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**Laboratory analysis of returned samples from the AMADEE-18 Mars
analog mission**

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Abstract

Between 01-28. February 2018, the Austrian Space Forum, in cooperation with the Oman Astronomical Society and research teams from 25 nations conducted the AMADEE-18 mission, a human-robotic Mars expedition simulation in the Dhofar region in the Sultanate of Oman As a part of the AMADEE-18 simulated Mars human exploration mission, the Remote Science Support team performed analysis of the Dhofar area, (Oman) in order to characterize it as a potential Mars analog site. The main motivation of this research was to study and register selected samples collected by the analog astronauts during the AMADEE-18 mission with laboratory analytical methods and techniques comparable to the techniques that will be used on Mars in the future. The 25 samples representing unconsolidated sediments obtained during the simulated mission were studied by using optical microscopy, Raman spectroscopy, X-ray diffraction, laser-induced breakdown spectroscopy, and laser-induced fluorescence. The principal results showed the existence of minerals and the detection of alteration processes related to volcanism, hydrothermalism, and weathering. The analogy between the Dhofar region and the Eridania Basin region of Mars is clearly noticeable, particularly as an analog for secondary minerals formed in a hydrothermal seafloor volcanic-sedimentary environment. The synergy between the techniques used in the present work provides a solid basis for the geochemical analyses and organic detection in the context of future human-robotic Mars expeditions. AMADEE-18 has been a prime test bed for geoscientific workflows with astrobiological relevance and has provided valuable insights for future space missions.

Keywords: Planetary exploration, combined instrumentation methods, Astrobiology, simulated space mission.

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Introduction

AMADEE-18 was an integrated Mars analog mission conducted by the Austrian Space Forum in the Dhofar region in the Sultanate of Oman between 01-28th February 2018. The activities were carried out in the proposed terrestrial Martian analog site and directed by a Mission Support Center in Innsbruck, Austria (Groemer *et al.*, 2019). During the planning stage of the mission, a sequential method for the detection of biomarkers for future human and robotic Mars missions was developed. Garnitschnig (2018) proposed a framework for the detection of potential biomarkers and compiled a list of potential test sites for future missions (Groemer *et al.*, 2019; Groemer, 2018a). These kinds of methods and frameworks pursue a double objective with strong implications in the search of life on Mars and optimization in the synergy of analytical techniques at different test sites for future Martian missions. The first objective was to evaluate the impact of carbon detection, water evidence, and characterize any biosignature. The second objective was the rational selection of technique to be used, which was based on the Martian mission such as Mars2020 and ExoMars suit equipment. Thus, the synergistic combination of analyses are of interest for target selection, deployment outcrop site and instrumentation capabilities for water and possible life detection (Garnitschnig, 2018; Gruber *et al.*, 2019; Stromberg *et al.*, 2019).

Several in-situ experiments were used during this study such as the FieldSpec and ScanMars systems, which utilize reflectance spectroscopy and ground-penetrating radar GPR (described in detail in this issue). Other lab-based techniques that we have used during our analysis are micro-imaging, Raman spectroscopy, X-ray diffraction (XRD), laser-induced breakdown spectroscopy (LIBS), and laser-induced fluorescence (LIF). The objective of these methods is to have a full biological and geological understanding of the surface. In this work, we present the results of the analysis done using lab-based techniques and illustrate how they can be used to qualitatively and quantitatively describe landscapes in situ, in addition to analyzing samples collected by humans during analogue and/or actual space missions. This may facilitate the inference of a surface suitability for subsequent analyses. For example, the lab-techniques can help in future selection of instrumentation and design of the workflow for a possible field-survey.

The main motivation of this research was to study selected samples collected by the analog astronauts during the AMADEE-18 mission with laboratory analytical methods and techniques proposed in the strategic framework developed by (Garnitschnig, 2018) for the AMADEE-18 mission and comparable to the techniques that will be used on Mars in the future. The analyses were performed at the Planetary Exploration Instrumentation Laboratory at York University and the Royal Ontario Museum, both in Toronto, Canada. The results and measurements exemplify the capabilities of each technique, and the combined advantages. Furthermore, the results emphasize the selection applicability of portable twin systems—those that resemble real flight

instruments used in past/present/future Martian missions, and also based in the same technology—for future analog missions. Table 1 compares laboratory-based rover instrumentation from past, current and future missions. Finally, SEM, EDS and XRD as described above, can be used to validate field experiments such as LIBS, Raman Spectroscopy and LIF.

Methods used in this study and their relevance to Mars studies

Raman spectroscopy is a non-destructive spectroscopic technique that does not require sample preparation. The technique is well suited for in-situ analyses of rocks and minerals, and has been used for mineral identification and organic detection of the target sample (Edwards *et al.*, 2013; Ferraris *et al.*, 2012). Raman spectroscopy has been proposed as a method for geological identification for Mars-related materials, and it will be used during the future Martian missions: SuperCam and SHERLOC (NASA Mars 2020) or the ESA-RLS System (ExoMars) (Beegle *et al.*, 2015; Rull *et al.*, 2017; Wiens *et al.*, 2016).

LIBS utilizes emissions from plasma created at the surface of a sample via high-power laser pulses to perform quantitative chemical analyses (Wiens *et al.*, 2012). One of the primary benefits of LIBS is that there is no required sample preparation prior to measurement and only optical access to the sample is required. This makes it a formidable candidate for stand-off chemometric analysis (Cremers and Radziemski, 2013; Konstantinidis *et al.*, 2019). For these reasons LIBS was selected for flight on Curiosity in the ChemCam instrument (Anderson *et al.*, 2015), and will fly on Mars 2020 (Wiens *et al.*, 2016).

LIF provides a method for organic and mineral detection through excitation of molecules by the absorption of laser light followed by spontaneous emission of light (Storrie-Lombardi *et al.*, 2009). Typically, LIF used to be understood as an undesired byproduct of a Raman measurement, but (Eshelman *et al.*, 2018) has shown it to be useful for detection of organic such as organic carbon and amino acids, among others. SuperCam and SHERLOC (NASA Mars2020 Rover) will also incorporate this technique in their search for evidence of past or present life on Mars (Beegle *et al.*, 2015; Wiens *et al.*, 2016). SHERLOC utilizes LIF as a supporting tool for selection of targets with possible organics via fast mapping. Subsequently, Raman spectroscopy will be able to determine the mineralogy and possible existence of biosignatures (Beegle *et al.*, 2015).

XRD allows the user to obtain mineral identification and structural characterization, and is commonly used for crystallography and mineralogy (Klein, 2008). In space, ChemMin, onboard the MSL Curiosity rover, combines XRD with X-ray fluorescence (Bish *et al.*, 2013). The ChemMin system can identify and quantify the minerals present in rocks and soil in order to understand the habitability of Mars.

Other supportive methods such as scanning electron microscopy (SEM-EDS) allow us to analyze the mineralogic composition which, in combination with the in situ petrological description and in lab optical mineralogy study, provides information about regolith material source, petrological composition, identification of weathering and hydrothermal processes affecting the Martian rocks. A summary of the techniques used to determine mineral and elemental composition, detect organics, and deduce the geological context is presented in Table 1.

Table 1. Comparison of the capabilities of the 5 systems employed in this investigation. (RLS=Raman Laser Spectrometer), (ChemCam=Chemistry and Camera), (Sherloc=Scanning Habitable Environments with Raman & Luminescence for Organics & Chemicals), (CheMin=Chemistry and Mineralogy), (MSL=Mars Science Laboratory)

Laboratory Technique	Mineral Identification	Elemental Composition	Organic detection	Geological Context	Laboratory instrumentation location in this study	Rover Instrument Equivalent
Raman Spectroscopy	Yes	Potentially (only major elements)	Yes	Yes (mapping)	Royal Ontario Museum	RLS (ExoMars) SuperCam (Mars 2020) Sherloc (Mars2020)
Micro-imaging	Yes	No	Yes	Yes	York University	CLUPI (ExoMars)
XRD	Yes	No	No	Potentially	Royal Ontario Museum	CheMin (MSL)
LIBS	No	Yes	Potentially	Yes (mapping)	York University	ChemCam (MSL) SuperCam (Mars 2020)
LIF	Potentially	No	Yes	No	York University	Sherloc (Mars 2020)
SEM-EDX	Yes (Imaging)	Yes (Qualitative)	Yes	No	York University	

Geology of the site and its analogy with Mars

In the Dhofar region a pre-rift, syn-rift and post-rift sequence ranging from Upper Cretaceous pre-rift sediments until Upper Middle Miocene post-rift deposits units can be found. These series involve three sedimentary groups (Roger *et al.*, 1989): The Hadhramaut Group, Paleocene to late Eocene in age that overlays in an unconformity over cretaceous strata consisting in shallow-marine carbonate units (Lepvrier *et al.*, 2002); The Dhofar Group, late Eocene, laying unconformably over Hadhramaut Group represented by two limestones units part of a shallow lacustrine to shallow marine deposits (Lepvrier *et al.*, 2002) and the Fars Group, early Miocene, compose by conglomerated and carbonate deposits units (Lepvrier *et al.*, 2002). Figure 1 and 2 show The Paleocene–Miocene stratigraphic units of the Dhofar region Simplified geological map of southern Oman (Dhofar area) (from (Robinet *et al.*, 2013).

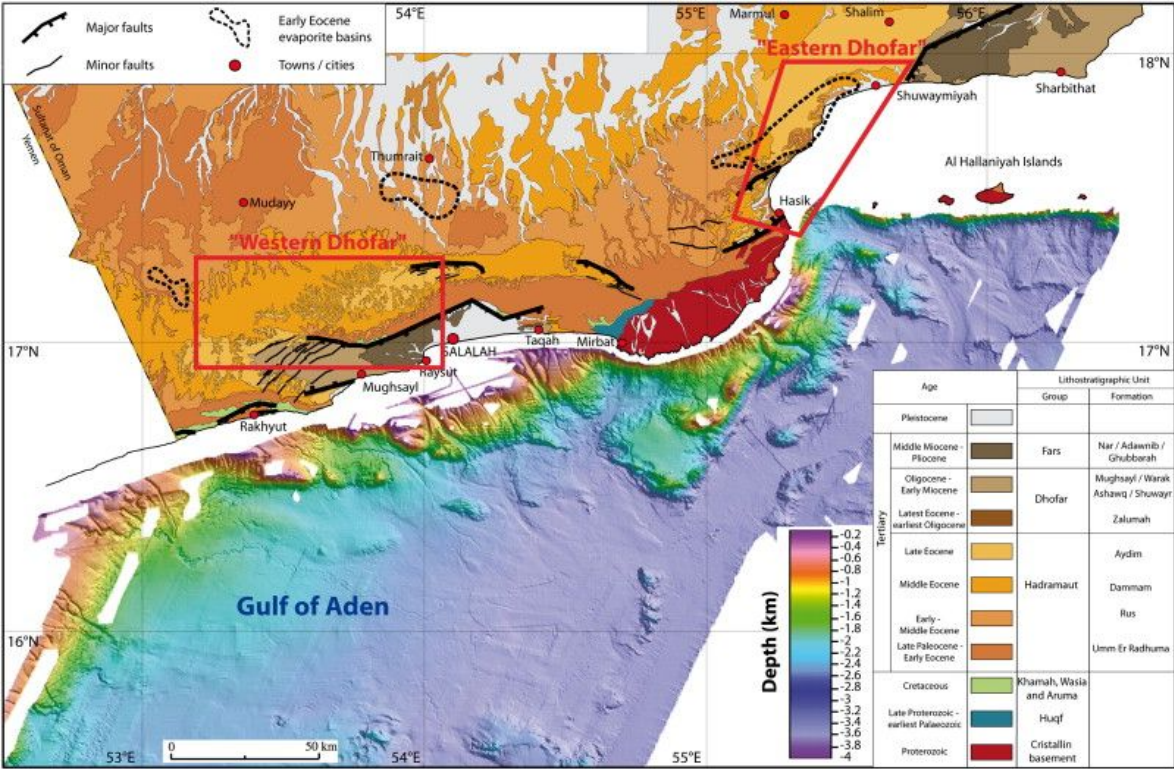


Figure. 1. Simplified geological map of southern Oman (Dhofar area) showing the two studied areas discussed in this paper from (Robinet *et al.*, 2013). (Reproduced by permission of Elsevier).

