm of ice) increase in water vapor concentration to some 60%, or the required 6.1 mbar in the 11 mbar environment [Levin and Weatherwax, 2004]. Therefore, early proposed research will use the Techshot simulator [N. Thomas, 2006] to test such hypotheses.

Table 1. Characteristics of potential Martian test venues.

<table>
<thead>
<tr>
<th>SITE NAME</th>
<th>LATITUDE</th>
<th>MAX DEPTH, m</th>
<th>MAX P, mb</th>
<th>MAX T °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elysium Planitia</td>
<td>3°N</td>
<td>45</td>
<td>6.2</td>
<td>32</td>
</tr>
<tr>
<td>Isidis Planitia</td>
<td>3.0-12.9°N</td>
<td>3,600</td>
<td>7.5*</td>
<td>26</td>
</tr>
<tr>
<td>Valles Marineris</td>
<td>13.9°S</td>
<td>7,000</td>
<td>11.0</td>
<td>26</td>
</tr>
<tr>
<td>Gale Crater</td>
<td>4.5°S</td>
<td>4,500</td>
<td>7.8</td>
<td>28</td>
</tr>
<tr>
<td>Triple Point</td>
<td></td>
<td>6.1</td>
<td>0.02</td>
<td></td>
</tr>
</tbody>
</table>

*Approximate, based on linear interpolation.

Improved simulator facility

The Mars simulation facility has been upgraded with an automated power supply that can control the Martian sol by automatically firing the arc lamp (and confirming it lit) and tying the power and temperature output to a data logger. Previously this was a manual process to confirm the arc lamp fired. We attained the use of a cryogenic hygrometer to monitor moisture continuously and subjected it to testing to verify that we can produce the pressures and vapor pressure requirements set forth for the research.

To maintain a Martian atmospheric environment, the sample is loaded into a 140 mm ID X 430 mm long quartz-tube chamber featuring a single open end. Once loaded, within the tube, the
open end of the tube is sealed with a stainless steel end cap that features all of the ports and valves necessary to maintain a vacuum, monitor temperature and pressure, and input a simulated Martian atmosphere or liquid water. A photograph of the Mars Simulator Chamber within the environmental cabinet is shown in Figure 1.

Figure 1. Interior of the Mars simulator environmental cabinet showing reflecting mirror, quartz chamber, sample tray, eighteen samples with twelve of them in direct illumination and six of them in shade, thermistor leads for temperature measurement in and out of regolith, end plate for pressure and electrical access, and cradle to hold quartz-tube Mars chamber. This configuration was used in an uninterrupted five-week simulation campaign.

Once the experiment loading is complete the quartz chamber is secured within a thermal cabinet and all external mechanical and data interfaces are connected to the sealing end plate. Upon completion of this process, the inner volume of the simulator chamber is drawn down to the desired Martian atmospheric pressure by a Welch Reitschle Thomas vacuum pump. To accommodate most very low pressure experiments the vacuum pump is required to run continuously. If minor pressure adjustment is required, a ball and needle valve are located downstream of the experiment between a low pressure hygrometer and the vacuum pump. When needed the valves can be adjusted to obtain the desired pressure.

In addition to control and measurement the Mars Simulator Chamber end plate contains a port through which liquid water or a gas can be periodically introduced to the test volume throughout the experiment. Periodically CO₂ or a specially mixed gas resembling the Martian atmosphere is introduced into the volume to flush any dry nitrogen that may have intruded into the volume and maintain an atmosphere consistent with that of the simulation objectives. This function is fully automated and its frequency is programmed into the thermal cabinet’s Watlow controller as part of the experiment profile.
The downstream hygrometer is used to continuously sample the moisture content of the atmosphere within the Mars simulator chamber. The hygrometer data are monitored along with pressure and temperatures and are recorded using a Fluke 1586A Super DAQ data logger.

