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# Infrared Spectroscopic Detection of Biosignatures at Lake Tírez, Spain: Implications for Mars

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## Abstract

The detection of potential biosignatures with mineral matrices is part of a multifaceted approach in the search for life on other planetary bodies. The 2020 ExoMars Rosalind Franklin rover includes within its payload three IR spectrometers in the form of ISEM (Infrared Spectrometer for ExoMars), MicrOmega, and Ma-MISS (Mars Multispectral Imager for Subsurface Studies). The use of this technique in the detection and characterization of biosignatures is of great value. Organic materials are often co-deposited in terrestrial evaporites and as such have been proposed as relevant analogs in the search for life on Mars. This study focuses on Ca-sulfates collected from the hypersaline Tírez Lake in Spain. Mid infrared and visible near infrared analysis of soils, salt crusts, and crystals with green and red layering indicative of microbial colonization of the samples was acquired from across the lake and identified the main mineral to be gypsum with inputs of carbonate and silica. Organic functional groups that could be attributed to amides and carboxylic acids were identified as well as chlorophyll; however, due to the strong mineralogical absorptions observed, these were hard to unambiguously discern. Taxonomical assignment demonstrated that the archaeal community within the samples was dominated by the halophilic extremophile Halobacteriaceae while the bacterial community was dominated by the class Nocardiaceae. The results of this research highlight that sulfates on Mars are a mixed blessing, acting as an effective host for organic matter preservation but also a material that masks the presence of organic functional groups when analyzed with spectroscopic tools similar to those due to fly on the 2020 ExoMars rover. A suite of complementary analytical techniques therefore should be used to support the spectral identification of any candidate extraterrestrial biosignatures. Key Words: IR spectroscopy—Mars—Sulfate—Biosignature—Lake Tírez. *Astrobiology* 20, xxx–xxx.

## 1. Introduction

THE SEARCH FOR LIFE on other worlds is a multifaceted endeavor. One of its key elements is the detection of potential biosignatures preserved within mineral structures that could indicate the presence of extinct or extant life. Cell-like morphologies and isotopic signatures resulting from metabolic reactions can also be produced by entirely abiotic mechanisms. The organic components comprising cells and their associated materials can be preserved either as degraded molecules trapped in mineralized cells or as disseminated molecules chemically bonded to mineral particles. Therefore, correlative detection of organic compounds is required before a case for life can be conclusively made.

Historically, the search for life on Mars has focused on the search for water. The presence of mineral phases such as phyllosilicates, sulfates, and opals (Carter *et al.*, 2013, 2015; Sun and Milliken, 2015), tells a story of aqueous alteration processes occurring on the surface and subsurface of early Mars (*e.g.*, Moore and Howard, 2005; Baker, 2006; Ehlmann *et al.*, 2011, Michalski *et al.*, 2013). Now the search includes the detection of organic molecules on or beneath the surface of Mars. Carbon is abundant as an atmospheric gas and can also be found in the form of carbon dioxide ice and carbonate minerals (Ehlmann *et al.*, 2008; Boynton *et al.*, 2009; Morris *et al.*, 2010). In addition, organic matter has now been found preserved within the 3 billion-year-old mudstones of Gale Crater (Eigenbrode *et al.*, 2018). This

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evidence indicates that complex organic molecules have been present and possibly available for life for most of Mars' history, which suggests that this planet has supported multiple habitable environments during the Noachian and early Hesperian periods (Vago *et al.*, 2017).

In astrobiology, minerals provide surfaces to support, concentrate, and preserve organic molecules, and act as templates for prebiotic reactions. This work focuses on sulfates, a class of hydrated minerals detected on Mars. Sulfates have been mapped by the ESA Mars Express OMEGA instrument (Arvidson *et al.*, 2005; Bibring *et al.*, 2005; Gendrin *et al.*, 2005) and the Compact Reconnaissance Imaging Spectrometer for Mars (CRISM) on board Mars Reconnaissance Orbiter (MRO) (Pelkey *et al.*, 2007; Wiseman *et al.*, 2010), and detected by the NASA rovers (Squyres *et al.*, 2004; Gellert *et al.*, 2006; Morris *et al.*, 2006; Ehlmann and Edwards, 2014; McAdam *et al.*, 2014). Varieties identified include iron sulfates (such as jarosite and ferricopiapite), magnesium sulfates (such as kieserite), and calcium sulfates (such as gypsum, basanite, and anhydrite). Among these, Calcium sulfates are more stable under martian conditions compared to other varieties. They have been found in veins and the lower strata of Mount Sharp at Gale Crater (Grotzinger *et al.*, 2012; Nachon *et al.*, 2014; Vaniman *et al.*, 2014; Rapin *et al.*, 2016), at Endeavor Crater, the Columbia Hills at Gusev Crater (Squyres *et al.*, 2006; Yen *et al.*, 2008), and at the North and South Poles of Mars. Sulfates are also not just restricted to ancient terrains; the well-mixed dust that covers most surfaces on Mars has high concentrations of sulfates, at an average concentration of 5.82 wt % (Gellert *et al.*, 2004).