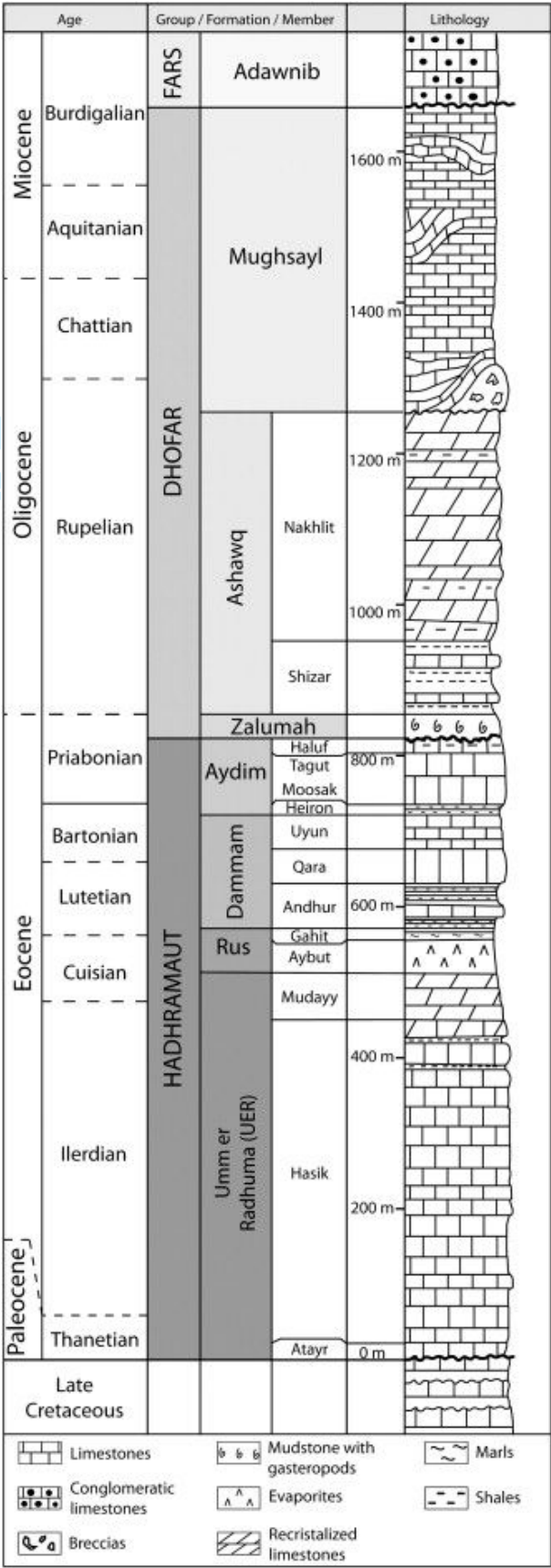


Figure. 2. The Paleocene–Miocene stratigraphic units of the Dhofar region from (Robinet *et al.*, 2013). (Reproduced by permission of Elsevier).

In the study area, the geological records are register by outcrops of the Huqf Supergroup, in the NE elongated salt basin in south Oman were uplifted blocks become sites of carbonate deposition, whereas basinal transtensional depressions were overlain with black shale and silicite (Amthor *et al.*, 2005). These belonging to the Ara Fm depositional times (Ramseyer *et al.*, 2013) as can be observed in Figure 3.

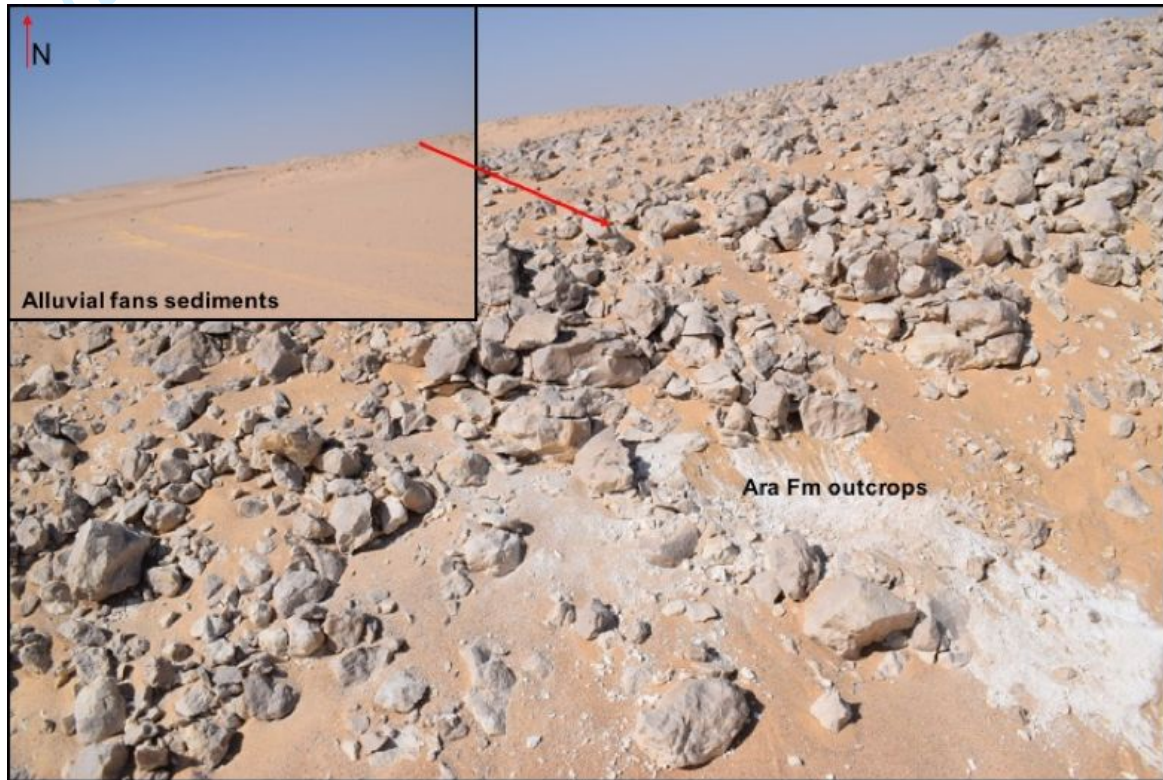


Figure 3. Picture of field site showing the outcrops and alluvial fans deposits from were samples were taking and study in laboratory.

Quaternary deposits of alluvial fans were found during the field study; these unconsolidated sediments are present together with outcrops ranging from cm up to 1 meter approximately of silicified dolomites and organic-rich laminated chert (silicilyte) belonging to the Ara Fm as observed in Figure 3. The alluvial fans are composed mostly by sand to pebbles to boulders and cemented by poorly sorted rolled clasts covered by thin dark oxidized coating (Yuan *et al.*, 2016) (Figure 3).

The feeding area of the sediments for the alluvial fans comes from the northwest-north region product of erosion and weathering of the Oman mountains outcrops and from the Huqf Supergroup sediments mention before outcropping in the area.

Part of those no consolidated sediments of the alluvial deposits includes volcanic rocks of alkaline to basaltic composition belonging to the volcanic rocks interbedded with the Precambrian-

Cambrian deposits, limestones and silicified dolomites fragments, and fragments of a hydrothermal altered silicified dolomite-carbonates and volcanic rocks that includes magnetite and sulfides like the remnant product of the hydrothermalism observed before in the Oman mountains (Reuning *et al.*, 2007)

The hydrothermalism was vaguely studied for Dhofar area, its believed to be related to magmatism in part with the intrusion of late-stage veinlets carrying metals of magnetite and sulphides. A possible supergene alteration was also identified on the samples during the petrological description and field studies with secondary mineralization of alunite, coesite and montmorillonite in the sediments together with oxides and secondary chlorite.

The syn-rift deposits observed in the regolith and in the Ara Fm could correspond to possible Martian analogs. The Precambrian-Cambrian carbonates interbedded with volcanic products, and the development of hydrothermal deposits due to active magmatism in a spreading region from the study area can be correlated to similar regions from Mars' surface, such as reported by Michalski *et al.* (2017). For example, Eridania Basin shows a 3.8 Billion-year record from a spreading area with active high magmatism and signatures of high Mg-Fe rich clay mineral products of hydrothermalism in an ancient lake base with high content of carbon and Ca-Fe-Mg rich carbonates (Michalski *et al.*, 2017). Even though the chemistry of Eridania Basin on Mars is still unknown, strong evidence from infrared spectroscopy and high-resolution imaging suggest that the Eridania Basin contains a complex suite of alteration minerals that likely formed in a hydrothermal seafloor volcanic-sedimentary setting with intrusions of sulfide-rich fluids interacting with ultramafic-mafic volcanic rocks and sediments (Michalski *et al.*, 2017). Even so, more areas in the Martian surface show similarities to geological records from selected places of ancient Earth (Léveillé, 2009). Therefore, our future understanding of Martian geo-chemistry and ancient development of organic rich carbon deposits in similar environments can be improved with the study of these analogs on Earth.

Experimental analysis and setup

In total, 25 geological samples, mostly consisting of various unconsolidated sediments were collected by the analog astronauts at the different locations where the in-situ experiments were carried out throughout the AMADEE-18 mission (See Figure 4a and 4b). The locations where samples were collected and the field cataloging procedures were conducted are presented in the Field Activity Plans from the mission (Groemer, 2018; Sejkora *et al.*, 2018). The detailed geographical positions of the geo-sampling are available in the online supporting material.

