Spectral features of biogenic calcium carbonates and implications for astrobiology

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Abstract: The ability to discriminate biogenic from abiotic calcium carbonate (CaCO3) would be useful in the search for extant or extinct life, since CaCO3 can be produced by both biotic and abiotic processes on Earth. Bioprecipitated CaCO3 material was produced during the growth of heterotrophic microbial isolates on medium enriched with calcium acetate or calcium citrate. These biologically produced CaCO3, along with natural and synthetic non-biologically produced CaCO3 samples, were analysed by reflectance spectroscopy (0.35–2.5 μm), Raman spectroscopy (532 and 785 nm), and laser-induced fluorescence spectroscopy (365 and 405 nm excitation). Optimal instruments for the discrimination of biogenic from abiotic CaCO3 were determined to be reflectance spectroscopy, and laser-induced fluorescence spectroscopy. Multiple absorption features in the visible light region occurred in reflectance spectra for most biogenic CaCO3 samples, which are likely due to organic pigments. Multiple fluorescence peaks occurred in emission spectra (405 nm excitation) of biogenic CaCO3 samples, which also are best attributed to the presence of organic compounds; however, further analyses must be performed in order to better determine the cause of these features to establish criteria for confirming the origin of a given CaCO3 sample. Raman spectroscopy was not useful for discrimination since any potential Raman peaks in spectra of biogenic carbonates collected by both the 532 and 785 nm lasers were overwhelmed by fluorescence. However, this also suggests that biogenic carbonates may be identified by the presence of this organic-associated fluorescence. No reliable spectroscopic differences in terms of parameters such as positions or widths of carbonate-associated absorption bands were found between the biogenic and abiogenic carbonate samples. These results indicate that the presence or absence of organic matter intimately associated with carbonate minerals is the only potentially useful spectral discriminator for the techniques that were examined, and that multiple spectroscopic techniques are capable of detecting the presence of associated organic materials. However, the presence or absence of intimately associated organic matter is not, in itself, an indicator of biogenicity.

Key words: biomarkers, biomineralization, carbonates, exobiology, Mars, spectroscopy.

Introduction

Current and upcoming Mars exploration missions are targeted towards habitability, and the detection of extant or extinct life through in situ exploration and future sample return missions. Currently, Mars is a cold desert considered to be inhospitable to life; it is postulated that if life were to exist on Mars, its presence could be detected through biomarkers indicating past life or biological activity (Banfield et al. 2001). There has been evidence for the presence of liquid water on the surface and subsurface of early Mars, which would have been conducive to the development and support of life (McKay 1997). It has been hypothesized that geological and environmental conditions of Early Mars were similar to those of Early Earth, in which conditions were warm and wet (Davis & McKay et al. 1996; McKay 1997; Cockell et al. 2000; Holm & Andersson 2005; Edwards 2010). It has been suggested that if life originated on Mars during this time, it may have been similar to microorganisms living in extreme environments on Earth today (McKay 1997; Raulin & McKay 2002; Simoneit 2004; Delaye & Lazcano 2005; Edwards 2010).

Recently, carbonates have been detected at several sites on Mars, such as in the Nili Fossae region (Ehlmann et al. 2008), in the Columbia Hills of Gusev crater (Morris et al. 2010) and in Leighton crater (Michalski & Niles 2010). Carbonates have also been detected in Martian meteorite ALH84001, and have been thought to be of biological origin (McKay et al. 1996), however not without dispute (e.g. Vecht & Ireland 2000). Of particular interest to our study, calcium carbonate has tentatively been detected in soils at the Phoenix Landing Site (Boynton et al. 2009).

On Earth, calcium carbonate (CaCO3) can be formed by biomineralization, diageneric processes, or abiogenic processes (Tucker & Wright 1990; Stalport et al. 2005). CaCO3 is one of the most commonly produced biominerals, and it can be produced under strict genetic control (biologically controlled
mineralization), as seen in eukaryotes, or in the form of a by-product (biologically induced mineralization), as seen in prokaryotes (Mann 2001). Six CaCO₃ polymorphs exist, in which calcite and aragonite are the most stable and most commonly formed by biominalization (Mann 2001; Ehrlich 2002).

There are numerous minerals that can be produced by both biogenic and abiogenic processes on Earth, which highlights the need for determining a method to differentiate between them. Factors that may indicate biogenicity of a mineral include, but are not limited to: specific phases, major element composition, trace elements, isotopic composition, crystal morphology, surface composition and morphology, spatial arrangement, and the presence of organic molecules indicating biominalization (Banfield et al. 2001). In order to determine the biogenicity of a sample from Mars, it would first be necessary to establish reliable criteria that differentiate minerals that are biologically produced from their non-biologically produced counterparts. Similarly, the determination of an optimal instrument able to detect these differences and to be included on future missions involving rovers or landers would be extremely useful in the detection of biomarkers in planetary exploration.