In this paper, we report on the mid infrared (MIR) and near infrared (NIR) spectroscopic study of the hypersaline lake environment, Tírez Lake, in Spain. On Earth, hypersaline environments impose severe stresses on endogenous microorganisms, such as high osmotic pressures and potentially low ( $a_w \sim 0.75$ ) water activities (Grant, 2004). Despite this, life has been found surviving at an unexpected level of diversity over a wide range of salt concentrations (Ley *et al.*, 2006). The potential of hypersaline environments and the evaporitic minerals they precipitate to preserve life and its biomolecules over geologically significant timescales makes Tírez Lake a compelling habitat to investigate biomolecular signatures, and for the search for life vestiges on Earth and beyond.

## 2. Materials and Methods

### 2.1. Tírez Lake

Tírez Lake is located in the province of Toledo, region of Castilla-La Mancha, central Spain, and covers an area of 0.8 km<sup>2</sup>. It is one of several endorheic hypersaline lakes originating under semi-arid conditions in the Iberian Peninsula (Fig. 1a) and is home to a thriving community of sulfate-reducing and methanogenic microorganisms, including haloarchaea and halobacteriaceae (Montoya *et al.*, 2011, 2013). Its salt composition is determined by the inflow of water coming from Triassic evaporites, dolomites, and calcium-sulfate marls of Tertiary age (Prieto-Ballesteros *et al.*, 2003). During the summer, salt content in the water column can reach saturation, causing crystallization of salts such as gypsum, halite, epsomite, and hexahydrate, with high con-

centrations of SO<sub>4</sub><sup>2-</sup>, Cl<sup>-</sup>, Ca<sup>2+</sup>, and Mg<sup>2+</sup> recorded (Prieto-Ballesteros *et al.*, 2003). During these months, the sediments are found to have a circumneutral pH around 6.9 (Fig. 1b).

Samples were collected during the summer of 2017 when dry conditions and high temperatures during the season had caused complete evaporation of the lake. Sampling occurred at regular intervals from the interior of the lake progressively toward the shore. This provided an environmental gradient designed to identify any changes in the mineralogy and potentially the microbiology of Tírez sediments. Four representative areas will be focused on here. Material collected from area 1 (Fig. 1c) is from the interior of the lake and is composed of white and brown sediments, dry crust, and soil from under the dry crust. The samples from areas 2 and 3 (Fig. 1d, 1e) are a mixture of dry salt crusts and mud located at two midway points between the interior and shoreline. Area 4 samples collected from the shore comprise dry crust (Fig. 1f) and whole salt crystals displaying subsurface zones of green and red coloration (Fig. 2). Samples were collected with sterile scoops, placed into sterile falcon tubes and in sterilized sampling bags (Whirl-Pak) and immediately sealed. The samples were kept at ambient temperature during field work, transportation, and storage on return to the laboratory. For analysis, evaporitic samples were ground in an agate pestle and mortar to a fine powder (grains varied in size to reflect the variable grain size distributions on Mars), handled minimally and only with nitrile gloves.

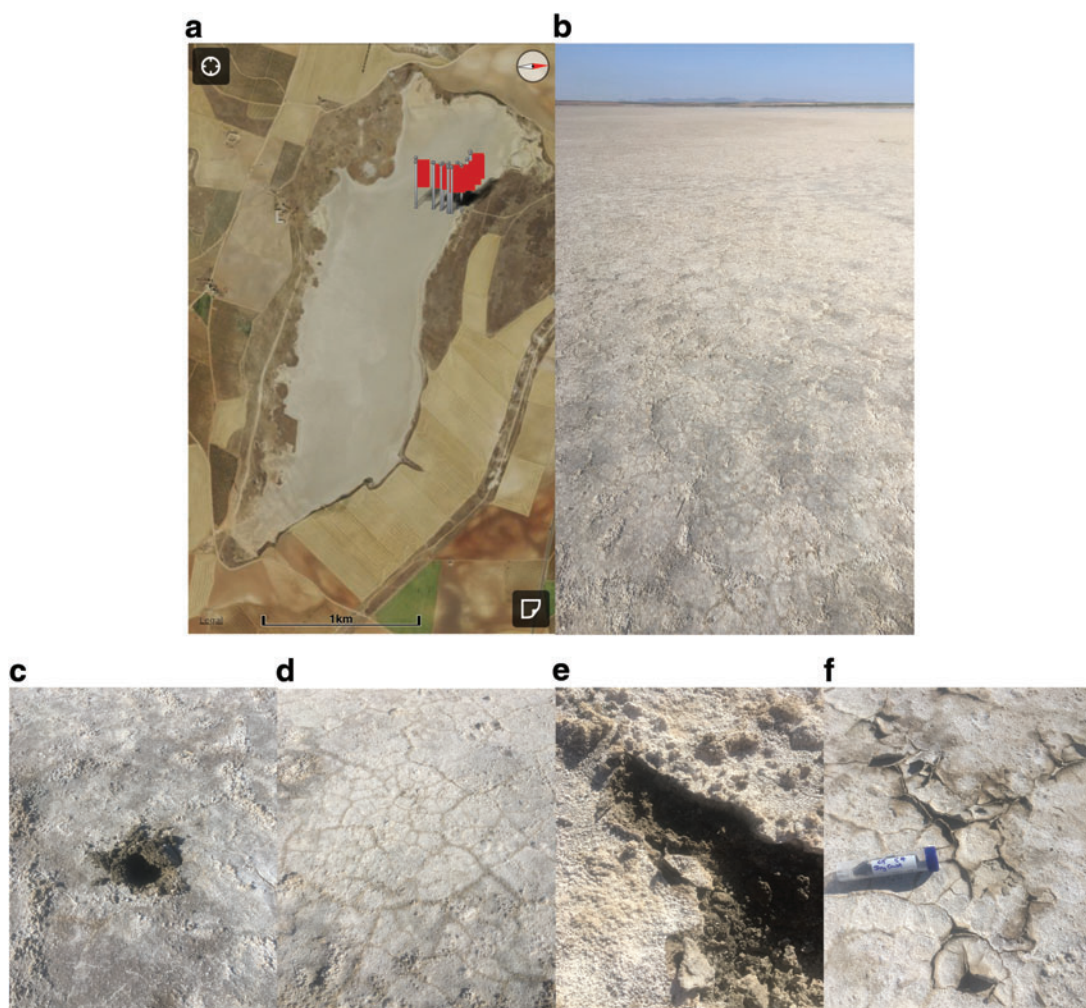
### 2.2. Fourier transform infrared spectroscopy