The Martian thermal environment is simulated by a highly insulated, modified, cryogenic thermal cabinet (Model ZBD-108 LN$_2$ Cooled Chamber, Associated Environmental Systems, Ayer, MA) that is thermally maintained by the evaporation of liquid nitrogen. Since the chamber is filled with dry nitrogen there is very little water present, and heavy frosting is avoided. Thermal cycling of the chamber, according to the Mars daily cycle, is accomplished by programming a Watlow F4 Controller that was provided as a standard component of the stock thermal chamber. This controller is programmed with a timeline that enables thermal ramping functions. The programming can be performed either on the user panel or with a GUI on the PC.

<table>
<thead>
<tr>
<th>Table 2. Customized Liquid Nitrogen Cooled Environmental Test Chamber Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Working Volume: 24” x 24” x 24”</td>
</tr>
<tr>
<td>Insulation: 4” Fiberglass</td>
</tr>
<tr>
<td>Power Requirements: 120 VAC, 1 phase, 60 Hz</td>
</tr>
<tr>
<td>Refrigeration System: LN$_2$ Cooled</td>
</tr>
<tr>
<td>Temperature Range: -135° C to +177° C</td>
</tr>
<tr>
<td>Temperature Stability: +/- 1/2° C at sensor</td>
</tr>
<tr>
<td>Temperature Rise Time: Ambient to upper limit – 20 minutes</td>
</tr>
<tr>
<td>Temperature Pull Down Time: Ambient to lower limit – 20 minutes</td>
</tr>
<tr>
<td>Interior: 18 Gauge 304 Stainless Steel</td>
</tr>
<tr>
<td>Illumination: Double-paned quartz window</td>
</tr>
<tr>
<td>18 Gauge cold rolled steel with two coats of textured epoxy paint</td>
</tr>
<tr>
<td>Watlow F4 Programmable Controller with RS232 communications</td>
</tr>
<tr>
<td>Dry Nitrogen Purge</td>
</tr>
</tbody>
</table>

Simulated Solar energy is provided to the sample by an automated Sciencetech Solar Simulator that has been equipped with an AM0 filter to more closely resemble the mildly filtered solar radiation which reaches the Martian surface. A diagram of the unfiltered (top) output of the xenon arc lamp solar and the output spectrum after filtering with an AM0 filter (bottom) is shown in Figure 2.
Figure 2. Light spectrum produced by the 1,000 W xenon arc illuminator without (above) and with (below) AM0 (Air Mass zero) light filter. The similarity to the solar spectrum below about 300 nm is important in terms of its known biologically damaging action.

As part of the improved solar simulator automation package the user can preprogram a light cycle and intensity. For this experiment a 12 hour, 1000 W cycle was preprogrammed prior to initiation. The ScienceTech illuminator and the added control technology is shown in Figure 3. Modifications to the stock liquid nitrogen cooled thermal chamber, enable the solar simulator’s light beam line to pass into the thermal chamber through a 4” x 4” (10 x 10 cm) double paned opening after which it is reflected by a front-surface mirror inside the cabinet to illuminate the quartz Mars jar chamber as shown in Figure 1. A photograph detailing the light path from the solar simulator is shown in Figure 4.
Nine variables were measured and logged as a function of elapsed time every 5 minutes for 5 weeks. These were ambient laboratory temperature, Mars-jar pressure, cabinet temperature, regolith temperature, Mars-jar temperature, hygrometer pressure, dew/frost point, illuminator on/off, and water concentration.
An example of measured values during days 10-20 is given in Figure 5. The pressure pattern is seen to have spikes due to the hourly introduction of fresh CO₂ to maintain the atmospheric composition. The purpose of this operational procedure was to prevent cabinet nitrogen from entering the simulated atmosphere inside the chamber.

![Figure 5](image.png)

**Figure 5.** Example of environmental data logged over a 10-day period, showing four measurements: regolith temperature (red), chamber internal temperature (green), chamber internal pressure (blue), and solar simulator on/off (violet). The “Days” on the abscissa are earth days and not Martian sols.