Some work has been done on characterizing differences between biogenic and abiogenic CaCO₃, however much of the work has focused on differential thermal analysis (Stalport et al. 2005, 2008), and analysis of thermally processed carbonates using infrared spectroscopy (Orofino et al. 2007, 2009; Blanco et al. 2013). Our study aims to establish discrimination criteria using samples that have not been thermally altered in order to provide a baseline for future studies. Instead, spectroscopic techniques were used since they allow for the non-destructive analysis of samples without sample preparation, facilitating in situ analyses. The instruments that were applied to the analysis of biogenic and abiogenic samples include some that are available on orbiting platforms, such as reflectance spectrometers (Bibring et al. 2005; Murchie et al. 2007), and others that have been proposed for future landers/rovers, including reflectance spectroscopy, Raman and laser-induced fluorescence (Storrie-Lombardi & Sattler 2009; Piorget et al. 2012; Korables et al. 2013; De Angelis et al. 2014). A companion study is evaluating how and if any spectral differences are altered after exposure to simulated Martian conditions.

In order to investigate potential criteria and optimal instruments for the discrimination of biogenic from abiogenic CaCO₃, we analysed bioprecipitated material collected from microbial isolates taken from Mars analogue sites within the Canadian High Arctic (Steven et al. 2007) and Antarctica (Goordial personal communication). In this study, biologically produced and non-biologically produced CaCO₃ were analysed by reflectance spectroscopy, Raman spectroscopy, and laser-induced fluorescence spectroscopy. The spectral differences between biogenic and abiogenic carbonates will be useful in evaluating how these minerals could be distinguished by a planetary rover or lander in the search for extant or extinct life.

Materials and methods

Biological precipitation of calcium carbonates

Each biogenic CaCO₃ sample was precipitated during the growth of several heterotrophic bacteria isolated from Polar soil and endolith samples, as outlined in Ronholm et al. (in press). A culture collection (~ 300 isolates) was screened for the ability to precipitate CaCO₃ crystals. B₄ Media (Boquet et al. 1973) (4 g of yeast extract, 10 g glucose, 15 g bacteriological agar and either 2.5 g calcium acetate or 2.5 g calcium citrate per litre of media) was streaked with each isolates and grown for 4 weeks in a humidity controlled environmental chamber at 22 °C. After 4 weeks each isolate was inspected for visual signs of CaCO₃ precipitation on both calcium acetate and calcium citrate supplemented media. Several, but not all, isolates produced macroscopic crystals on one or both media types (Table 1). Negative control B₄ plates were incubated under the same conditions without being inoculated with bacterial cultures; crystals did not form on these plates.

Biogenic CaCO₃ was separated from growth media and bacteria by scraping the mineral/bacteria mixture off the surface of solid media using a sterile inoculation loop. Biogenic CaCO₃ was separated from the bacteria by successive rounds of sedimentation in millipure water (Sánchez-Román et al. 2011). Precipitates were then dried on the low setting using a rotary evaporator and stored at room temperature until examination. The abiogenic carbonates included for comparison were naturally occurring (likely hydrothermal) monocristalline or coarsely-cristalline aggregates of aragonite (CRB120 and CRB125), calcite (CRB131) and a synthetic calcite (PIG004). Descriptions of these samples and the biogenic samples are provided in Table 2.

X-ray diffractometry

The identities of the biogenic and abiogenic samples included in this study were confirmed by X-ray diffractometry (XRD), and shown in Tables 1 and 2. XRD analysis was carried out using a Bruker D8 Discover diffractometer at the University of Winnipeg using Co Kα₂,2 radiation (λ = 1.78897 Å), accelerating voltage 40 kV and 40 mA beam current. X-ray diffraction patterns were collected with a step size 0.02° and counting time of 10 s per step covering 10–70°20.