Fourier transform infrared spectroscopy has been used successfully for the identification of microorganisms and microscopic biological structures (Naumann *et al.*, 1991; Schmitt and Flemming, 1998; Igisu *et al.*, 2009; Preston *et al.*, 2015 and references therein), including analysis of bacterial samples by Helm *et al.* (1991), comparison of Tírez brines with Galileo NIMS spectra data by Prieto-Ballesteros *et al.* (2003), and discrimination of halophilic microorganisms by Güven *et al.* (2015).

Mid infrared spectra were obtained with a Thermo Scientific Nicolet iS5 FTIR Spectrometer and an iD5 Single-Bounce attenuated total reflectance (ATR) attachment equipped with a diamond crystal, housed in the Planetary Spectroscopy Laboratory at Birkbeck, University of London. Reflectance measurements, published as absorbance units, were obtained over a spectral range of 500–4000 cm<sup>-1</sup> at a resolution of 4 cm<sup>-1</sup>. A diamond calibration standard was used and atmospheric water (H<sub>2</sub>O) and carbon dioxide (CO<sub>2</sub>) subtractions made. No further processing of data (*e.g.*, smoothing or Fourier self-deconvolution) was used, to avoid the introduction of artifacts. Band positions are reported simply as the observed maxima, rather than maxima obtained from second derivations or from curve fits. Six analyses were taken of each powdered sample of Tírez sediments which were re-homogenized after each spectral acquisition to obtain an accurate spectral profile of the material and its associated organic functional groups.

### 2.3. Visible near infrared (Vis-NIR) spectroscopy

Near infrared spectroscopy has been used to study the mineralogy of a wide variety of soil profiles and minerals ranging from volcanic glass alteration products and clays



**FIG. 1.** (a) Aerial view over Tírez Lake with area and individual sampling spots labeled by red flags. (b) Photograph on the ground of the sampling area. (c) Site 1 soil sampling. (d) Site 2 salt crust with desiccation patterns visible. (e) Site 3 salt crust and subsurface soil sampling. (f) Site 4 dry salt crusts. Color images are available online.

**FIG. 2.** Magnified image of gypsum crystals and salt crust from site 4 displaying red and green microbial endolithic layers. Color images are available online.



(e.g., Orenberg and Handy, 1992; Viscarra Rossel *et al.*, 2009) to sulfates (e.g., Cloutis *et al.*, 2006). It has also been included in remote sensing payloads to map the mineralogy of planetary surfaces including that of Europa (McCord *et al.*, 1998), Mars (Bibring *et al.*, 2005), and the Moon (Foing *et al.*, 2006).

Three analyses of each unprocessed sample were performed by using an RxSpec 700Z spectrometer housed at the University of Westminster, London. NIR reflectance spectra were obtained over a range of 0.35–2.5  $\mu\text{m}$ .

#### 2.4. Nucleic acid analysis

Prior to extraction, the samples ( $n=3$ ) were rinsed in sterilized MiliQ water to dissolve the outer layer of the salt and then sterilized by using the method described in the work of Jaakkola *et al.* (2014), except the NaCl was replaced with NaSO<sub>4</sub>. The samples were crushed with a sterilized pestle and mortar, and 5 g of sample was added to 30 mL of sterilized MiliQ water. The samples were vortexed for 10 min, and any undissolved material was precipitated by centrifugation, at  $4000\times g$  for 5 min. The supernatant (containing the nucleic acids) was removed and directly added to Amincon Ultra 0.5 mL centrifugal filter units (30 kDA). Twelve filters were used in total to accommodate the volume of supernatant. The filters were centrifuged at  $12,000\times g$  for 5 min at 4°C, inverted, and placed into a clean sterilized centrifuge tube. The nucleic acids were eluted by centrifugation again, as before, and eluted volume was adjusted to 50  $\mu\text{L}$ .