**Moisture Measurement and Control in the Laboratory Test Bed**

Nine parameters were logged during test and experimental operation of the laboratory test bed. Of these, the CR-1A cryogenic hygrometer (Buck Research Instruments, LLC, Boulder, Colorado) recorded temperature, pressure and dew/frost point in extracted chamber atmosphere. Figure 6 is a chart record of nine logged parameters during days 11-14 of a 35-day campaign. The dew/frost point never exceeded the regolith temperature except when a pulse of liquid water was injected when the temperature was below -75 C. At all other times humidity was well below saturation and the partial pressure of water was below the triple point (see Figure 7).
Figure 6. Chart record of nine logged parameters during days 11-14 of a 35-day campaign. The dew/frost point never exceeded the regolith temperature except when a pulse of liquid water was injected when the temperature was below -75 C. At all other times humidity was well below saturation and the partial pressure of water was below the triple point (see Figure 8).

Spikes in pressure and dew/frost point that appear on the chart are due to the periodic injections of fresh dry CO\textsubscript{2} for maintenance of atmosphere composition. From the data set of Figure 6 were extracted correlated time points for chamber pressure (as measured by the CR-1A hygrometer) and regolith temperature. Figure 7 represents the repeated journey around the P-T diagram through three daily cycles. The total pressure never exceeded values measured on the Martian surface.
The CR-1A Cryogenic Hygrometer measured, and the data logger recorded dew/frost point at each sampling time. From the recorded dew/frost points the partial pressure of pure water was calculated using the empirical formula

$$p_w = 6.11 \times 10^3 \left[7.5 \frac{T_d}{273.3 + T_d}\right]$$

using $p_w =$ water partial pressure in mbars and $T_d =$ dew/frost point in °C. For 814 time points accumulated over days 11-14 of a 35-day campaign this partial pressure is plotted against temperature on a traditional P-T diagram, on which the liquidus line for pure water is also shown (Figure 8).
Performance record: A review of all data sets was made to create a list of “bumps in the road”. The CR-1A hygrometer must be cooled below the chamber temperature at all times, and cooling was lost on a small number of occasions during the 35-day campaign; however, the maintenance of environmental conditions did not depend on continuous hygrometer readings, and its temperature and pressure readings are redundant. The hygrometer only contributed data, and hygrometer-related events are noted by an asterisk in Table 2. Some maintenance of the solar illuminator, including xenon arc lamp replacement was required toward the end of the campaign.

Table 3. Descriptions of individually recorded data sets during 35-day Mars surface simulation campaign.

<table>
<thead>
<tr>
<th>DAYS</th>
<th>OBSERVATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,2</td>
<td>Breaking in. Chamber pressure rose to 14-20 mbar for 5 min during hourly CO₂ injection</td>
</tr>
<tr>
<td>2-4</td>
<td>Pressure maintained &lt;10 mbar. Hygrometer cooling lost; hygrometer data not taken*</td>
</tr>
<tr>
<td>4,5</td>
<td>Same as days 11-14, which were used as model for detailed report data</td>
</tr>
<tr>
<td>5-8</td>
<td>Loss of hygrometer function on day 8*</td>
</tr>
<tr>
<td>8</td>
<td>Hygrometer recovery period; tube pressure spike to 70 mbar for 2 h*</td>
</tr>
<tr>
<td>9-11</td>
<td>Same as days 11-14</td>
</tr>
<tr>
<td>11-14</td>
<td>Chosen as source for model data</td>
</tr>
<tr>
<td>14-16</td>
<td>Same as days 11-14; spike in measured dew/frost point on day 15</td>
</tr>
<tr>
<td>16-18</td>
<td>Same as days 11-14</td>
</tr>
<tr>
<td>18-21</td>
<td>Same as days 11-14</td>
</tr>
<tr>
<td>21-24</td>
<td>Loss of hygrometer cooling, days 23,24*</td>
</tr>
<tr>
<td>25-30</td>
<td>Loss of hygrometer cooling, day 29*</td>
</tr>
<tr>
<td>30-36</td>
<td>Loss of light day 31, ½ of day 32, day 35. Repression recorded when campaign ended.</td>
</tr>
</tbody>
</table>

*Losses of hygrometer function have no impact on environment control.