Reflectance spectroscopy

Reflectance spectra over the 0.35–2.5 μm range were acquired with an Analytical Spectral Devices (ASD) FieldSpec Pro high-resolution (HR)® spectrophotometer. The spectral resolution of the instrument varies between 2 and 7 nm and spectral sampling is done at 1.4 nm intervals. The instrument internally resamples the data to 1 nm intervals with a third-order polynomial. Spectra were acquired at ι = 30° and ε = 0° using the FieldSpec Pro. Illumination was provided by an in-house 100 W quartz-tungsten-halogen (QTH) bulb directed into a fibre assembly through an aluminium light pipe. A total of 500 spectra were averaged to improve the signal-to-noise ratio (SNR). Reflectance spectra were measured relative to a calibrated Spectralon® disk. Wavelength calibration was
<table>
<thead>
<tr>
<th>Sample Description</th>
<th>Source of Microbial Isolate</th>
<th>Optimal Substrate for CaCO₃ Precipitation</th>
<th>CaCO₃ Polymorphs Identified by µXRD</th>
<th>Major Mineral Morphologies Identified (SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arthrobacter sp.</td>
<td>Antarctic Soil</td>
<td>Calcium Acetate (CA)</td>
<td>Calcite</td>
<td>Euhedral</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Chloride (CC)</td>
<td>Calcite</td>
<td>Euhedral</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Nitrate (CN)</td>
<td>Calcite</td>
<td>Anhedral (Diskoid)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Sulfate (CS)</td>
<td>Calcite</td>
<td>Anhedral (Flakey)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Sulfate (CS)</td>
<td>Calcite</td>
<td>Euhedral, Anhedral (Globular, Hemi-spherulite, diskoid)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Acetate (CA)</td>
<td>Calcite</td>
<td>Euhedral</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
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<td>Calcium Chloride (CC)</td>
<td>Calcite</td>
<td>Anhedral (Flakey)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Arctic Soil</td>
<td>Calcium Chloride (CC)</td>
<td>Calcite</td>
<td>Anhedral (Irregular, Flakey)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
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<td>Calcium Acetate (CA)</td>
<td>Calcite</td>
<td>Euhedral</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Antarctic Soil</td>
<td>Calcium Chloride (CC)</td>
<td>Calcite</td>
<td>Anhedral (Flakey)</td>
</tr>
<tr>
<td>Arthrobacter sp.</td>
<td>Antarctic Soil</td>
<td>Calcium Chloride (CC)</td>
<td>Calcite</td>
<td>Anhedral (Spherulite)</td>
</tr>
<tr>
<td>Rhodococcus sp.</td>
<td>Antarctic Soil</td>
<td>Calcium Acetate (CA)</td>
<td>Calcite</td>
<td>Anhedral</td>
</tr>
</tbody>
</table>

**Raman spectroscopy**

Raman spectra were collected at the University of Winnipeg over the 175–4000 cm⁻¹ range at a resolution of ~4 cm⁻¹ at 614 nm with the B&W Tek (Newark, DE) i-Raman-532-S instrument. Excitation was provided by a 532 nm ~50 mW BWN solid-state diode laser. Raman-scattered light was detected by a Glacier™ T, a high spectral resolution (0.08 nm) thermoelectrically cooled (14°C) charge-coupled device (CCD) detector. The automatic integration time function (which increases integration time incrementally, until the response is close to saturation) was used, yielding an optimal SNR. Initial sample measurements were acquired with the BAC102 fibre optic attachment with a working distance of 5.9 mm that gives a theoretical spot size of 85 μm. The BAC102 fibre optic probe contains a long-pass OD6 notch filter, which filters reflected excitation light prior to interaction with the detector for an increased SNR. Samples were placed on an adjustable stage and were approached directly to the distance regulator prior to collection. Measurements for each sample were made by first acquiring a dark current spectrum, followed by measurement of the sample. Both measurements were made using an identical viewing geometry and integration time, and 10 averaged spectra. Raman-shift calibration was monitored through regular measurements of a polystyrene standard.

A DeltaNu Rockhound Raman spectrometer was used for complementary data collection in the 200–2000 cm⁻¹ range with a thermoelectrically cooled (0°C) CCD detector. A solid-state diode laser provided excitation of 100 mW at 785 nm. Theoretical spot size measured by the instrument is 25 μm. The instrument has a long pass OD6 notch filter. A spectral resolution of 7 cm⁻¹, an optimized integration time approaching signal detector saturation, and 10 averaged spectra were used to increase the SNR within each sample. The Rockhound instrument was mounted securely with the probe pointed 90° downwards. Samples were placed on an adjustable stage and were approached directly to the probe prior to collection. Prior to each sample collection, the instrument was calibration checked with a polystyrene standard.

**Laser-induced fluorescence spectroscopy**

Sample fluorescence was measured at the University of Winnipeg in the 200–1150 nm range with an Ocean Optics (Dunedin, Fl) Maya2000 PRO miniaturized spectrometer equipped with an HC-1 grating and 30 μm slit width, giving...
us an effective spectral resolution between 0.48 nm at 200 nm and 0.46 nm at 400 nm. The detector is a 2D back-thinned linear CCD-array Hamamatsu S10420. Illumination was provided by one laser: a 75 mW 405 nm (Wicked Lasers, Tsim Sha Tsui, Kowloon, Hong Kong) and one spot light: UVP (Upland, CA) Black-Ray® B-100AP 100 W (21 700 uW cm⁻² at a distance of two inches) spotlight with a 365 nm bandpass filter, that were independently directed through a bifurcated fibre optic bundle consisting of six illumination fibres surrounding a central pick-up fibre feeding into the detector array. This assembly consisted of 400 μm diameter solarization-resistant (XSR) fibres. A viewing geometry of \( \theta = 0° \) and \( \phi = 0° \) was used for all collections. An integration time of 150 ms with 500 averaged spectra were used to determine sample fluorescence at 365 nm. Fluorescence spectra were collected and interpreted in conjunction with the obtained Raman spectra, as discussed below.