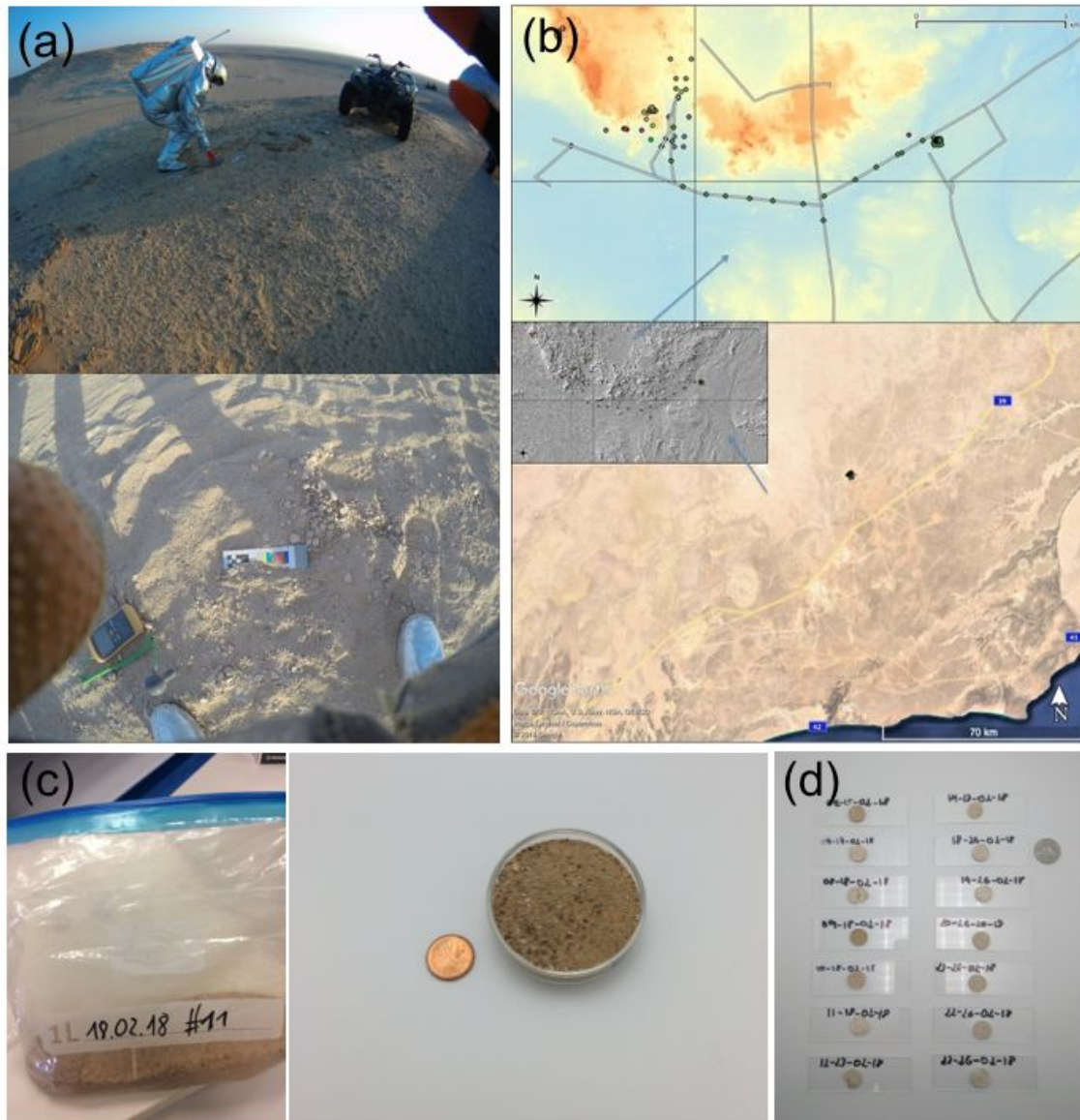


Figure 4. (a) Analog astronauts doing a geo-sampling in the field during the AMADEE-18 mission, (b) Digital Model Elevation (DEM) map of the sampling points with the location, and Digital Elevation Map (DEM) (the coordinates of sampling location are on the supporting material), (c) example of one the geo-samples cataloged at York University, and (d) pelletized sand samples.

The samples were composed of unconsolidated sediments with grain sizes ranging from $25\ \mu\text{m}$ to a few mm in diameter. For practical reasons, LIBS and SEM-EDS measurements need to have a pelletized sample before carrying out the measurements. The sample preparation was conducted to ensure the same mineral and organic compositional properties. A small amount (5-10 g) from each sample was powdered manually in an agate mortar and sieved to obtain particles ranging from $45\text{--}150\ \mu\text{m}$ in size. Powders were mixed with 1 ml of distilled water and compressed. These humidified pellet samples were left to dry by evaporation at high vacuum and room temperature.

This resulted in the formation of stable pellets. To carry out the measurements, the pellet was glued to a microscope slide for easy handling (See Figure 4c).

Micro-Raman measurements were made with a Horiba Lab-RAM Aramis Raman spectrometer. The analyses employed a 532 nm 50 mW laser focused to a 1.3 μm spot (FWHM). The spectrometer configuration used a 100 μm slit, and a 1200 grooves/mm diffraction grating. The location of the points to be analyzed were selected by the operator, along with several automatic mappings. A total of 40 different points per sample were acquired. The routine for obtaining the spectra was: selecting the point, taking a preview spectrum (fast measurement for good signal and focusing condition), and finally acquiring the spectra. The measurement conditions (acquisition and accumulations) for the manual location were varied from point to point ranging between several second to several minutes. The automatic mapping conditions per sample were 20 points separated 0.05 cm. Measurements at each point are a cumulative spectrum of five 10 s exposures. Spectra were corrected following the standard procedure available on the OPUS software from Bruker. A background subtraction was done using the Rubberband correction method and 65 baseline points. The normalization to the maximum value =1 was carried out and spectral smoothing was used when the spectra was noisy for band identification. The Levenberg-Marquardt curve fitting was done when in some of the spectra because several Mineral Raman bands were overlapping. Mineral identification was performed by comparison to the RRUFF Database using Crystal Sleuth (Downs *et al.*, 2015; Laetsch and Downs, 2006). Voight fittings were used to deconvolute the main spectral features when the Raman bands of several minerals overlapped.

XRD measurements were carried out with a Bruker D8 Advance Diffractometer equipped with a two-circle goniometer setup and CuK_α radiation source. The system was operated with a voltage generator of 40 kV and current of 40 mA. The data was collected using a Ni-filter, low background plate, and a LYNXEYE detector. The XRD diffractograms were acquired in range $10 < 2\theta < 70$ with a step size of 0.02° and acquisition time of 1 s per step and a rotation spin of 15 per minute. Analysis of resulting diffractograms was performed using the X Powder 2004.04.71 software with PDF-2 (2010) and the American Mineralogist Crystal Structure Database (AMCSD) crystallographic databases. A background correction for each diffractogram was achieved with the Splin-autoroller and polynomial tools available X Powder 2004.04.71 that allows to calculate a background polynomial subtraction function (Martin, 2004). XRD mineral quantification, and theoretical density determination of the mixture was achieved using the reference intensity ratio (RIR) from pattern matching results with X Powder 2004.04.71 (Martin, 2004).

The micro-imaging analyses and petrological descriptions were made with a standard binocular Zeiss microscope.

Laser-induced fluorescence measurements were carried out using a 266 nm Raman-LIF system designed by (Eshelman *et al.*, 2015, 2014) satisfying selected requirements of a putative flight-worthy instrument. The excitation power was provided by a quadrupled Team Photonics diode-pumped solid state Nd:YAG laser with a 0.6 ns pulse width and 13.8 μ J pulse energy at 1000 Hz. The acquisition system was composed of an Andor Shamrock 163 spectrometer. A 600 lines/mm grating allowed for an observation window between 270 to 550 nm. The spectrometer is coupled to an intensified, cooled CCD (Andor iStar 334T). The 1 cm long, 20-point LIF measurements are each an accumulation of 40 spectra with 0.01 s exposures. The spectral corrections were done in a same way as the Raman analysis.

The elemental composition of the AMADEE-18 geo-samples was measured using a Vega TESCAN Scanning Electron Microscope (SEM) equipped with a Bruker Quantax energy dispersive X-ray (EDS) detector. The beam voltage used for secondary electron imaging (SEI) and backscatter electron imaging (BSE), as well as EDS spectra acquisition, was 10 kV. SEM-EDS data were collected on the pelletized sand samples without any coating.

LIBS measurements were achieved with a breadboard system. Plasma excitation was obtained with a 1064 nm Quantel ULTRA Nd:YAG laser with 10 mJ pulse energy and 10 ns pulse width, focused to a 33 μ m spot (FWHM). The peak power density at the target is on the order of 100 GW/cm². The light produced from the plasma is collected through an off-axis f/7 collection system and delivered via fiber to the Andor ME5000 echelle spectrograph combined with the ICCD Andor iStar 334T camera (covering 230-850 nm). The spectrometer was wavelength calibrated using an Ocean Optics HG1 Calibration Light Source and intensity calibrated using an Ocean Optics DH-3P-CAL Calibration Light Source. Pelletized samples were used for LIBS analysis. Each LIBS measurement consists of 15 cumulative spectra with 0.1 s exposures (one spectrum per laser shot, for a total of 15 shots). An intensity correction was performed using the standard ANDOR Solis software procedure for the ME5000. Identification of element present within the spectra were carried with the Solis software and manual identification with the NIST Spectral Lines Database (National Institute of Standards and Technology, 2016).

Results

Field samples petrological description

14 of the most representative regolith samples from the Quaternary alluvial fans sediments, silicified veinlets intruding Ara Fm outcrops, and the silicified dolomite and carbonate rocks samples from the Ara Fm outcrops were selected in order to describe the mineralogy and