The DNA was extracted by using the XS buffer protocol, as described in Leuko *et al.* (2008) and pooled ( $n=3$ ). The V4 to the V5 of the hypervariable region of the 16S rRNA gene was amplified with universal primers using PCR conditions that have previously been described (Olsson-Francis *et al.*, 2016). The PCR products were purified with ethanol and sequenced using the Illumina NovaSeq platform (Mr DNA, US). The QIIME2 pipeline was used to process the raw data. All chimeric sequences were removed by using the DADA2 noise algorithm, and the remaining sequences were trimmed (<15 bp and >270 bp). The sequences were clustered into amplicon sequences variants (ASVs), and phylogeny was assigned using Scikit-learn classifier, which was based on the Greengenes database (DeSantis *et al.*, 2006) (confidence threshold of  $p=0.7$ ).

### 3. Results

Mid infrared and visible near infrared spectra averaged from each site of interest are shown in Fig. 3a and 3b, respectively. Only one Vis-NIR spectrum is presented, as all analyses taken showed the same features. The fundamental vibrational modes of gypsum (CaSO<sub>4</sub>·2H<sub>2</sub>O) were found to correspond to the main peaks in the MIR and the overtone and combinational modes to the absorption bands in the Vis-NIR. The assignments for the mineralogical components of the samples are consistent with previous studies such as Liu (2018).

#### 3.1. Mineralogy

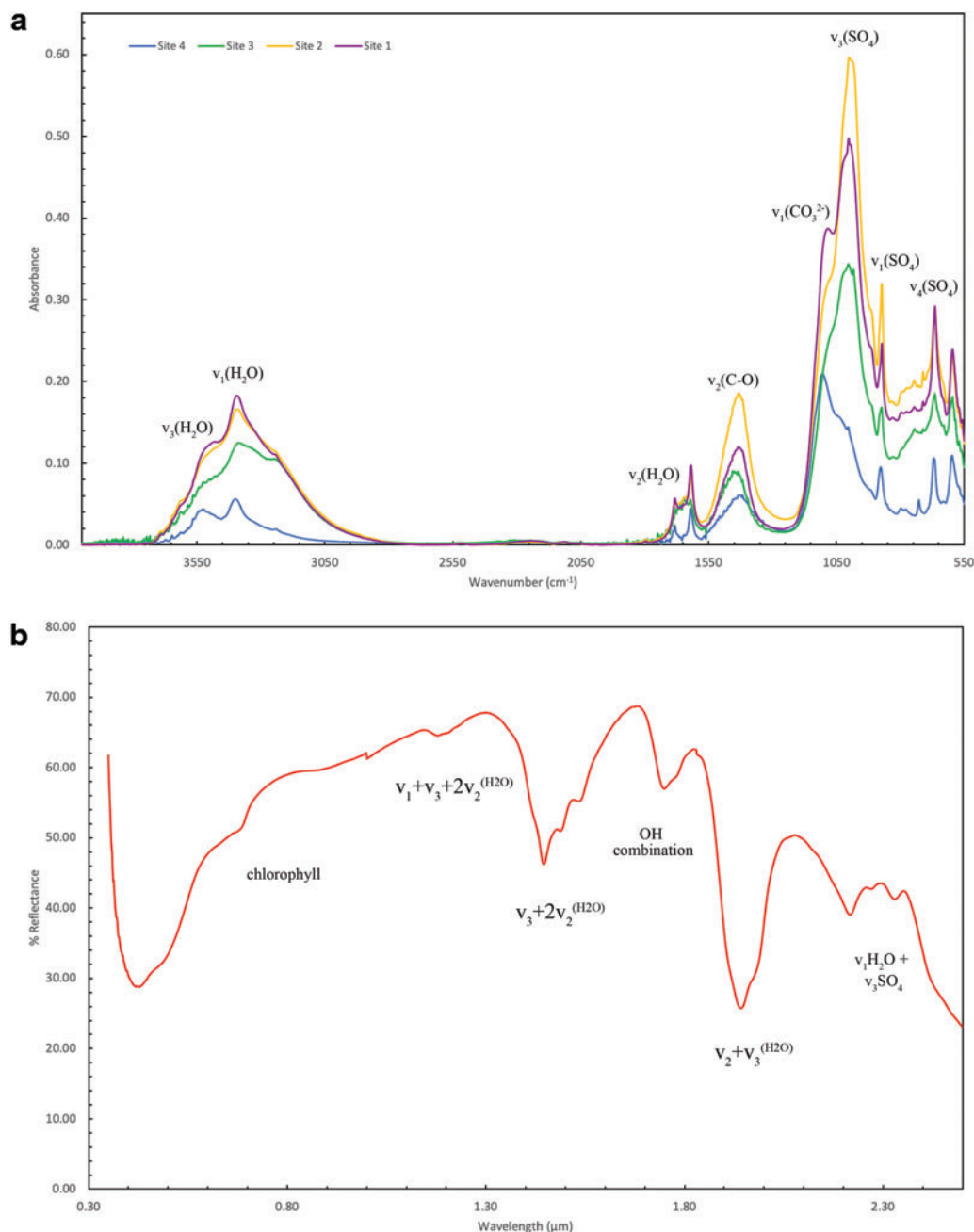
In the MIR, the strongest absorption peaks in samples from site 1 to 4 were found at 1002, 1001, 1003, and 1105  $\text{cm}^{-1}$  respectively. This corresponds to the  $\nu_1$  symmetric and  $\nu_3$