### Results and discussion

Our findings suggest that there are differences in spectral features between biogenic and abiogenic carbonates using a number of the applied spectroscopic techniques. There was no apparent correlation between medium type on which a given bacterial isolate was grown and CaCO₃-associated absorption band positions in reflectance, or emission bands in Raman or fluorescence spectra of their bioprecipitated material. Also, the genus of bacterial isolate responsible for CaCO₃ precipitation did not predict which spectral features are present. It should be cautioned that the attribution of a given peak to a given pigment or compound is not always definitive, and further analyses are planned to confirm the presence of specific compounds. The heterotrophic bacteria whose bioprecipitated material was analysed in this study are not known to carry out photosynthesis, therefore the spectral features observed cannot be attributed to photosynthetic pigments, such as chlorophylls, bacteriochlorophylls, or phycobilins.

### Reflectance spectroscopy

All abiotic and biogenic CaCO₃ samples showed an absorption feature at \( \sim 2.34 \mu m \), which can be attributed to the second overtone of the asymmetric stretching mode (3ν₃) of the carbonate ion (Table 3, Fig. 1(a)–(c)) (Gaffey 1987; Workman & Weyer 2012). Band depths for the 2.3 μm feature were similar for both biogenic and abiogenic carbonates. The band minimum wavelength for this feature varied slightly between samples, with the largest deviation seen between abiotic aragonite (CRB120 and 125) and calcite (CRB131, PIG004), as expected (Gaffey 1987).

All biogenic samples and abiogenic aragonite sample CRB125 showed absorption features at both \( \sim 1.4 \) and \( \sim 1.9 \mu m \), which were attributed to the presence of water within these samples (Fig. 1(a)–(c)) (Clark et al. 1990). It should be noted that this water is likely adsorbed, as pure CaCO₃ is anhydrous; therefore, these bands are not diagnostic of anhydrous carbonates.

Below 0.7 μm, biogenic carbonates showed a decrease in absolute reflectance and an increase in number of absorption features, except in *Rhodotorula mucilaginosa* (JG1), both features which were not seen in the abiogenic carbonates (e.g. Fig. 1(d)). Absorption features in the visible spectral range were attributed to the presence of pigmented compounds within the samples, possibly in trace amounts. All biogenic carbonates showed absorption features in the \( \sim 0.406–0.417 \mu m \) range. All biogenic carbonates except *R. mucilaginosa* (JG1), *Arthrobacter* sp. (oc825) and *Rhodococcus* sp. (eur3 9.01.3) showed absorption features in either or both of the \( \sim 0.444–0.459 \mu m \) and \( \sim 0.472–0.477 \mu m \) ranges. All biogenic carbonates except *R. mucilaginosa* (JG1) and *Arthrobacter* (ofe 10.10) had absorption features in either or both the 0.537–0.541 μm and 0.571–0.576 μm ranges. Some biogenic carbonates (*Ralstonia* sp. (ofe 1.3), *Arthrobacter* sp. (oc825), *Arthrobacter* sp. (cy7), *Arthrobacter* sp. (JG9), *Rhodococcus* (cy5), *Rhodococcus* sp. (eur3 9.01.3) and *Rhodococcus* sp. (JG12)) also showed absorption features in the \( \sim 0.498–0.503 \mu m \) region.