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3 petrology. These studies were later used to compare with the data acquired from LIBS, Raman
4 and SEM-EDS laboratory analyses.
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7 A microscope magnifier Zeiss lens was used for the petrological description. Varying grain sizes
8 from sand to pebbles and gravel were observed (Figure 5 and 6).
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11 During the petrological study of the sediment samples, sand and pebble sizes of calcite, dolomite,
12 silicified dolomite rocks and quartz rocks fragments were observed. These present rounded to
13 sub-rounded shapes with sizes up to 1 mm. All of them show an irregular oxidized coating on
14 their surfaces. They represent more of the 80% of the sediment components.
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18 In minor quantities, less than 10%, sub-rounded to sub-angular fragments of volcanic rocks from
19 intermediate to basic composition (andesites and basalts) are present. These rocks mostly show
20 an altered matrix, with presence of secondary clay alteration (possible montmorillonite and
21 coesite), plagioclase, amphiboles, pyroxenes crystal observed embedded in matrix.
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25 Sediments also show single crystals in less than 8% and 0.5 to 0.1 mm sizes range belonging to
26 unidentified feldspars in angular to sub-angular crystals; unidentified plagioclases, prismatic sub-
27 angular crystals, many of them with clay alteration on their borders. Volcanic quartz sub-rounded
28 to rounded with dark inclusions, prismatic crystals of amphiboles, chloritized and with presence
29 of oxides inclusions on their borders, and transverse basal sections of pyroxenes, prismatic crystal
30 of chlorite and zircons were also identified.
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35 Between 2 to 5 % oxides and sulfides were present in the sediments. Alunite was observed likely
36 as replacement of sulfides, possible pyrite, in small sizes (less than 0.5 mm) of silicified veinlets
37 (Figure 5 and 6)
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41 Samples belonging to silicified veinlets and silicified dolomite Ara Fm sediments were also
42 described. The first ones show a matrix of fine microcrystalline white silica, with presence of
43 anorthoclase plagioclase crystals in sizes ranging from less than 0.1 mm up to 0.5 mm, which
44 were fractured and rotated; showing evidence of flow movement in the matrix. Magnetite and
45 sulfides were also observed. For the silicified dolomite rock fragments, a fine microcrystalline
46 matrix of dolomite replaced by silica was clear via microscope mineral identification. Some of
47 them show a small zebra pattern, related to possible hydrothermal activity as can be observed in
48 Figure 5 and 6.
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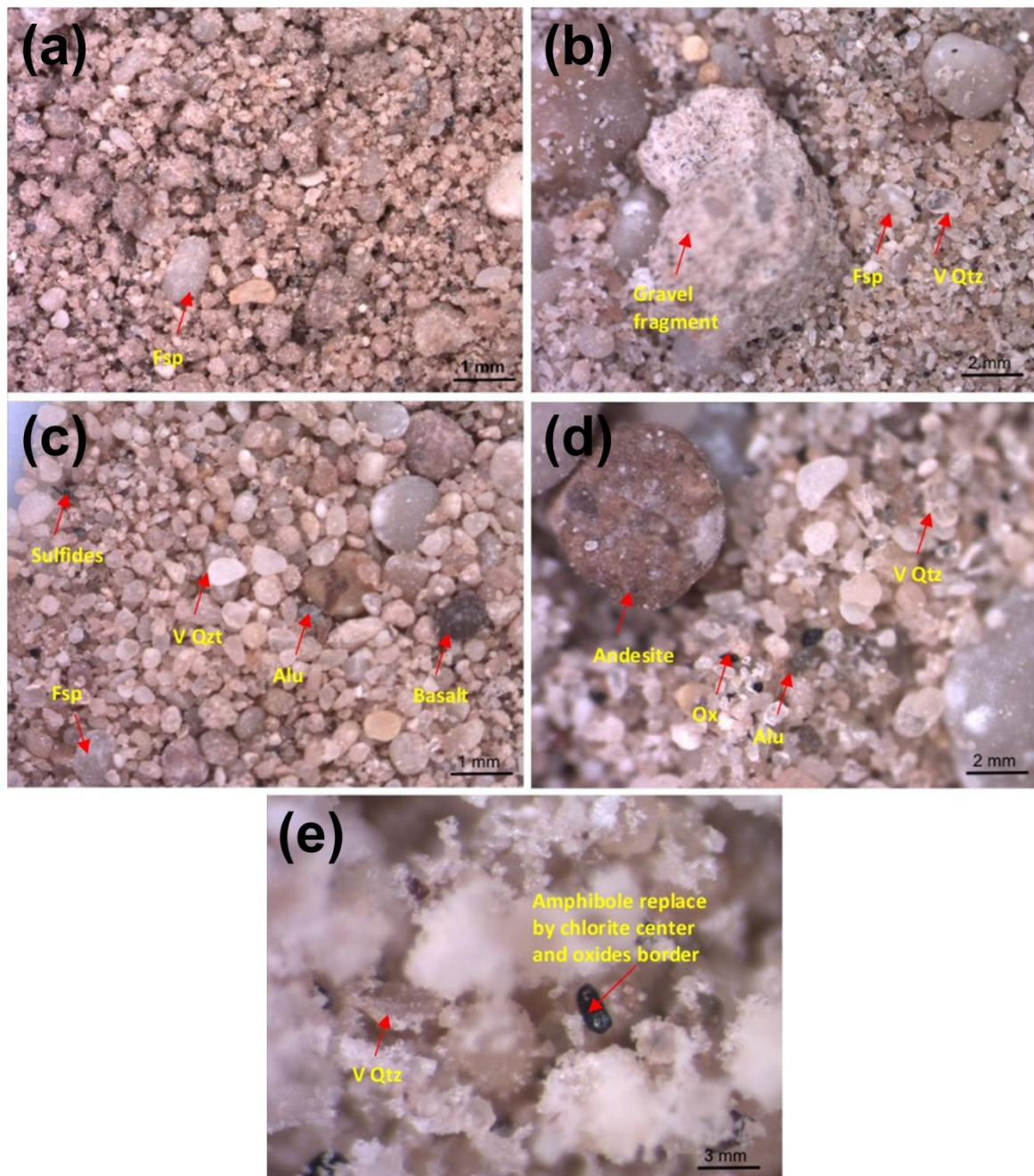


Figure 5. Magnified images of the alluvial fan sediments and sand to gravel size material. (a) Dolomites, silicate, carbonate clasts with feldspar crystals; (b) sand with presence of volcanic quartz, feldspar, disseminated oxides and gravel size clast; (c) rounded volcanic quartz, dolomite clasts, rounded basalt clast and alunite and feldspar crystals; (d) andesite sub-rounded clast with abundant presence of volcanic quartz, and oxides and disseminated alunite crystals; (e) amphibole in fine grained silica with chloritized mineralization in center and oxides in borders. Abbreviations: Fsp (feldspar), V Qtz (volcanic quartz), Alu (alunite) and Ox (oxide).

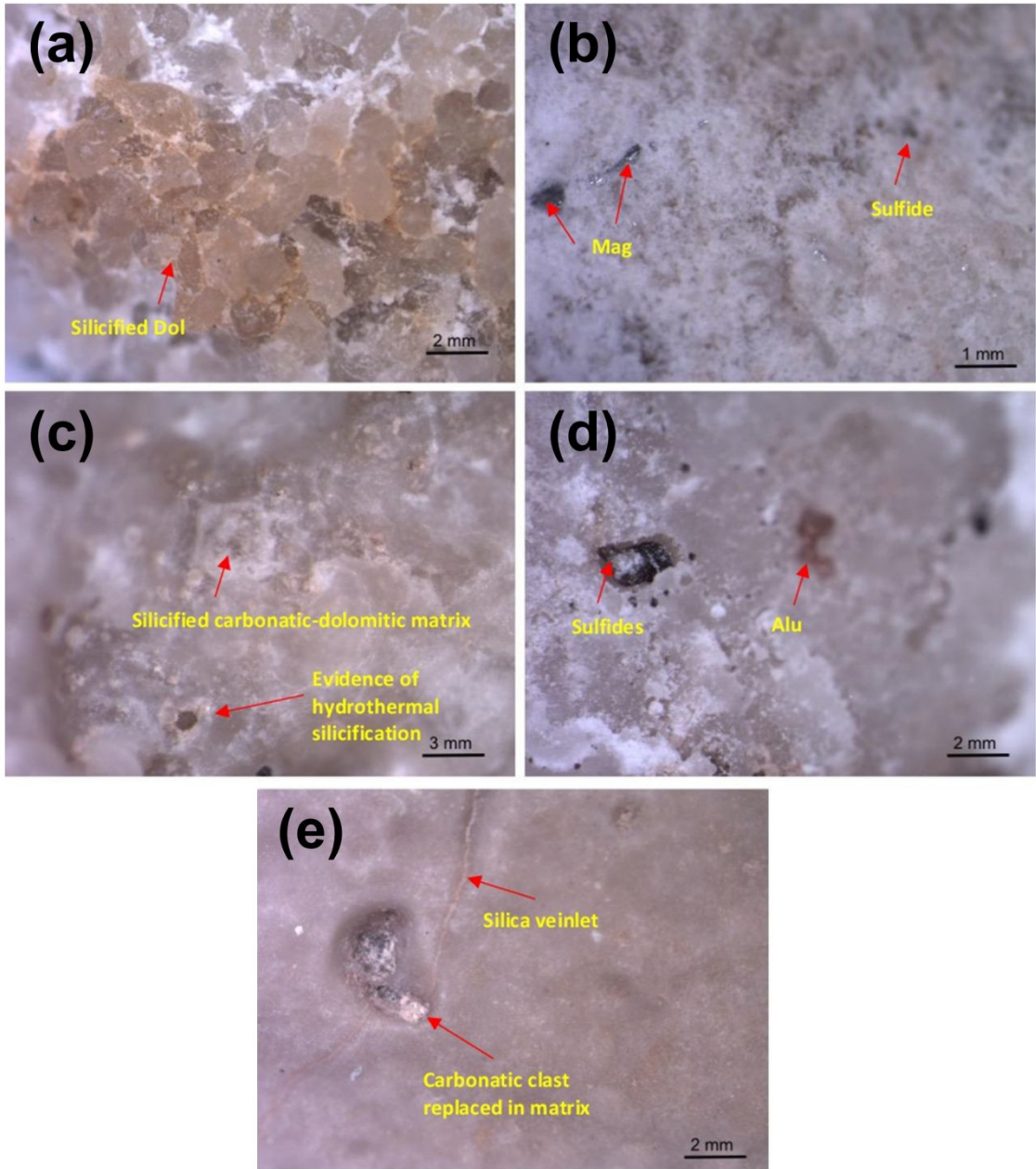


Figure 6. Magnified images of the alteration processes. (a) The silicate replacing dolomite rock from Ara Fm with oxide coating on the surface; (b) silica vein intruded in Ara Fm carrying magnetite and sulfides; (c) silicate replacing carbonate rocks from Ara Fm (possible evidence of supergen activity); (d) random dissemination of oxides and alunite crystals in silica vein matrix; (e) fine to very fine silica matrix replacing carbonate rocks (details of carbonate clast and silica veins in process of replacement can be observed). Abbreviations: dolomite (Dol), magnetite (Mag), and alunite (Alu).

Raman Analysis

The mineralogical composition results are listed in Table 2 following the ordering of Dana’s classification method (Palache *et al.*, 1952). Figure 7 shows representative Raman spectra

obtained from the samples. Identification of mineral species is done using the RRUFF Database within the Crystal Sleuth software (Downs *et al.*, 2015; Laetsch and Downs, 2006). The following references for each minerals were considered alongside the spectral analyses: oxides (Balachandran and Eror, 1982; Markovski *et al.*, 2017; Wang *et al.*, 2004), carbonates (Buzgar and Apopei, 2009; Rull-Perez and Martinez-Frias, 2003), pyroxenes (Huang *et al.*, 2000; Wang *et al.*, 2001), feldspar (Freeman *et al.*, 2008; Lalla *et al.*, 2019, 2015), clays (Apopei and Buzgar, 2010; Black and Hynek, 2018; E.A. Lalla *et al.*, 2016), and organics (Ferrari, 2007). The organics present in the main Raman bands in the range of 2800 to 3000 cm^{-1} correspond to C-H bonding and the other less intense bands near 1300 cm^{-1} correspond to C-C vibrations (Beegle *et al.*, 2015).

Table 2. Micro-Raman mineral and other material detection on the different sampling points from AMADEE-18 Mission. X indicates that the mineral was found to be present in the respective sample.