asymmetric stretching vibrational modes of SO<sub>4</sub> tetrahedra. Two antisymmetric  $\nu_4$  bending vibrations of SO<sub>4</sub> tetrahedra were also observed around 664–667 and 596  $\text{cm}^{-1}$ . Within the gypsum spectrum there are also stretching and bending  $\nu_2$  vibrational modes of C-O of carbonate located at 1455 and between 872 and 877  $\text{cm}^{-1}$ . The presence of water in the gypsum can be discerned by peaks in the region near 3500 and 1600  $\text{cm}^{-1}$ . The  $\nu_1$  symmetric stretching vibrational modes of H<sub>2</sub>O are located around 3400  $\text{cm}^{-1}$  and the  $\nu_3$  antisymmetric stretching vibrational modes at 3520 and 3640  $\text{cm}^{-1}$ . There are two peaks caused by the O-H bending vibrational modes located at 1682 and 1619  $\text{cm}^{-1}$ . This double peak indicated there are two distinct crystallographic types of water—one hydrogen bonded to the sulfate ions and one linked to calcium ions. In addition to gypsum, there are also vibrational modes indicative of carbonate ions (CO<sub>3</sub><sup>2-</sup>) within samples from sites 1 and 2. A  $\nu_1$  stretching vibration is located at 1082  $\text{cm}^{-1}$ , and the  $\nu_4$  vibration is split with peaks observed at 746 and 711  $\text{cm}^{-1}$ . A minor  $\nu_3$  stretching vibration of CO<sub>3</sub><sup>2-</sup> ions is observed in samples from site 4. In samples from site 3, a strong shoulder is observed on the 1003  $\text{cm}^{-1}$  peak, assigned to the Si-O asymmetric stretching vibration of SiO<sub>4</sub> tetrahedra.

In the NIR, all spectra displayed the same absorption bands indicative of gypsum, albeit at variable intensities. Reflectance minima were observed at 2.22, 2.27, and 2.33  $\mu\text{m}$  due to the combinational mode of  $\nu_1\text{H}_2\text{O} + \nu_3\text{SO}_4$ . A strong absorption at 1.94 and 1.97  $\mu\text{m}$  is that of H<sub>2</sub>O combinations ( $\nu_2 + \nu_3$ ) H<sub>2</sub>O vibrational modes produced by structural water molecules, at 1.75  $\mu\text{m}$  due to an OH combination, and at 1.18  $\mu\text{m}$  of the combination of the H<sub>2</sub>O bending fundamental and the first overtone of the OH stretching fundamental. A diagnostic yet complicated band shape of gypsum is observed between 1.4 and 1.5  $\mu\text{m}$  as a reflectance minima triplet caused by OH stretching.

#### 3.2. Organics

There are very few absorptions in the MIR that can be unequivocally attributed to organic molecules within the samples. A discussion later will look into the difficulties of organic versus mineral absorption band interpretations. There are a few bands, however, that cannot be attributed to the mineralogical host of the samples and so have the potential to be due to organic materials. Across all samples there are absorption bands at 1747 and 1795  $\text{cm}^{-1}$  which may be caused by C=O stretching vibration of a carbonyl group. In addition, a minor peak at 1434  $\text{cm}^{-1}$  potentially of a C-O-H in-plane bending functional group is observed. A number of absorptions could be assigned to nitrogen-bearing functional groups of amides and amines. In samples from sites 1, 3, and 4 a C=O stretching band is observed at 1651–1655  $\text{cm}^{-1}$  sandwiched between absorption bands around 1681 and 1621  $\text{cm}^{-1}$ . These may be created by the presence of a primary amide within the samples. In samples from sites 3 and 4 located toward the edge of the lake, a peak around 1563  $\text{cm}^{-1}$  is observed of the N-H in-plane bending vibration of secondary amides. In all spectra, a shoulder peak is observed around 3236  $\text{cm}^{-1}$ . This may be due to the N-H stretching vibrational mode of secondary amines. Finally, in spectra from sites 1 and 2 in the center of the lake, twin absorptions at 746 and 711 may be attributed to the N-H out-of-plane bending vibration of secondary amides and the



**FIG. 3.** (a) Mid infrared spectra from sites 1–4 with the main mineralogical functional group vibrations identified. (b) A Vis-NIR spectrum from site 4 with the main mineralogical combination and overtone absorptions identified. The absorption band of chlorophyll at 0.67  $\mu\text{m}$  is also highlighted. Color images are available online.