Absorption features within the range of 0.420–0.550 μm may be due to the presence of carotenoids, which are isoprenoid pigments that absorb strongly in the blue-green region (0.420–0.550 μm), which could lead to the appearance of yellow, orange or red coloration (Armstrong & Hearst 1996; Dworkin & Falkow 2006). The number of conjugated double bonds in the polyene chain of a given carotenoid plays an important role in the determination of its spectral properties (Armstrong & Hearst 1996). Although carotenoids are often associated with photosynthetic organisms, it has been shown that heterotrophic bacteria are capable of producing carotenoids for protection against environmental stress, such as cold temperatures and solar radiation (Dier et al. 2010), as seen in the hostile environments from which isolates were sampled.
Table 3. Band positions in reflectance spectra (2.3 μm, visible region, and near-infrared band minima), Raman spectra (ν1 symmetric stretch maxima and fluorescence peaks), and laser-induced fluorescence spectra for abiogenic and biogenic samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Reflectance: 2.3 μm band minima</th>
<th>Reflectance: visible absorption band minima (μm)</th>
<th>Raman: ν1 symmetric stretch maxima (cm⁻¹)</th>
<th>Raman: fluorescence peaks (nm)a</th>
<th>Laser-induced fluorescence: peak positions (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRB120 (aragonite)</td>
<td>2.32</td>
<td>None</td>
<td>1088</td>
<td>~ 594b</td>
<td>None</td>
</tr>
<tr>
<td>PIG004 (synthetic calcite)</td>
<td>2.34</td>
<td>None</td>
<td>1085</td>
<td>~ 596b and ~ 618b</td>
<td>None</td>
</tr>
<tr>
<td>CRB131 (calcite)</td>
<td>2.34</td>
<td>None</td>
<td>1085</td>
<td>~ 596b and ~ 618b</td>
<td>None</td>
</tr>
<tr>
<td>CRB125 (aragonite)</td>
<td>2.32</td>
<td>None</td>
<td>1085</td>
<td>~ 596b</td>
<td>None</td>
</tr>
<tr>
<td>Arthrobacter sp. JG9</td>
<td>2.34</td>
<td>0.411d, 0.459i, 0.500i, 0.538i and 0.574i</td>
<td>1085f</td>
<td>~ 596b and ~ 618b</td>
<td>576b, 620b and 682b</td>
</tr>
<tr>
<td>Arthrobacter sp. ocf825</td>
<td>2.34</td>
<td>0.411d, 0.499d, 0.539d and 0.574d</td>
<td>1085f</td>
<td>~ 596b and ~ 618b</td>
<td>503b, 620b and 682b</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 4.2</td>
<td>2.34</td>
<td>0.417d, 0.446d, 0.477d, 0.539d and 0.573d</td>
<td>1092f</td>
<td>~ 596b</td>
<td>466b and 520b,d</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 7.23</td>
<td>2.34</td>
<td>0.413, 0.459i, 0.537 and 0.573</td>
<td>1081f</td>
<td>~ 596b and 622</td>
<td>None</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 10.25</td>
<td>2.34</td>
<td>0.414, 0.451d, 0.474, 0.541 and 0.576</td>
<td>1075f</td>
<td>~ 582 and ~ 630b</td>
<td>466b,d, 520b,d, 578b, 620 and 682f</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 1.1</td>
<td>2.34</td>
<td>0.410, 0.450d, 0.475d, 0.503, 0.540 and 0.574</td>
<td>1082f</td>
<td>~ 580b and ~ 620</td>
<td>466b, 498b, 578b, 620 and 682f</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 10.10</td>
<td>2.34</td>
<td>0.415d, 0.444d and 0.474d</td>
<td>1085f</td>
<td>~ 596b</td>
<td>466b,e, 520b,e, 578d, 620 and 682f</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 3.1</td>
<td>2.34</td>
<td>0.411, 0.444, 0.476, 0.537 and 0.571</td>
<td>1085f</td>
<td>~ 596b and ~ 618</td>
<td>466, 498b, 580b,d, 620 and 682d</td>
</tr>
<tr>
<td>Arthrobacter sp. ofe 8.9</td>
<td>2.34</td>
<td>0.410, 0.445, 0.477, 0.537 and 0.571</td>
<td>1075f</td>
<td>~ 596b and ~ 616</td>
<td>466, 498b, 580b,d, 620 and 682d</td>
</tr>
<tr>
<td>Arthrobacter sp. eur3 7.19</td>
<td>2.34</td>
<td>0.410, 0.444, 0.476, 0.537 and 0.571</td>
<td>1085f</td>
<td>~ 596b and ~ 618</td>
<td>466, 498b, 580b,d, 620 and 682d</td>
</tr>
<tr>
<td>Arthrobacter sp. cy7</td>
<td>2.34</td>
<td>0.410, 0.459i, 0.501d, 0.539 and 0.574</td>
<td>1085f</td>
<td>~ 596b and ~ 618</td>
<td>466b,e and 520b,e</td>
</tr>
<tr>
<td>Ralstonia sp. ofe 1.3</td>
<td>2.34</td>
<td>0.410, 0.477d, 0.503d, 0.540d and 0.574d</td>
<td>1085f</td>
<td>~ 596b</td>
<td>503b, 620b and 682d</td>
</tr>
<tr>
<td>Rhodococcus sp. JG12</td>
<td>2.34</td>
<td>0.411d, 0.459i, 0.500i, 0.538i and 0.574i</td>
<td>1085f</td>
<td>~ 585b, ~ 596i and ~ 618b</td>
<td>466b, 520b, 578b, 620 and 682f</td>
</tr>
<tr>
<td>Rhodococcus sp. ofe 9.29</td>
<td>2.34</td>
<td>0.412, 0.447, 0.472, 0.538 and 0.574</td>
<td>1075f</td>
<td>~ 608b,g</td>
<td>466b, 498b,d, 580 and 682d</td>
</tr>
<tr>
<td>Rhodococcus sp. cy5</td>
<td>2.34</td>
<td>0.411d, 0.459i, 0.500i, 0.538i and 0.574i</td>
<td>1085f</td>
<td>~ 596b</td>
<td>503b, 620b and 682d</td>
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<tr>
<td>Rhodococcus sp. eur3 9.01.3</td>
<td>2.34</td>
<td>0.406d, 0.498i, 0.538i and 0.572i</td>
<td>1085f</td>
<td>~ 596b</td>
<td>503b, 620b and 682d</td>
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<tr>
<td>Rhodotorula mucilaginosa JG1 (yeast)</td>
<td>2.34</td>
<td>None</td>
<td>1088</td>
<td>~ 596b</td>
<td>503b, 620 and 682</td>
</tr>
</tbody>
</table>

a Raman spectra plotted in wavelength rather than inverse centimetres.
b Maximum of a broad band.
c Minor/relevant fluorescence.
d Weak.
e Very weak.
f Inadequate signal above fluorescence to assign a reliable position.
g No clear maximum.