Mineral	Samples														
	04-15	05-17	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Magnetite	X	X				X		X		X			X	X	X
Hematite						X		X	X	X	X			X	X
Goethite						X									
Anatase		X					X				X	X		X	
Chromite												X			
Quartz	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Calcite	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Dolomite		X						X					X		
Titanite						X									
Olenite												X			
Enstatite		X									X				
Diopside							X				X				
Epidote													X		X
Microcline				X				X						X	
Orthoclase													X		
Sanidine				X	X	X	X					X			
Albite		X			X	X		X	X		X	X	X		X
Andesine					X			X			X				
Labradorite	X														X
Clays						X									
Carbonaceous Material and organics	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X

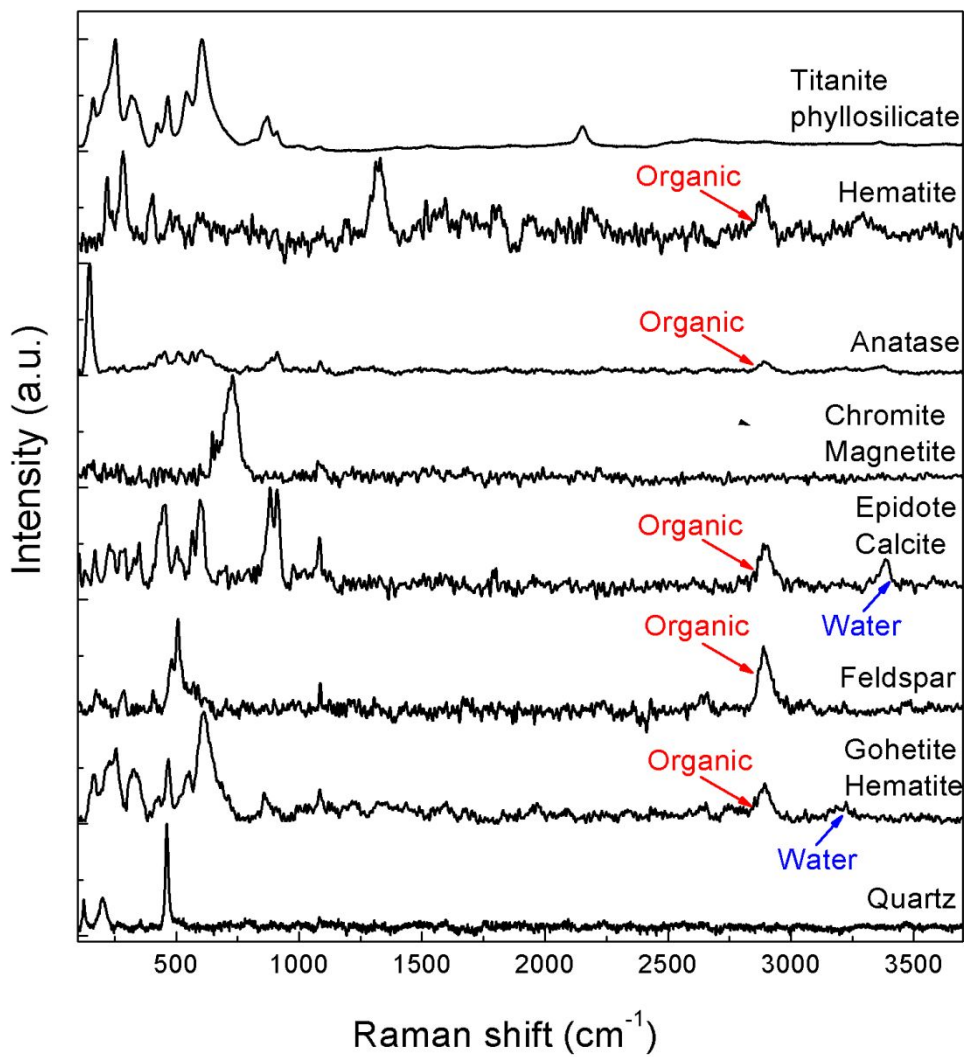


Figure 7. Micro-Raman spectra of the most significant mineral phases detected.

XRD analysis

The minerals found within the collected samples using XRD are compiled in Table 3. Figure 8 compiles the XRD diffractograms of the different sampling points and the higher concentration minerals detected on the different samples are also indexed in the figure. The detailed mineral quantification of each transect point is attached as supporting material in weight percent (wt%) including all the minor minerals.

Table 3. XRD mineral detection on the different sampling points from AMADEE-18 Mission.
X corresponds to the detected mineral in each collected sample.

Mineral	Samples													
	04-15	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Iron-Oxides					X									
Anatase	X												X	
Rutile	X												X	
Quartz		X	X	X	X	X	X		X	X	X	X	X	X
Calcite		X	X	X	X	X	X	X	X	X	X	X	X	X
Dolomite		X	X	X		X				X				
Ferrosilite								X				X		
Olenite											X			
Diopside	X													
Pigeonite							X							
Enstatite			X					X						
Epidote														X
Pyrope			X								X			
Coesite							X							
Olivine						X			X					
Feldspar									X					X
Anorthoclase													X	
Orthoclase				X										
Sanidine							X							
Anorthite					X					X				
Albite	X		X	X				X				X		
Bytownite											X			
Andesine	X	X												
Labradorite	X							X						

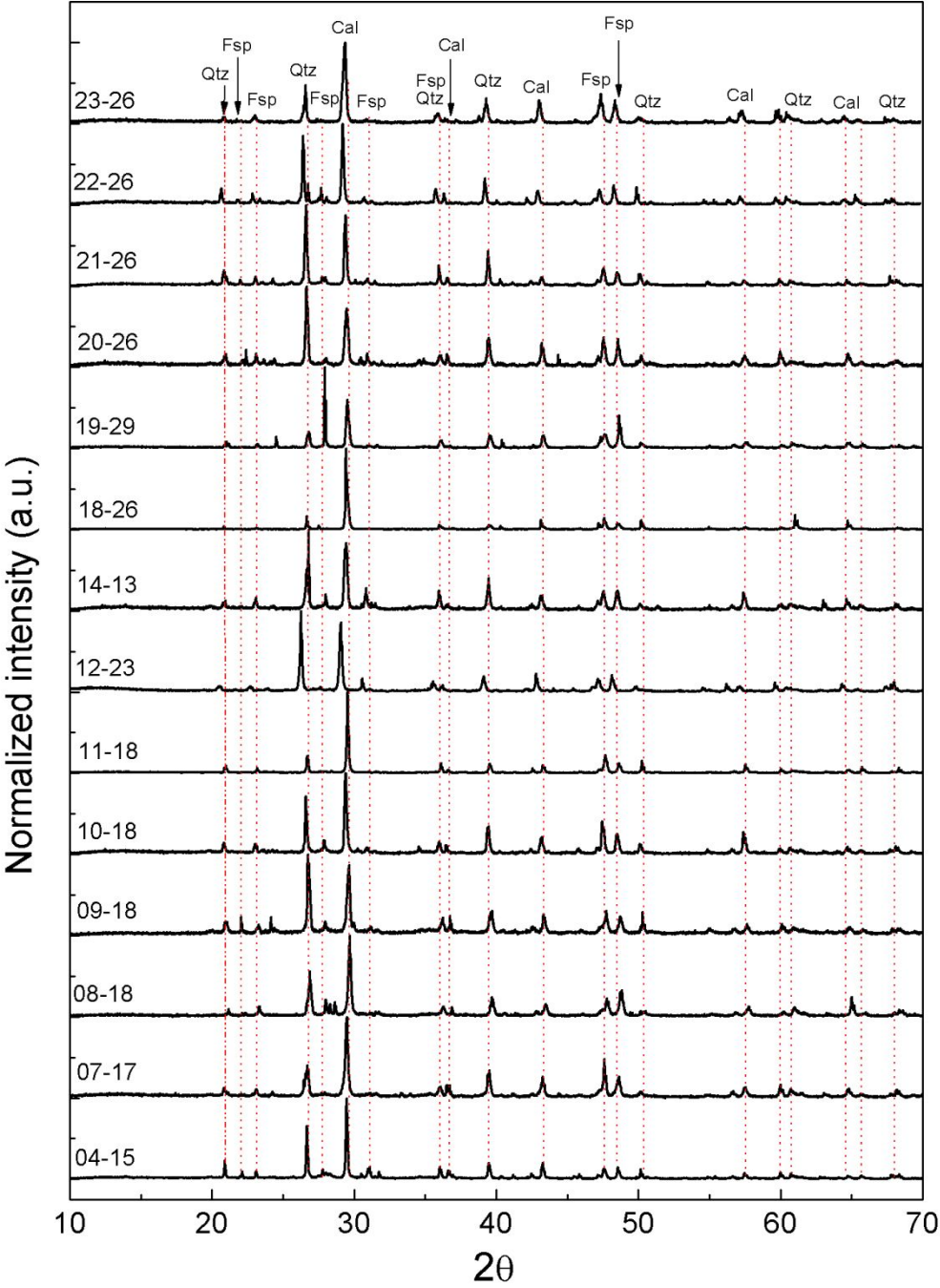


Figure 8. X-Ray diffractogram of the selected transect points from the AMADEE-18 Mission. The minerals shown are the higher concentration obtained from the quantification (see supporting materials). Abbreviations: quartz (Qtz), calcite (Cal) and feldspar (Fsp).

Laser Induced Fluorescence

A 1 cm long, 20-point LIF measurement was carried out on unprocessed sediment. Results of the LIF measurements for the different sampling points are shown in Figure 9a-d. The most intense LIF bands for organic detection are located between 300 to 450 nm (Eshelman *et al.*, 2015, 2014). The spectra from the organics detected at the collected samples were compared with the internal PIL (Planetary Exploration Instrumentation Laboratory) database (Cote *et al.*, 2018; Lymer, 2018). The results coincide with previous analyses of amino-acids like tyrosine and possibly tryptophan. Figure 9e shows a comparison between the different amino-acids used as standards and the detected organics from AMADEE-18 mission. The minerals detected coincide with carbonates (dolomite and calcite) and silica (quartz) LIF spectra from the PIL database. The dolomite is clearly differentiable on the spectra considering broad bands at ~350 and ~460 nm. The quartz presents a weak band at ~325 and the strongest broad band between 450 to 475 nm. Also, Figure 9e. presents some selected bands from the different minerals detected and AMADEE-18 results. Table 4 shows the minerals and organic signature detected for the different samples. The minimum amount of minerals and organic signatures were considered only for more than 15% of detection of the 20-point, being in more than 3 points

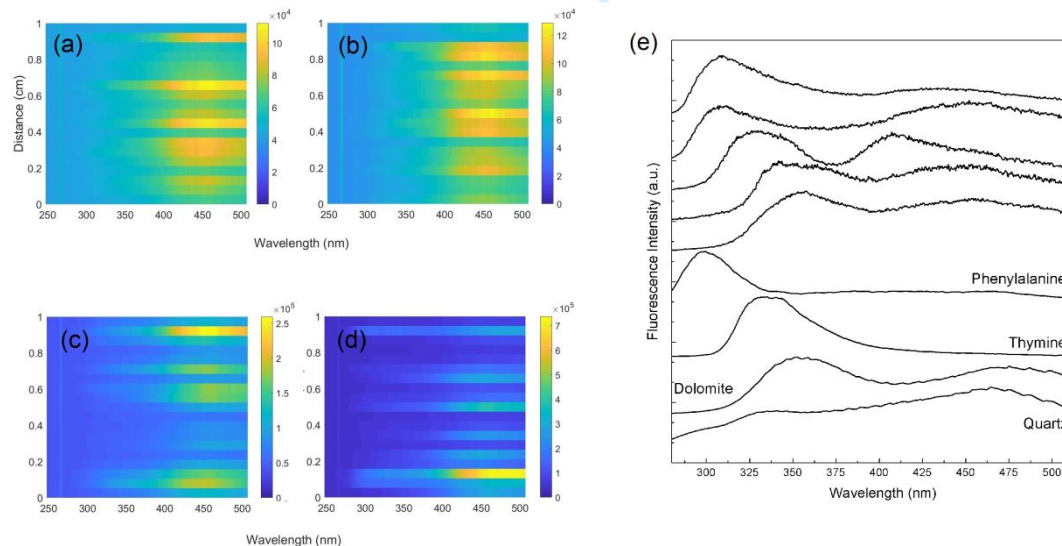


Figure 9. Representative LIF measurements (20 points along a 1 cm line) from the different sampling points from AMADEE-18: (a) 09-18; (b) 10-18; (c) 12-23; (d) 18-26; (e) selected spectral comparison with some organics and minerals