The data above demonstrate the continued presence of water in the solid and vapor phases throughout this 35-day campaign. At the end of the campaign the chamber was repressurized for sample recovery, at which time two observations on liquid water were made. There had been an accumulation of some 10 cm³ of liquid water, melted from the solid and condensed from the vapor phases of the chamber upon its return to earth-ambient conditions, and the regolith simulant in the shaded specimen jars was dark due to moisture while the regolith in the shaded specimens were the color of dry regolith.

Objective 2. Identify current candidate pioneer organisms for testing, and initiate a selection program.

The project officially kicked off internally with the convening of the advisory panel consisting of Dr. Larry Kuznetz (Advisory council chair), Dr. Christopher House and Dr. David Thomas. Unfortunately, Dr. Chris McKay was unable to attend the first meeting because of commitments with Curiosity Rover results.

The first key outcome of the meeting was the identification of test organisms. Due to the size of our Mars simulator and past experience, we decided to proceed with 3 cyanobacteria and an alga. Space
permitting, we would add two species of heterotrophic eubacteria, which actually was done. The final selections are given in the following list.

**Organisms:**
*Anabaena* sp. (cyanobacteria)
*Chroococcidiopsis* sp. CCMEE171 (cyanobacteria)
*Plectonema boryanum* UTEX485 (cyanobacteria)
*Chlorella ellipsoidea* (algae)
*Bacillus subtilis* (bacteria)
*Pseudomonas aeruginosa* (bacteria)

A single test was designed for a 5-week period, to start after simulator improvements and physical tests were completed.

A single 5-week simulation was performed using 100% CO₂ at pressures between 3 and 10 mbar and water supplementation as described in Figures 2-10. Multiple samples of each species were subjected to four conditions for 5 weeks for comparison: Mars simulator, -80 C in darkness, +4 C in darkness and 25 C in diurnal illumination.

The following test organisms were used:
Cyanobacteria: *Anabena sp.*, *Chroococcidiopsis* CCMEE171, *Plectonema boryanum*
Eubacteria: *Bacillus subtilis*, *Pseudomonas aeruginosa*
Eukaryota: *Chlorella ellipsoidea*

As in previous work the production of fluorescence by samples resuspended in aqueous solution was used as a test for the presence of intact cells containing esterases using the fluorescein diacetate (FDA) test. Figure 20 is a summary of the resulting measurements on suspended regolith samples.
Figure 20. Rate of FDA hydrolysis by 1 gram of regolith containing each of the six species exposed, measured as rate of increase in fluorescence intensity.

When specimens were returned to ambient temperature and pressure liquid water present in the samples was tested for biological activity. Figure 21 is a summary of the resulting measurements of FDA conversion in liquid water samples.
Figure 21. Rate of FDA hydrolysis by 1 mL of water residue obtained from each of the six species exposed, measured as rate of increase in fluorescence intensity.

Samples were streaked for growth on semisolid media and assessed for colony counts on a 4-unit scale. Many plates had too many colonies to count, so an area-coverage scale of 0 – 4 was used to characterize growth, using 25 C diurnal samples as control, identified as a score of 4. These results are summarized in Figure 22.
Figure 22. Relative phototrophic growth of cyanobacteria on nutrient plates based on a 4-unit scale with 25°C diurnal controls defining a score of 4.

The colony-forming ability of the two heterotrophic bacteria was assayed directly on the basis of colony counts on nutrient agar. The results are shown in Figure 23 and indicate the survival of reproducing cells under all treatment conditions.
Figure 23. Counts of cfu/mL of two species of heterotrophic eubacteria. 

Chlorophyll was extracted from samples of the four autotrophs and measured spectrophotometrically. Figure 24 indicates that chlorophyll could be detected in all specimens under all treatments, and Mars simulation was found less destructive of chlorophyll than the other two test treatments.

Figure 24. Chlorophyll concentrations in extracts of phototrophs after treatments.
Nitrite tests were performed to determine remaining functions for denitrification and converting nitrate to nitrite. Table 3 summarizes the results of nitrite tests (vs. optical standards) for all six species tested. Only the heterotrophic eubacteria retained denitrification ability after exposure to Mars conditions.