Note: 2.3 μm region absorption bands are reported to the nearest 10 nm. Visible region absorption bands are reported to the nearest 1 nm.
However, the reflectance spectrum for minerals formed by *R. mucilaginosa* (JG1), a strain of yeast that normally produces carotenoids and appears pink during growth (Moore *et al.* 1989), did not show absorption features in the visible region. One possible explanation for this exception is that the concentration of carotenoids was too low to result in measurable absorption features.

Absorption features between 0.400 and 0.630 µm may have also been due to the presence of cytochromes, which are pigments containing an iron-containing porphyrin (haem) prosthetic group involved in cellular respiration and electron transfer (Thöny-Meyer 1997). In reduced cytochromes, there are three strong absorption features in the ultraviolet to visible spectrum called the $\alpha$, $\beta$ and $\gamma$, or Soret, peaks.
Fig. 1. (a) Reflectance spectra (0.35–2.5 µm) of abiogenic aragonite (CRB120 and CRB125) and calcite (CRB131 and PIG004). (b) Reflectance spectra (0.35–2.5 µm) of abiogenic aragonite (CRB120 and CRB125), abiogenic calcite (CRB131 and PIG004) and various biogenic CaCO₃ samples: ofe 7.23, eur3 7.19, ofe 8.9, ofe 3.1, ofe 10.25, ofe 1.1, oc825 and JG9. Absorption bands in the 1.4 and 1.9 µm regions are likely due to adsorbed water and are not diagnostic of anhydrous carbonates. Characteristic carbonate absorption features are present near 2.3 microns. Absorption bands attributable to organic compounds are present in the visible spectral region (narrow absorption bands below ~0.7 µm, as well as in the 1.7 and 2.0–2.2 µm regions, due to various organic (C–H) molecules (e.g. Izawa et al. 2014). (c) Reflectance spectra (0.35–2.5 µm) of abiogenic aragonite (CRB120 and CRB125), abiogenic calcite (CRB131 and PIG004) and various biogenic CaCO₃ samples: ofe 4.2, ofe 10.10, JG1, ofe 9.29, ofe 1.3, cy7, cy5, eur3 9.01.3 and JG12. (d) Reflectance spectra (0.35–2.5 µm) of abiogenic aragonite (CRB120 and CRB125), abiogenic calcite (CRB131 and PIG004) and biogenic CaCO₃ (represented by sample ofe 1.1). Multiple absorption features in the visible light region, and a decrease in absolute reflectance are present for biogenic CaCO₃, but are absent in abiogenic samples.
Different spectral features exist for different types of cytochromes, in which the α peak (0.550–0.630 μm) can be used to classify the type of cytochrome present in a sample (Thöny-Meyer 1997). Oxidized cytochromes absorb in the 0.390–0.420 μm range (Soret region), and may also be related to the observed absorption features in the studied biogenic carbonates (Barton 2005).

An alternative explanation for the absorption of light in the visible and near-infrared region could be due to colour centres within the lattice structure of the bioprecipitated carbonates. Colour centres are caused by an unpaired electron, in which there can be an extra electron at a vacancy, a missing electron where there is usually a pair of electrons, or an unpaired electron can move to an excited state upon light absorption (Schulman & Compton 1962; Nassau 1978). Crystal lattices of

Fig. 2. For legend see next page.
Fig. 2. (a) Raman spectra (175–4000 cm$^{-1}$) of abiogenic aragonite (CRB120 and CRB125) and calcite (PIG004 and CRB131) collected with a 532 nm laser. (b) Raman spectra (175–4000 cm$^{-1}$) of abiogenic calcite (PIG004) and biogenic CaCO$_3$ (represented by sample ofe 1.1) collected with a 532 nm laser. (c) Raman spectra (175–4000 cm$^{-1}$) of abiogenic aragonite (CRB120 and CRB125) and calcite (PIG004 and CRB131) collected with a 785 nm laser. (d) Raman spectra (175–4000 cm$^{-1}$) of abiogenic calcite (PIG004) and biogenic CaCO$_3$ (represented by sample F4B3) collected with a 785 nm laser.
minerals have defects that occur naturally, and include vacancies, interstitials, and substituted impurities (Schulman & Compton 1962). However, colour centres usually result in broad and weak absorption features (Nassau 1978).