Table 4. Main LIF minealogical detection for the different sampling points from AMADEE-18 Mission (20-point mapping of 1 cm line). (** The carbonate and organic signatures were considered only for more than 15% of detection – more than 3 points)

	Samples													
	04-15	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Silicate (Quartz, Amorphous sand)	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Carbonates (Calcite or dolomite)	X	X	X					X	X	X	X	X	X	
Organic signature**	X	X		X	X	X	X	X			X	5	X	X

Scanning Electron Microscopy Analysis (SEM-EDS)

The results of the semi-quantitative EDS analyses are summarized in Table 5. The elemental analysis, representing bulk composition of the mixed powder samples was performed on areas of approximately 10 mm² on each pellet representing a qualitative approximation of the elemental distribution. The quantification of EDS spectra was performed with the P/B-ZAF QUANTAX analysis strategy (Bruker-Nano, 2011). This is a standard-less, self-calibrating spectrum analysis procedure that makes use of ZAF matrix correction formulas, enabling simple processing of the EDS spectra collected for each sample. Since the accuracy of EDS quantification is relatively low, especially with standard-less quantification, the normalized elemental mass percentages were used to allow comparison within and between samples for the relative elemental concentrations. With these quantifications, a qualitative analysis of the sample elemental composition was performed on the data. Figure 10 shows an example of the elemental distribution and BSE profile of a sample. Most samples appeared relatively homogeneous in BSE images, however, distinct mineral phase boundaries appeared in some samples, as evidenced in the BSE image and corresponding EDS spectral map of sample 05-17. The stark contrast seen between the left and right halves of both images are a result of large sodium chloride grains, interpreted as halite. The EDS spectra of the soil samples were more homogeneous across all samples, reflecting the random mixing involved in their preparation. The wt% oxide concentrations (rounded to the nearest percent to reflect the low accuracy of method) in Table 5 were calculated from the normalized mass percentages derived through the quantification procedure. All of the elemental iron present was converted to wt% FeO (see supporting data).

The spectra for some samples gave erroneous results, likely due to the large error associated with the quantification, and only the elements with identifiable peaks were quantified. Two poor quality spectra (4-15 and 8-18) have been omitted in Table 5.

Table 5. EDS elemental quantification converted to equivalent wt. % oxides for the different sampling points from the AMADEE-18 mission. Powder sample spectra collected over ~10 mm² acquisition areas and normalized.

oxide wt%	Samples												Average (oxide wt%)
	07-17	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26	
SiO ₂	24	41	33	16	28	32	27	12	22	31	28	28	27
Al ₂ O ₃	5	9	7	2	5	6	5	3	4	6	5	5	5
FeO	2	4	3	1	2	3	1	1	2	3	2	2	2
MgO	3	8	5	2	3	5	3	2	3	5	4	3	4
CaO	27	13	21	34	27	22	20	28	28	23	18	29	24
Na ₂ O	1	1	1	0	1	1	1	0	1	0	1	1	1
K ₂ O	1	1	1	1	1	1	1	1	1	2	1	1	1
CO ₂	37	23	29	44	33	30	42	53	39	30	41	31	39
Total	100	100	100	100	100	100	100	100	100	100	100	100	103

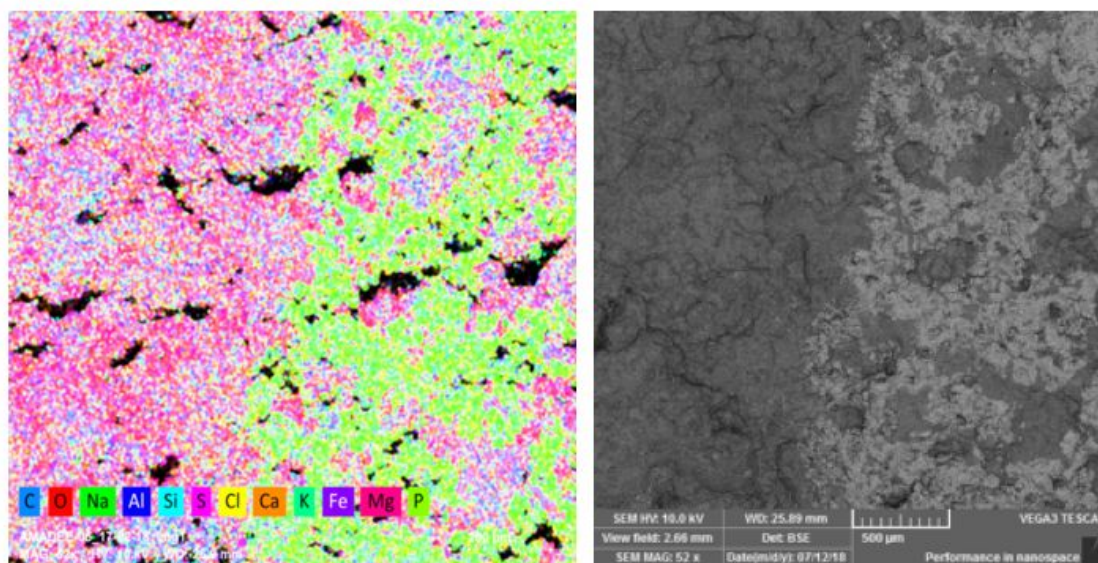


Figure 10. An EDS elemental map of 05-17 (right). The left half of the image is primarily composed of oxygen, silicon, and calcium whereas the right is dominated by sodium and chlorine. BSE image corresponding to area of EDS elemental map (left).

Laser Induced breakdown spectroscopy (LIBS)

LIBS spectra are shown in Figure 11. The spectral range covers almost all the near-UV, visible, and near-infrared. The main compositional elements are Al, Ca, Ti, Si, K, Fe, and Mg, were identified using the NIST Spectral Lines Database (National Institute of Standards and

Technology, 2016). In the near-UV region, the most intense peaks are Ca II located at 315.89, 317.93, 393.36 and 396.85 nm, Al I peaks at 308.31 nm, and Ti II within the 310-360 nm region, and 368.5 nm. Si II peaks are present at 385.6 nm and 390 nm approximately, as well as, Fe I peaks at 370.73 nm and 373.8 nm. The region from 480 to 660 nm contains Mg I (517.2 nm and 518.3 nm), Fe I and Fe II (526.6 nm, 527 nm, and 534.97 nm), Ca I (487.81 nm and 585.74 nm), and Na I (588.9 nm and 589.59 nm) characteristic lines. In the region of the near-infrared, the major peaks observed correspond to Ca I (714.8 nm, 720.2 nm, and 732.6 nm), K I (doublet at 766.49 and 769.89 nm), and O I (777.4 nm and 844.6 nm).

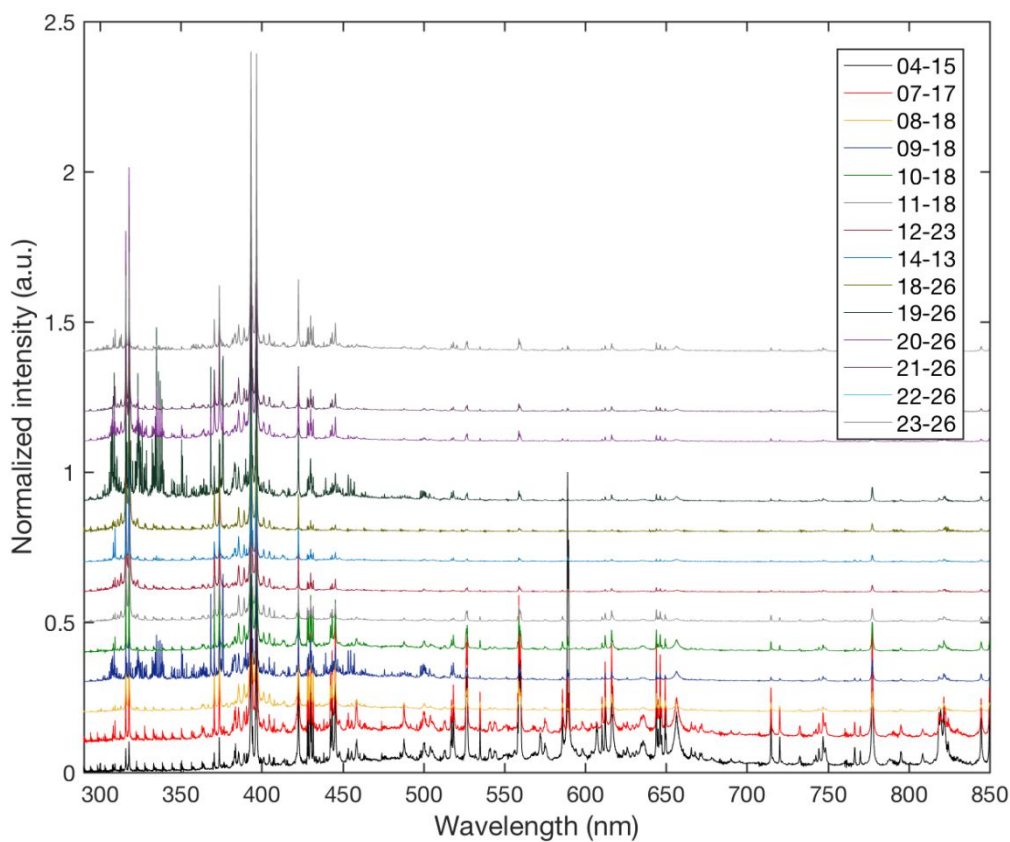


Figure 11. LIBS spectra from the pelletized AMADEE-18 sand samples.

Discussion

The petrological study of these samples shows evidence of hydrothermal deposits in the Ara Fm previously undescribed. This hydrothermalism is probably related to the syn-rift volcanism in the late stage of formation (Ramseyer *et al.*, 2013; Reuning *et al.*, 2007; Rollinson *et al.*, 2014). It is represented by the existence of the silica veinlets carrying sulfides and oxides intruding the dolomitic rocks studied in the present work. Part of this alteration has generated secondary clays

being observed in the present samples. Also, a supergene alteration has been observed on the samples reflected by an oxidized coating over the surface of almost all the clasts described. These alterations are supported by the presence of tertiary clays. In general, the supergene alteration was produced by the alluvial water intruding into the outcrops from the surface. The supergene process is combined with other alterations like the weathering and the alteration of the post-rift processes. All of the alteration processes described above are believed to have occurred on Mars. Because of this, this area has been proposed as a key terrestrial Mars analog for alteration processes (Chevrier and Mathé, 2007). The hydrothermalism and supergene alteration observed in the samples are comparable to similar geological data identified by the NASA MSL-Curiosity mission in Mars (Le Deit *et al.*, 2013; Marzo *et al.*, 2010; Popa *et al.*, 2015).