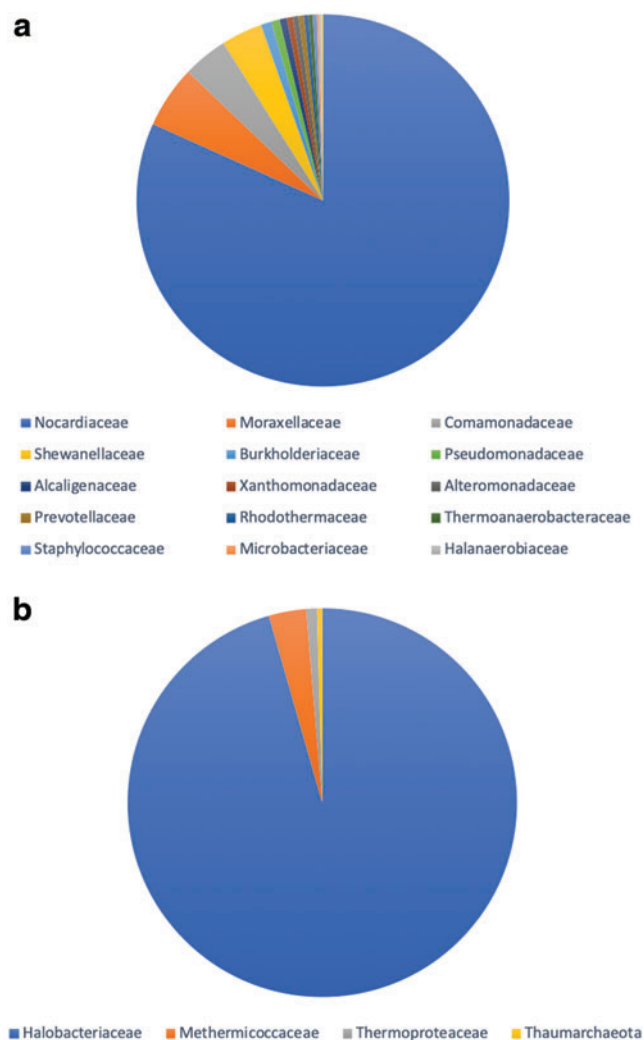
NH<sub>2</sub> out-of-plane bending vibration of primary amides; however, it is more likely that of carbonate ions.

In the Vis-NIR there is a strong drop in reflectance shortward of  $\sim 0.8 \mu\text{m}$  which corresponds to the “red edge” of chlorophyll, and a chlorophyll absorption feature is observed at  $\sim 0.67 \mu\text{m}$ . There are weaker features below this wavelength that can probably be attributed to chlorophyll and other photoprotective pigments such as carotenoids. Absorption bands near 1.75 and 2.2–2.35  $\mu\text{m}$  were observed that could be identified to arise from the combination and/or overtone modes of the fundamental CH stretching vibration,

which although a possibility in these samples are more likely consistent with fundamental vibrational modes and overtones/combination bands of gypsum.

### 3.3. Composition of the microbial community

Partial sequencing of the 16S rRNA gene was used to identify the major bacterial and archaeal classes within the sample. Quality filtration of the raw data resulted in a total of 50954 sequences (medium read length 390 bp). Due to the length of the sequences, identification was carried out to a



**FIG. 4.** Relative abundance of the most common microbial (**a**) and archaeal (**b**) families identified in evaporitic material collected from Tírez Lake. Color images are available online.

family level. Taxonomical assignment demonstrated that the archaeal community was dominated by Halobacteriaceae (96%) while the bacterial community was dominated by the class Nocardiaceae (80%), as shown in Fig. 4a and 4b.

## 4. Discussion

### 4.1. The preservation of organics within gypsum

The stability of any biosignature is dependent on its initial form, the matrix in which it is hosted, and the chemical and physical processes it is subjected to over time. Based on observations of terrestrial analog sites, there is a general consensus that sediments, evaporites, and hydrothermal systems provide the highest preservation potential for biosignatures. On Earth, entombment of biosignatures within mineral matrices constitutes one of the most efficient preservation mechanisms against degradation of both microorganisms and organic molecules. This occurs through mineralization processes, including those driven by evaporation and precipitation, and enrichment inside mineral

matrices as evidenced within sedimentary rocks and phyllosilicate minerals. Here the reduced permeability limits the exposure of entrapped organics to migrating fluids and gas, hence preserving their composition or structure. However, the persistence of the sedimentary (or other) material is a key factor for long-term preservation; if the mineralogical host matrix fails, then the entombed biosignatures may be lost. Sulfates, such as the gypsum described in this study, are not the optimal material for biosignature preservation, as they are susceptible to dissolution and mineralogical transformations; however, it has been observed that they can offer fossil stabilities for timescales of up to  $1 \times 10^6$  years (Farmer and Des Marais, 1999) and more given the right conditions. The ultra-dry martian surface environment means that, after deposition, sulfate minerals such as gypsum on Mars are likely to persist for longer than on Earth with its extensive hydrological cycle.

Organic materials are often co-deposited in terrestrial evaporites and as such have been proposed as relevant analogs in the search for life on Mars. A study conducted by Aubrey *et al.* (2006) investigated concentrations of organic matter along with amino acids in natural terrestrial sulfate mineral samples. They found that sulfate minerals contain between 0.03% and 0.69% organic carbon as well as high parts-per-billion to low parts-per-million abundances of amino acids and their degradation products in samples dating back 40 million years. Evaporites have been found to preserve biosignatures by the entrapment of halophiles in fluid inclusions in both modern systems (Foster *et al.*, 2010) and potentially those up to 250 million years old (Schubert *et al.*, 2009). A study by Parnell *et al.* (2004) focused on microbial colonization in impact-generated hydrothermal crystalline gypsum deposits in the Haughton Crater, Devon Island, Canadian High Arctic. The authors demonstrated the presence of cyanobacteria in endolithic habitats located up to 50 mm from the crystal margins of clear gypsum crystals. These endolithic niches provide organisms with UV-shielding and temperature and water-activity moderation (Rothschild, 1990; Hughes & Lawley, 2003; Wierzchos *et al.*, 2006; Cockell *et al.*, 2008).