The presence of various organic molecules is also evident from absorption features seen in the biogenic sample spectra in the 1.7 and 2.0–2.2 μm regions in particular. These are generally attributable to various organic molecules, specifically C–H overtone and combination bands, although other compounds such as N-bearing molecules may also contribute to absorptions in these regions (e.g. Izawa et al. 2014).

**Raman spectroscopy**

Raman spectra of abiogenic CaCO₃ samples with the 532 nm laser showed the presence of Raman peaks, whereas any potentially present Raman peaks were overwhelmed by sample fluorescence in spectra for biogenic carbonates (Table 3, Fig. 2(a) and (b)). Raman spectra for all abiogenic CaCO₃ had a major peak at ~1088 cm⁻¹, due to the symmetric stretching mode (ν₁) of the carbonate ion (White 2009). Additionally, spectra for abiogenic aragonite samples showed minor peaks at ~208 and ~702 cm⁻¹, while spectra for abiogenic calcite samples showed minor peaks at ~281 and ~712 cm⁻¹ (Fig. 2(a) and (b)). The peaks in the 200 cm⁻¹ range are attributed to lattice modes, and those in the 700 cm⁻¹ range were attributed to the in-plane bending mode (ν₄) of the carbonate ion (Bischoff et al. 1985; Urmos et al. 1991; Stopar et al. 2005; White 2009).

Raman spectra collected with the 785 nm laser exhibited a peak between 1075 and 1092 cm⁻¹ for both biogenic and abiogenic CaCO₃ (Table 3, Fig. 2(c) and (d)). These results indicated that there is reduced sample fluorescence when using a longer excitation wavelength in Raman measurements, which is ideal in the characterization of minerals containing organic matter; however, additional fluorescence suppression is still needed to obtain peaks for any organic compounds present. The selection of an optimal laser excitation wavelength is an important factor in Raman spectroscopy since different compounds have different sensitivities to a given laser wavelength, and wavelength can affect the level of sample fluorescence emission to the extent in which higher intensity fluorescence peaks can overwhelm the weaker Raman bands (Villar & Edwards 2006). The use of an even longer wavelength (>785 nm) may be successful in reducing sample fluorescence to observe Raman peaks in spectra of these particular biogenic CaCO₃ samples.

**Laser-induced fluorescence spectroscopy**

Abiogenic carbonates did not show any laser-induced fluorescence using the 365 and 405 nm lasers, whereas all biogenic carbonates except *Arthrobacter* sp. (ofe 7.23) exhibited fluorescence (Table 3, Fig. 3(a) and (b)). All biogenic samples except *Arthrobacter* sp. (ofe 7.23, ofe 4.2, ofe 9.29 and cy7) showed fluorescence peaks at ~620 and ~682 nm. *Arthrobacter* sp. (ofe 10.10, ofe 8.9, ofe 3.1, ofe 9.29, ofe 1.1, JG9 and ofe 10.25), *Agromyces* sp. (eur3 7.19), and *Rhodococcus* sp. (JG12) showed fluorescence peaks at either ~576, ~578 or ~580 nm. *Agromyces* sp. (eur3 7.19), *Arthrobacter* sp. (ofe 8.9, ofe 3.1, ofe 9.29 and ofe 1.1), *Ralsenia* sp. (ofe 1.3), *Rhodococcus* sp. (cy5 and eur3 9.01.3), *Arthrobacter* sp. (oc825) and *R. mucilaginosus* (JG1) showed fluorescence peaks at either ~498 or ~503 nm. Many samples (ofe 10.25, ofe 10.10, eur3 7.19, ofe 8.9, ofe 3.1, ofe 10.25, ofe 9.29, ofe 1.1, cy7 and JG12) showed fluorescence peaks at ~466. Some samples (ofe 4.2, ofe 10.10, ofe 10.25, cy7 and JG12) also showed peaks at ~520 nm.

Fluorescence peaks observed in the Raman spectra (532 nm) are listed in Table 3 Raman spectra for abiogenic aragonite carbonate samples (CRB120 and CRB125) showed fluorescence peaks at ~596 nm (2018 cm⁻¹) and ~594 nm (1962 cm⁻¹), respectively, while abiogenic calcite (CRB131) showed a peak at ~611 nm (2430 cm⁻¹) (Fig. 2(a) and (b)). Raman spectra for all biogenic carbonates except ofe 9.29 showed fluorescence peaks at either ~582, ~585 or ~596 nm. Raman spectra for many of the biogenic carbonates (isolates ofe 9.29, ofe 7.23, eur3 7.19, ofe 8.9, ofe 3.1, ofe 10.25, cy7, oc825, JG9 and JG12) had a fluorescence peak at either ~618, ~620, ~622 or ~630 nm.