The spectroscopic analyses performed here detected minerals ranging from oxides to complex clays. The detected minerals match those observed using optical microscopy. The detected carbonates like dolomite and calcite correspond to the hydrothermal alteration. The oxides such as hematite and quartz are also significant in the context of understanding the hydrothermal processes. Also, minerals from old volcanism of the region such as pyroxene, feldspar, and olivine were detected. The meteoric alterations have also been detected in the form of clays and the water content on certain samples. The present results also confirm that Raman spectroscopy is able to detect certain mineral species that are not visually differentiable such as olenite, pyrope, or coesite minerals, among others. Also, the results agree with the mineralogy previously reported (Amthor *et al.*, 2005; Ramseier *et al.*, 2013). The other powerful application of the Raman analysis was the detection of organic materials on the different samples. However, the measurements must be complemented with the capabilities of other laser techniques to get semiquantitative concentrations of the organics present. Given the fast measurements and high sensitivity to target biomolecular structures that are present in the different samples, the best complementary method for detection of organics is LIF. The wavelength range for the detected organics match the range amino-acids like lysine or phenylalanine (Eshelman *et al.*, 2018, 2014). Also, the LIF method helped to confirm the existence of carbonates such as calcite or dolomite and quartz.

The XRD measurements identified and quantified the mineralization present in the different samples. Indeed, the pattern matching facilitated easy identification of quartz, feldspars and carbonates. The detection of some feldspars like albite helped to confirm the hydrothermal alterations. Other minerals (pyroxene and certain feldspars) confirmed the existence of past volcanic activities. The high concentration of carbonates in the analyzed samples was confirmed with LIF, LIBS and EDS. The high concentration of calcite and dolomite agree with the petrological description of the sedimentary rocks.

The LIBS technique mainly detected Ca, Mg, Al and other metal ions like Fe. These elements could be the cations in the identified carbonates (e.g. dolomite and calcite). However, Mg and Fe could also be associated with the volcanic parent mineralogy such as pyroxene, olivine or Fe-oxides (magnetite). This agrees with the Si detected on the samples which could represent the ionization of the pyroxene, olivine, and feldspars. Peaks corresponding to Al can be assigned to the different alkali-feldspar and/or clay mineralization present along all the different sampling points. The K and Na peaks probably belong to feldspar group minerals because this mineral group presents higher concentrations of K and Na compared to other minerals.

The semi-quantitative EDS analysis of the uncoated pellets detected high concentrations of Si, Ca, and C, which is consistent with the dominant mineralogy of calcite, dolomite, and quartz determined through the other experimental techniques applied to the samples. Throughout the laboratory analysis, samples were handled with gloves and stored in sealed containers to minimize contamination. Despite these efforts, it is likely that some of the C measured by EDS for the samples is due to contamination; however, high concentrations of ~40% indicate that contamination cannot be the only responsible mechanism, and that the presence of carbonates is probable. Possible contamination factors could include in-situ biological contamination, or sample handling at the Kepler base station during the mission. (Abed *et al.*, 2010) reported that the bacterial community in the region is mainly composed of cyanobacterium (e.g. Deltaproteobacteria, Bacteroidetes, Gemmatimonas and Planctomycetes). However, we have shown that EDS provided a relatively quick and semi-non-destructive mineralogical and geochemical analysis of samples, with elemental results consistent with those derived from the other instruments.

From an astrobiological perspective, carbonates are most commonly associated with potentially habitable zones such as hydrothermal areas, marine or lacustrine sediments, or biominerals (Bish *et al.*, 2013). The carbonates form not only through hydrothermal or diagenetic processes, but they can have an origin of weathering byproducts or from serpentinization. Carbonates provide an energy source for chemosynthetic microbes on Earth. Considering the magnesium-iron rich carbonates detected by the Spirit rover (Morris *et al.*, 2010), the sand studied during the AMADEE-18 mission is a good geological candidate for an in-situ instrument analysis of future space-qualified equipment prototypes and twin systems for future research.

The approach described in this section provides a geo-chemical instrumentation and methodological suite for the comprehensive analysis of a sample, yielding a mineralogical, elemental, organic, petrological, and geomorphological investigation. The results obtained from the different techniques illustrate the applicability of portable systems based on the real flight instruments for future analog missions (E. A. Lalla *et al.*, 2016; Sehlke *et al.*, 2019; Warren

Rhodes *et al.*, 2019). Further, the combined measurements of this comprehensive geo-chemical suite in analog space missions will yield insight into how our present methods can be improved upon for future missions to Mars and beyond. Particularly, given the large degree of instrument overlap and complementation, continuous use of the geo-chemical suite in space exploration will facilitate a unique combination in which the instruments ought to be used (Foing *et al.*, 2011; Sehlke *et al.*, 2019; Warren-Rhodes *et al.*, 2019).

Among the possible proposed objectives, the suite of used lab-techniques demonstrated that these analyzes provided us useful information when the “Selection of Instrumentation” comes. It should be noted, that the objective of this work and subsequent conclusions are not in the means by which the samples were collected and archived, nor in how the capability of these specific instruments in the terrestrial analogue compare to potential capabilities on Martian exploration. Rather, we strive to demonstrate how the synthesis of information derived from the aforementioned instruments (whose techniques are highly comparable to those in present and future missions) can be used to maximize mineralogical, petrological and astrobiological inference in a rigorous and efficient way. The information of capabilities from Table 1 combined with the “Data from orbiters and previous missions” could help us to decide an optimized workflow when it is required (see Figure 1). When the traverse plan is included in the workflow, this feature information could help to: (1) determine the logical orders of experiments according to the relevance of the acquired data; (2) duration of the experimental procedure (e.g. how much time and how many astronaut are required, time-lining, among others); and (3) risk assessments from landing site geomorphology, required sensing range, and energy duties (robotic and human).

As we have discussed over the course of the article, there are great mineralogical and astrobiological implications from studying terrestrial analogues, not only from the point of view of instrumentation validation, but also in systematizing methods of inference in the analogue context in addition to the setting of real space exploration. Indeed, data comes at many stages in space exploration; ranging from site selection, standardization of surrounding factors (such as passive reflectance), to the final mineralogical and astrobiological conclusions. Having accepted this, the only feasible way in which the utility of subsequent missions can be optimized is by having the integral processes and methods for inference rigorously established in the terrestrial domain. This is particularly true as we begin to travel with higher frequency beyond Mars.

Conclusion

1- The different techniques proposed facilitated the petrological, mineralogical and chemistry goals, chemical determination and organic detection for possible habitability within in the framework in Martian research. These analyses are presented to provide guidance for future analog missions like the AMADEE-18 mission. Also, the combined analytical methods will

improve the methods and protocols of in the detection of life for future space systems and Human missions to Mars.

2- The petrological analysis allowed us to determine the mineralogy, and geochemical origin of the minerals and alteration processes such as volcanism, hydrothermalism, and weathering among others, and to propose this area as a Mars Analog for future field simulated missions and instrumental experimentation for samples study.

3- The mineral identification obtained from Raman spectroscopy and XRD agree with the petrological analysis. The results show that the mineralogy of the different samples is dominated by the carbonates, silica (quartz) and feldspars.

4-The LIBS detection of Al, Ca, Ti, Si, K, Fe, Si and Mg and the subsequent elemental analysis obtained by SEM-EDS were in high agreement. The detected elements could be assigned to carbonates, clays, Fe-ores, and feldspar. These results give the understanding of the elemental distribution between mineral phases detected by Raman and XRD.

5-The LIF measurements detected a high concentration of carbonates as did the EDS, XRD, LIBS and Raman.

6-The detection of the organic samples was mainly achieved by Raman spectroscopy and LIF. Thus, the combined Raman-LIF measurements would provide the capabilities to identify organics on Mars.

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Supporting Data

The complete results of the measurements and results presented on the current research manuscript can be found online in the OeWF Multi-Mission Science Data Archive of AMADEE-18 Mission at <https://mission.oewf.org/archive/>

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Figure 3. Picture of field site showing the outcrops and alluvial fans deposits from where samples were taken and study in laboratory.

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Table 1. Comparison of the capabilities of the 5 systems employed in this investigation. (RLS=Raman Laser Spectrometer), (ChemCam=Chemistry and Camera), (Sherloc=Scanning Habitable Environments with Raman & Luminescence for Organics & Chemicals), (CheMin= Chemistry and Mineralogy), (MSL=Mars Science Laboratory)

Laboratory Technique	Mineral Identification	Elemental Composition	Organic detection	Geological Context	Laboratory instrumentation location in this study	Rover Instrument Equivalent
Raman Spectroscopy	Yes	Potentially (only major elements)	Yes	Yes (mapping)	Royal Ontario Museum	RLS (ExoMars) SuperCam (Mars 2020) Sherloc (Mars2020)
Micro- imaging	Yes	No	Yes	Yes	York University	CLUPI (ExoMars)
XRD	Yes	No	No	Potentially	Royal Ontario Museum	CheMin (MSL)
LIBS	No	Yes	Potentially	Yes (mapping)	York University	ChemCam (MSL) SuperCam (Mars 2020)
LIF	Potentially	No	Yes	No	York University	Sherloc (Mars 2020)
SEM-EDX	Yes (Imaging)	Yes (Qualitative)	Yes	No	York University	

Table 2. Micro-Raman mineral and other material detection on the different sampling points from AMADEE-18 Mission. X indicates that the mineral was found to be present in the respective sample.

	Samples														
Mineral	04-15	05-17	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Magnetite	X	X				X		X		X			X	X	X
Hematite						X		X	X	X	X			X	X
Goethite						X									
Anatase		X					X				X	X		X	
Chromite												X			
Quartz	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Calcite	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Dolomite		X						X					X		
Titanite						X									
Olenite												X			
Enstatite		X									X				
Diopside							X				X				
Epidote													X		X
Microcline				X				X						X	
Orthoclase													X		
Sanidine				X	X	X	X					X			
Albite		X			X	X		X	X		X	X	X		X
Andesine					X			X			X				
Labradorite	X														X
Clays						X									
Carbonaceous Material and organics	X	X	X	X	X	X	X	X	X	X	X	X	X	X	X

Table 3. XRD mineral detection on the different sampling points from AMADEE-18 Mission. X corresponds to the detected mineral in each collected sample.

Mineral	Samples													
	04-15	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Iron-Oxides					X									
Anatase	X												X	
Rutile	X												X	
Quartz		X	X	X	X	X	X		X	X	X	X	X	X
Calcite		X	X	X	X	X	X	X	X	X	X	X	X	X
Dolomite		X	X	X		X				X				
Ferrosilite								X				X		
Olenite											X			
Diopside	X													
Pigeonite							X							
Enstatite			X					X						
Epidote														X
Pyrope			X								X			
Coesite							X							
Olivine						X			X					
Feldspar									X					X
Anorthoclase													X	
Orthoclase				X										
Sanidine							X							
Anorthite					X					X				
Albite	X		X	X				X				X		
Bytownite											X			
Andesine	X	X												
Labradorite	X							X						

Table 4. Main LIF minealogical detection for the different sampling points from AMADEE-18 Mission (20-point mapping of 1 cm line). (** The carbonate and organic signatures were considered only for more than 15% of detection – more than 3 points

	Samples													
	04-15	07-17	08-18	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26
Silicate (Quartz, Amorphous sand)	X	X	X	X	X	X	X	X	X	X	X	X	X	X
Carbonates (Calcite or dolomite)	X	X	X					X	X	X	X	X	X	
Organic signature**	X	X		X	X	X	X	X			X	5	X	X

Table 5. EDS elemental quantification converted to equivalent wt. % oxides for the different sampling points from the AMADEE-18 mission. Powder sample spectra collected over ~10 mm² acquisition areas, and normalized.