The surface of Mars is a hostile place with extremes of atmospheric composition and pressure, temperature, and UV light irradiation. The preservation of potential biosignatures is governed by the threat of destruction by chemical oxidants, a high ionizing radiation environment, and the exposure age of the minerals they are contained within. In addition to a UV-radiation refuge, evaporites can provide protection from oxidative surface processes (Stivaletta and Barbieri, 2009) and desiccation. Cloutis *et al.* (2007) found that minerals such as gypsum that contain structural  $H_2O$  are more resistant to  $H_2O$  loss under martian conditions than phyllosilicates. In a spectral study on gypsum endoliths exposed to the UV, anoxia, low pressure and desiccating conditions of Mars, Stromberg *et al.* (2014) found there was little change in the NIR spectra of the samples, with all major spectral features of both gypsum and chlorophyll preserved after exposure to UV irradiation equivalent to  $\sim 300$  years at present martian surface conditions.

### 4.2. The identification of organics within deposits at Tírez Lake

It has been confirmed that the samples collected from Tírez Lake contain both halophilic bacteria and archaea,

combined with visual evidence from site 4 of green and red horizons millimeters beneath the surface of the samples. Montoya *et al.* (2013) observed that archaeal communities in sediments at Tírez Lake showed higher diversity than within water samples from the area. They observed that around 40% of archaeal clones from both rainy and dry season sediments showed a high similarity with *Halobacterium* species with 16S rRNA gene sequence techniques showing a predominance of Haloarchaea, particularly members of the Halobacteriaceae. *Halobacterium* species, also generically known as halophiles, are commonly present in high-salinity environments and appear reddish due to the pigment bacterioruberin. It was also observed by Montoya *et al.* (2013) that Gammaproteobacteria are the dominant sequences in the sediments. Our study agrees with this observation, as the diversity within our own samples was dominated by bacteria.

The detection of organic matter in sulfate minerals, particularly gypsum, by using IR spectroscopy is challenging due to (i) the relative proportions of minerals and organics, with the former greatly outnumbering the latter, and (ii) the dominance of the sulfate ion vibrational modes and hydroxyl overtones in both the NIR and MIR spectra which occur at similar vibrational frequencies to many organic functional groups. Given the amplitude of the vibrational frequencies and excellent match between the observed spectra and previously documented examples cited above, it is logical to assign most absorbance data belonging to that of gypsum with minor contributions of carbonate and silica. This does not mean that there are no organic vibrations detectable within the sulfate samples, just that their absorptions are weaker and masked by the minerals present. There are a few peaks that are not mineralogically assignable and could belong to a few carboxyl group esters, and to some primary and secondary amides and amines. Despite the spectrally low organic content of the samples, amino acids are known to be excellent indicators for the presence of other organic material because they can be detected at very low levels, so their potential presence in the FTIR data may be significant. It should be noted that there is no observation of aliphatic structures as would be indicated by C-H stretching vibrations in the region 2960–2850  $\text{cm}^{-1}$ . C-H bonds exist in virtually all organic compounds, and the C-H stretch vibrations for methyl and methylene are the most characteristic in terms of recognizing a compound as organic. Since this region is free of any mineralogical overprint, if these aliphatic hydrocarbon vibrations were present within the samples, we would expect to see them here.

A successful interpretation of IR data both on Earth and Mars is based not only on the presence of particular bands within the spectrum but also the absence of other important bands. The 16S rRNA analyses imply that organic molecules, as either part of the microbial community present or associated with them, should be expected to be detectable within the samples, so it is at least possible that we are seeing minor representations of these within the FTIR spectra of Tírez Lake. However, the lack of the C-H stretching region, given its almost universal observation within organic compounds, implies that any organic assignments in the IR data should be viewed with caution and that the most likely interpretation is that the mineralogy of the samples is flooding the spectra.

#### 4.3. The identification of organics within gypsum on Mars

The difficulty in distinguishing organics within a sulfate-dominated system has profound implications for the search for evidence of past life on Mars. The robotic exploration of Mars has revealed that sulfate minerals are ubiquitous on the planet's surface (*e.g.*, King and McLennan, 2010). The mineralogical transition on Mars from clay-rich to sulfate-rich strata occurred as the atmospheric pressure on Mars decreased in the Late Noachian and the amount of sulfur dioxide degassing from lavas increased (Gaillard *et al.*, 2013). Sulfate-forming conditions are likely to have persisted well into the Hesperian (Bibring *et al.*, 2006; Gaillard *et al.*, 2013). Analyses at both Rocknest and the Yellowknife Bay study site by the Chemistry and Mineralogy (CheMin) instrument of the Mars Science Laboratory found only calcium sulfates (Nachon *et al.*, 2014; Downs *et al.*, 2015, Vaniman *et al.*, 2018), and oxidized sulfur species are widespread in martian meteorites, with calcium and magnesium sulfates being the most common forms (Gooding, 1992). On Earth, sulfate-forming conditions and sulfates themselves are inhabited by extremophilic microorganisms, so it is important to investigate the possibility of these deposits on Mars hosting evidence of similar life.