Table 4. Nitrate reduction and denitrification by the tested microbial species.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Mars simulator</th>
<th>-80°C Dark control</th>
<th>4°C Dark control</th>
<th>25°C Diurnal control</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Anabaena</em> sp.</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>NO₂</td>
</tr>
<tr>
<td><em>Chlorella ellipsoidea</em></td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>NO₂</td>
</tr>
<tr>
<td><em>Chroococcidiopsis</em></td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>NO₂</td>
</tr>
<tr>
<td>CCMEE171</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>NO₂</td>
</tr>
<tr>
<td><em>Plectonema boryanum</em></td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>NO₂</td>
</tr>
<tr>
<td><em>Bacillus subtilis</em></td>
<td>N₂</td>
<td>None</td>
<td>N₂</td>
<td>N₂</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>N₂</td>
<td>NO₂</td>
<td>N₂</td>
<td>N₂</td>
</tr>
</tbody>
</table>

Images of the streaked plates all showed evidence of surviving reproducing cells after exposure to the simulated Mars conditions. Figures 25 and 26, respectively, consist of images of plates streaked with phototrophic and heterotrophic cells from the Mars exposure (bottom row) compared with plates streaked with cells from cultures kept at 25°C with diurnal lighting (top row).

Figure 25. Images of plates streaked with phototrophic cells from the Mars exposure (bottom row) compared with plates streaked with cells from cultures kept at 25°C with diurnal lighting (top row).
Objective 3. Create a mechanical and electronic design concept for Mars surface shallow penetrator with planetary protection.

Proposed details of this task are seen in Figure 1. The advisory team held substantial discussions on how we would seal the bottom of the testbed after boring it into the Martian regolith as well as understanding the size scale. Since the organisms of choice are photosynthetic, we anticipate depths of no more than 3 cm are required so an overall depth of 15 cm was considered adequate. Requirements are clear that it must prevent bacterial escape but allow the hollow shaft to be filled with Martian regolith prior to sealing. Sealing concepts ranged from a chemical and civil engineering approach using regolith-based composite or concrete to mechanical approaches that lock the system closed after penetration and filling. The Techshot mechanical design team took this on and produced a number of CAD concepts. A design was selected, and 3-D printing of prototypes of all parts was undertaken in order to assemble a partially functional mock-up of the penetrator at an enlarged scale.
Selection of penetrator design

Selection of final components:
A ceramic filter that allows atmosphere into, but not organisms out from, the testbed was originally proposed. However, filter material with 0.5 µm pore size is also available in PEEK (polyetheretherketone) and stainless steel, and these materials are preferred because they are less frangible than porous ceramic filters.

Identification of a miniaturized biosensor approach was the goal of Objective 4 study, which follows.

Objective 4. Identify electronic biological activity tests (O₂ sensor, for example); initiate testing in a laboratory simulator.

Lengthy discussions on sensor design and function to “detect” biological activity. Simply measuring O₂ may not be adequate so further discussions are scheduled to look into biomarkers that may serve as surrogates for O₂ production or an optical method of adding Fluorescein to detect the presence of liquid water enabling the production of O₂. It is the goal of the Phase 1 project to identify at least 1 method to move forward into phase 2. Further discussions led toward more versatile sensing of volatile biomarkers and the use of spectrophotometry, gas chromatography or mass spectrometry (MS). The molecular mass of a compound is an almost completely unambiguous identifier, and the best hand-held technology for adaptation for bioproduct sensing in the Mars Ecopoiesis Test Bed is micromanufactured MS. This is not current off-the-shelf technology but will be the technology of the time frame in which launching and operating the Mars Ecopoiesis Test Bed are expected to occur. The development of an integrated hand-held micromanufactured MS system, including sampling [Hill et al., 2014], ionization source [Dong et al, 2015; Chen et al., 2011], quadrupole separator, event detector and even an on-chip
vacuum pump are presently becoming reality. Therefore Techshot will collaborate, to the extent possible, with developers of microminiaturized MS on a parallel track to adapt this new and promising technology to bioproduct sensing on Mars.

Publications

http://www.nasa.gov/feature/planting-an-ecosystem-on-mars

References

The following references formed the basis for the project’s concept and research, but not all are cited in this report.