Fluorescence may be attributed to the presence of trapped electrons, organic compounds, or trace amounts of transition metals in CaCO₃ (Bischoff et al. 1985; Urmos et al. 1991; Wang & Mullins 1997; Bozlee et al. 2005). Organic compounds that emit fluorescence are generally aromatic or conjugated, such as the enzyme cofactors FAD and NADPH, which have emission maxima at ~535 and 440/460 nm, respectively (Ammor 2007). In characterizing biogenic and abiogenic magnesium carbonates with Raman spectroscopy, Bischoff et al. (1985) also noted fluorescence in spectra for biogenic samples that at times overwhelmed Raman peaks, and was attributed to the incorporation of trace metal ions, since they are found in virtually all biogenic materials (Bischoff et al. 1983, 1985; Veizer 1983). This explanation could account for the lack of fluorescence emission peaks in spectra for our abiogenic samples since they should be of higher purity than the biogenic samples. Raman spectra for biogenic samples often show a weak peak at ~1014 cm⁻¹ due to the bicarbonate ion (HCO₃⁻), which may be present in order to balance Na⁺ charges in biological matrices (White 1975; Bischoff et al. 1983, 1985; Veizer 1983).

**Implications for astrobiology**

Our results indicate that reflectance spectroscopy would be able to discriminate biogenic from abiogenic CaCO₃ by numerous absorption features in the visible light region in spectra of biogenic carbonates. However, there are two important caveats: (1) the spectroscopic features that seem to enable discrimination of biogenic from abiogenic carbonates are all associated with the co-existing organic matter in the biogenic samples; and (2) this presumes that the carbonates under investigation are pure and no spectral interferences would be present from any phases that are not of interest for discriminating biogenic from abiogenic carbonates. Also, laser-induced fluorescence spectroscopy could be a useful tool for
detecting biomarkers, since many fluorescence emission peaks occurred in the visible region upon excitation with 365 and 405 nm light sources; however, it depends on what is causing these peaks. If the fluorescence peaks are in fact due to incorporation of trace elements, then it would be difficult to attribute to biogenicity based on fluorescence spectroscopy alone. Therefore, further analyses would be required in order to confirm the specific organic compounds present.

Fig. 3. (a) Laser-induced fluorescence spectra (200–1150 nm) of abiogenic aragonite (CRB120 and CRB125) and calcite (PIG004 and CRB131) collected with a 405 nm light source. (b) Laser-induced fluorescence spectra (200–1150 nm) of abiogenic calcite (PIG004) and biogenic CaCO₃ (represented by sample ofe 1.1) collected with a 405 nm light source. The prominent peak near 405 nm and weaker peak near 810 nm are attributable to the laser.
According to our findings, Raman spectroscopy with a 532 or 785 nm laser may result in sample fluorescence, and overpowering of any Raman peaks for organic material. This could be used as an indicator of the presence of organic molecules which would be consistent with, but not necessarily indicative of, biogenically precipitated carbonates.

One concern in differentiating between biogenic and abiogenic minerals is that a compound having biomimetic properties could lead to the misinterpretation of a mineral as having biological origin. Biomimetic carbonates have been found to be indistinguishable from their biogenic counterparts when analysed by differential thermal analysis, since both exhibit a highly strained crystal structure and similar thermal resistance difference from abiogenic carbonate (Thompson et al. 2014). The increased strain causing a lower phase transformation temperature, seen in both biomimetic and biogenic carbonates, may be due to incorporation of organic molecules (e.g. amino acids) (Thompson et al. 2014). Therefore, it has been suggested that results from differential thermal analysis of carbonates may not necessarily indicate the presence of life; it may only provide evidence for pre-biotic organics that could have been previously used to form life (Thompson et al. 2014). This concern may be extended to spectroscopic discrimination of carbonates, and studies concerning the spectral features of biomimetic carbonates should be carried out in order to detect any similarities between biomimetic and biogenic carbonates.

Biomarkers are variable in stability; therefore the degradation of compounds and their derivatives should be taken into account in the search for extant or extinct life on Mars. On Mars, fossilization and preservation of organic material is enhanced by cold temperatures, and anaerobic, hypersaline and arid conditions (Ellery et al. 2002). Over geological time, carotenoids are degraded to isopenoids via modification of the carotenoid polynye chain by various reactions, with hydrogenation being the most common degradation pathway (Ellery et al. 2002; Marshall & Marshall 2010). Haem redox centres of aerobic metabolism are degraded to porphyrins (Suo et al. 2007). Isopenoids and porphyrins are considered to be important as biomarkers since they have been found to remain in the fossil record for extended periods of time and are indicative of biogenicity (Ellery et al. 2002).

**Future studies**

The degradation of biomarkers should be considered when distinguishing biogenic from abiogenic carbonates, since any remnant of life or biological activity on Mars would be subject to degradation over geological time. We are currently conducting further studies investigating the effects of Martian conditions on the spectral and morphological features of biogenic and abiogenic carbonates. These studies will be useful in identifying if and how harsh conditions can modify mineralogical biomarkers in order to better understand the limits of investigative techniques in the discrimination of biogenic from abiogenic minerals.

**References**