For *in situ* analysis of martian rocks and soils, including those rich in sulfates, organic matter can be, and so far has been, mainly identified by using thermal extraction protocols such as the Viking gas chromatograph–mass spectrometer (GC-MS), the Phoenix Thermal and Evolved Gas Analyzer (TEGA), and Mars Science Laboratory's Sample Analysis at Mars (SAM) instrument suite (Biemann *et al.*, 1977; Grotzinger *et al.*, 2012; Mahaffy *et al.*, 2012). For the upcoming ExoMars rover, a similar process will be achieved through the Mars Organic Molecule Analyzer (MOMA) instrument (Goesmann *et al.*, 2017). It has been shown, however, that, despite their high preservation potential, sulfate-bearing minerals pose a significant threat to the detection of organic compounds during thermal extraction. When sulfates decompose, they release oxygen on heating which may oxidize organic matter present in the sample of interest. A benefit of searching for biosignatures on Mars within gypsum and magnesium sulfates is that they are unlikely to yield oxygen at the temperatures commonly employed for the thermal extraction of organic matter, as under these conditions gypsum only dehydrates (West and Sutton, 1954; Gordon and Sephton, 2016). Due to the potential issues with organic detection and pyrolysis, however, it is important to be able to analyze and search for biosignatures by using nondestructive techniques, such as the IR spectroscopy utilized in this study.

The results of this work are very timely and appropriate given that the ExoMars Rosalind Franklin rover includes within its payload three IR spectrometers in the form of ISEM (Infrared Spectrometer for ExoMars), MicrOmega, and Ma-MISS (Mars Multispectral Imager for Subsurface Studies). ISEM is a pencil-beam spectrometer designed to study the mineralogical composition of the martian surface in the vicinity of the rover over the NIR spectral range between 1.15 and 3.3  $\mu\text{m}$  (Korablev *et al.*, 2017). This range is highly sensitive to the hydration state of minerals and has the ability to observe, if present, strong absorption features



associated with a number of PAH functional groups, particularly C-H molecules (Izawa *et al.*, 2014). Absorptions can be observed at 1.69  $\mu\text{m}$  due to CH overtones, N-H stretching overtones at 1.50  $\mu\text{m}$ , and many further overtones and combination bands beyond 2.1  $\mu\text{m}$ . MicrOmega is a Vis-NIR hyperspectral microscope located within the rover's analytical laboratory drawer. It is designed to characterize the texture and composition of martian samples with the ability to identify minerals, ices/frosts, and organics over the spectral range 0.5–3.65  $\mu\text{m}$  (Bibring *et al.*, 2017). Finally, there is the Ma-MISS experiment, a miniaturized spectrometer within the drill system that will perform IR spectral reflectance investigations in the 0.4–2.2  $\mu\text{m}$  range. Its goal is to study the subsurface of Mars, to characterize mineralogy and ices (De Sanctis *et al.*, 2017).

One of the key findings in this study is the ability, or lack thereof, to observe known microbial components within the samples when using IR spectroscopy, most likely due to the dominance of gypsum. The amount of organic material needed for it to be spectrally visible has been investigated by ISEM-based simulations and by Dos Santos *et al.* (2016). In the later study, the authors studied the preservation of 25 amino acids spiked onto Mars analog minerals that included clays and the sulfates jarosite and gypsum under Mars UV conditions. They found that gypsum had the highest amino acid preservation when spiked with 10 M amino acid solution equivalent to 1 mmol of amino acid per gram of mineral, whereas at 1 M solution no amino acids were detected in any of the samples, given the amino acid detection limit of the GC-MS used is 3 ppb. Overall, Dos Santos *et al.* (2016) found that the smectite and sulfate samples provided the best preservation of amino acids. In the case of gypsum this may be due to partial dissolution and recrystallization trapping the amino acids. It is important to consider the detection limits for Mars-bound spectrometers where it concerns multiple components, be it mineral mixtures, targets covered in wind-blown dust, or mineral-organic assemblages. Work by Korablev *et al.* (2017) has included studying the detection capabilities of the ISEM instrument. They found that detection limits are affected by factors such as grain size and the availability and intensity of diagnostic absorption bands as well as end-member abundances. They found for example that 1 wt % adenine is detectable in the NIR range of ISEM in the presence of 99 wt % nontronite or hematite. Further work needs to be done to study the spectral effects and detectability of sulfate-organic mixtures.

## 5. Conclusions

It appears that sulfates on Mars may represent something of a mixed blessing, acting as an effective host for organic matter preservation but also a potentially aggressive material that is difficult to analyze and masks the presence of organic functional groups when viewed using IR spectroscopy. In the case of Tírez Lake it is known that the sediments contain bacterial and archaeal communities, and as such spectral fingerprints of organic molecules should be visible. However, due to the band positions of the mineralogical host materials, these organic absorptions are hidden or are so small compared to the volume of sulfate present that their detection limit has been exceeded. Despite the difficulties encountered extrapolating organic molecular IR

fingerprints from the gypsum at Lake Tírez, future astrobiology missions should not be discouraged from targeting Ca-sulfate deposits due to their high biosignature preservation potential. Instead, a suite of complementary analytical techniques should and will be used to support the spectral identification of extraterrestrial biosignatures in the presence of gypsum.

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## Author Disclosure Statement

No competing financial interests exist.

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#### Abbreviations Used

GC-MS = gas chromatograph–mass spectrometer  
ISEM = Infrared Spectrometer for ExoMars  
Ma-MISS = Mars Multispectral Imager for  
Subsurface Studies  
MIR = mid infrared  
NIR = near infrared  
Vis-NIR = visible near infrared