	Samples												
oxide wt%	07-17	09-18	10-18	11-18	12-23	14-13	18-26	19-26	20-26	21-26	22-26	23-26	Average (oxide wt%)
SiO ₂	24	41	33	16	28	32	27	12	22	31	28	28	27
Al ₂ O ₃	5	9	7	2	5	6	5	3	4	6	5	5	5
FeO	2	4	3	1	2	3	1	1	2	3	2	2	2
MgO	3	8	5	2	3	5	3	2	3	5	4	3	4
CaO	27	13	21	34	27	22	20	28	28	23	18	29	24
Na ₂ O	1	1	1	0	1	1	1	0	1	0	1	1	1
K ₂ O	1	1	1	1	1	1	1	1	1	2	1	1	1
CO ₂	37	23	29	44	33	30	42	53	39	30	41	31	39
Total	100	100	100	100	100	100	100	100	100	100	100	100	103

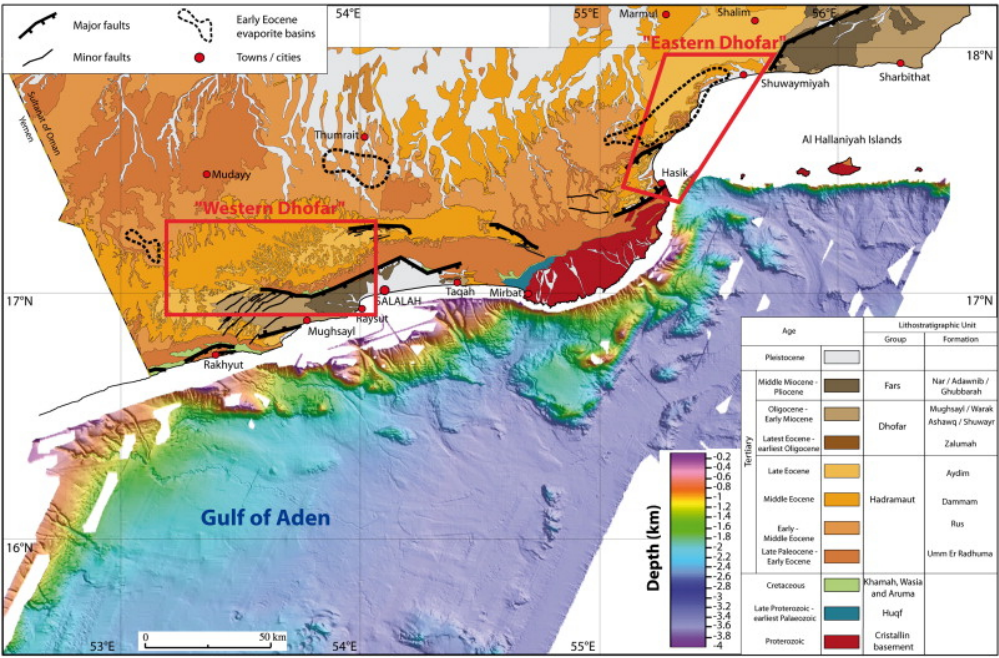


Figure 1

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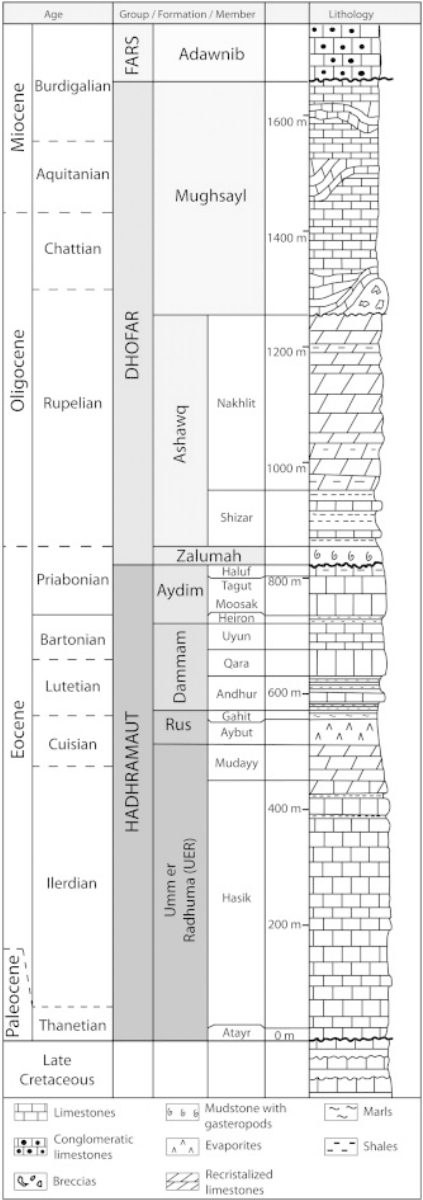


Figure 2

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Alluvial fans sediments

Ara Fm outcrops

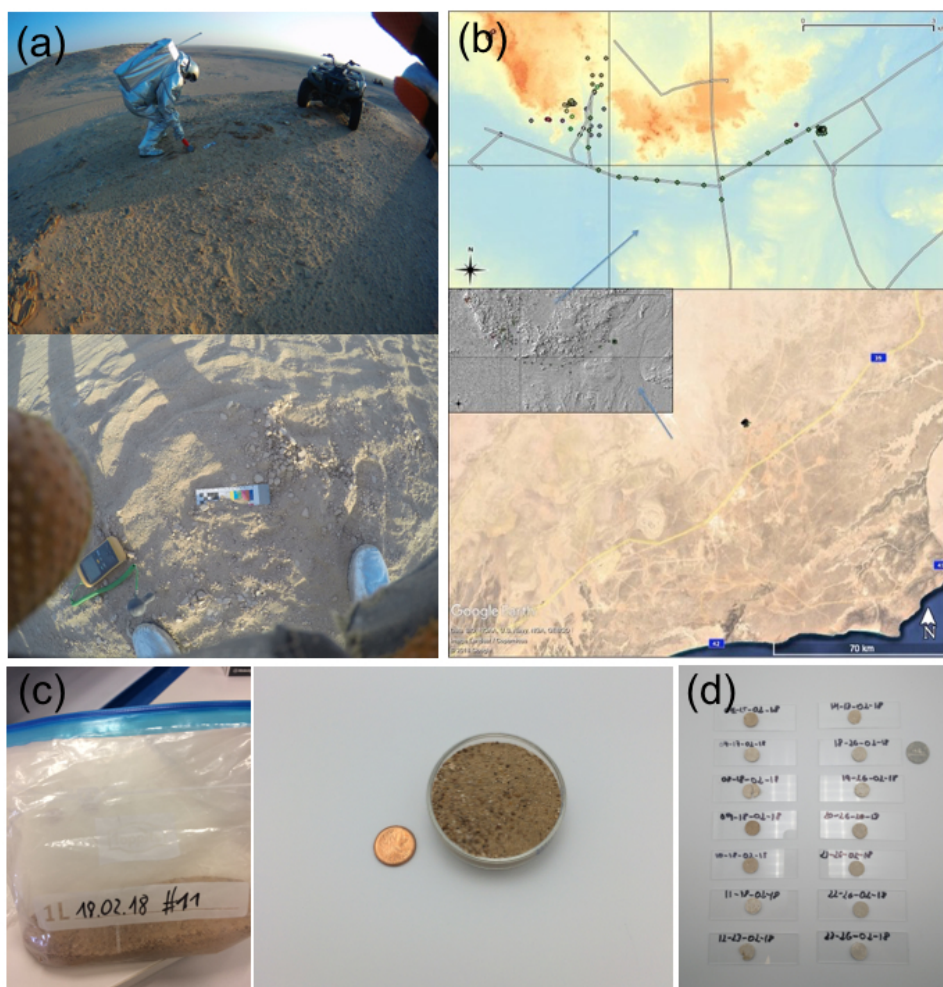


Figure 4

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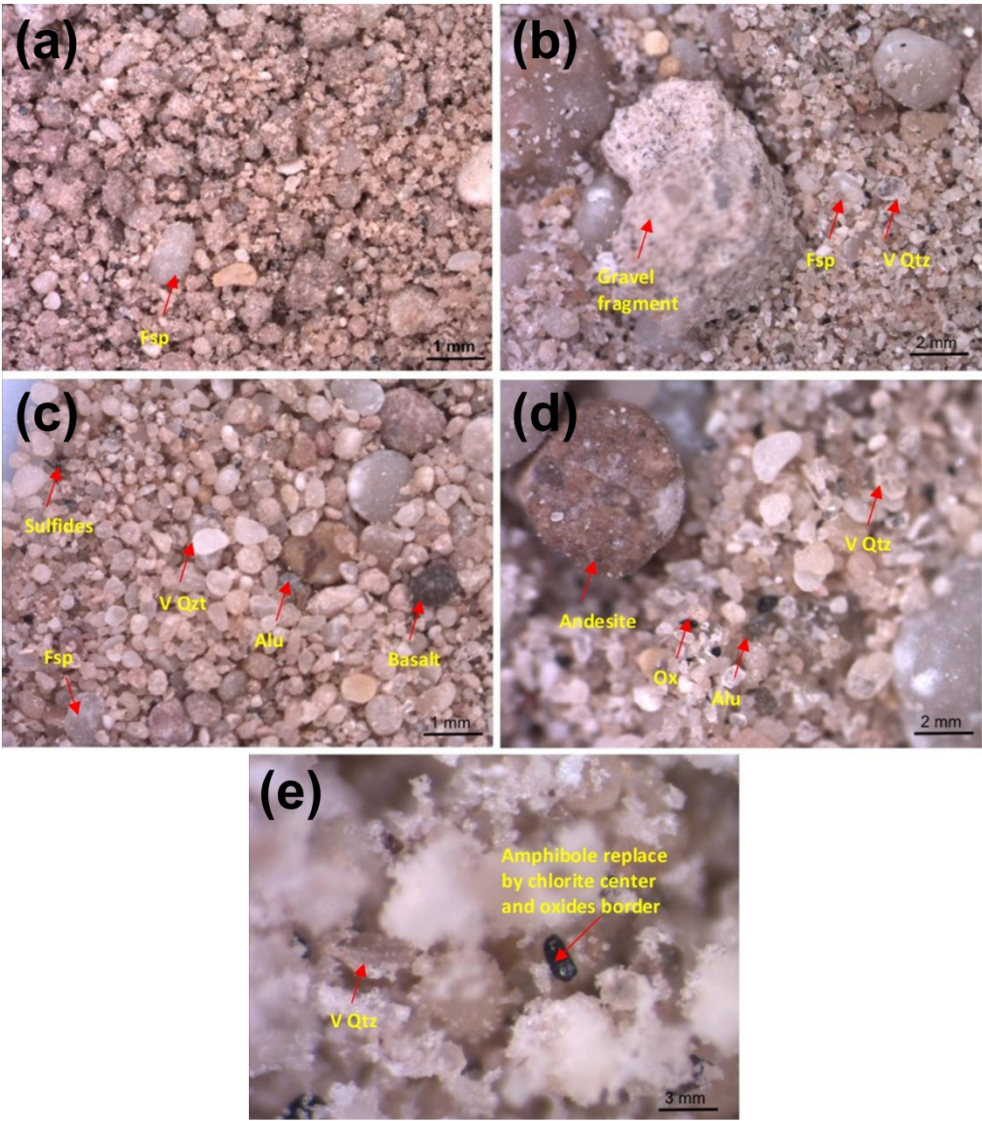


Figure 5

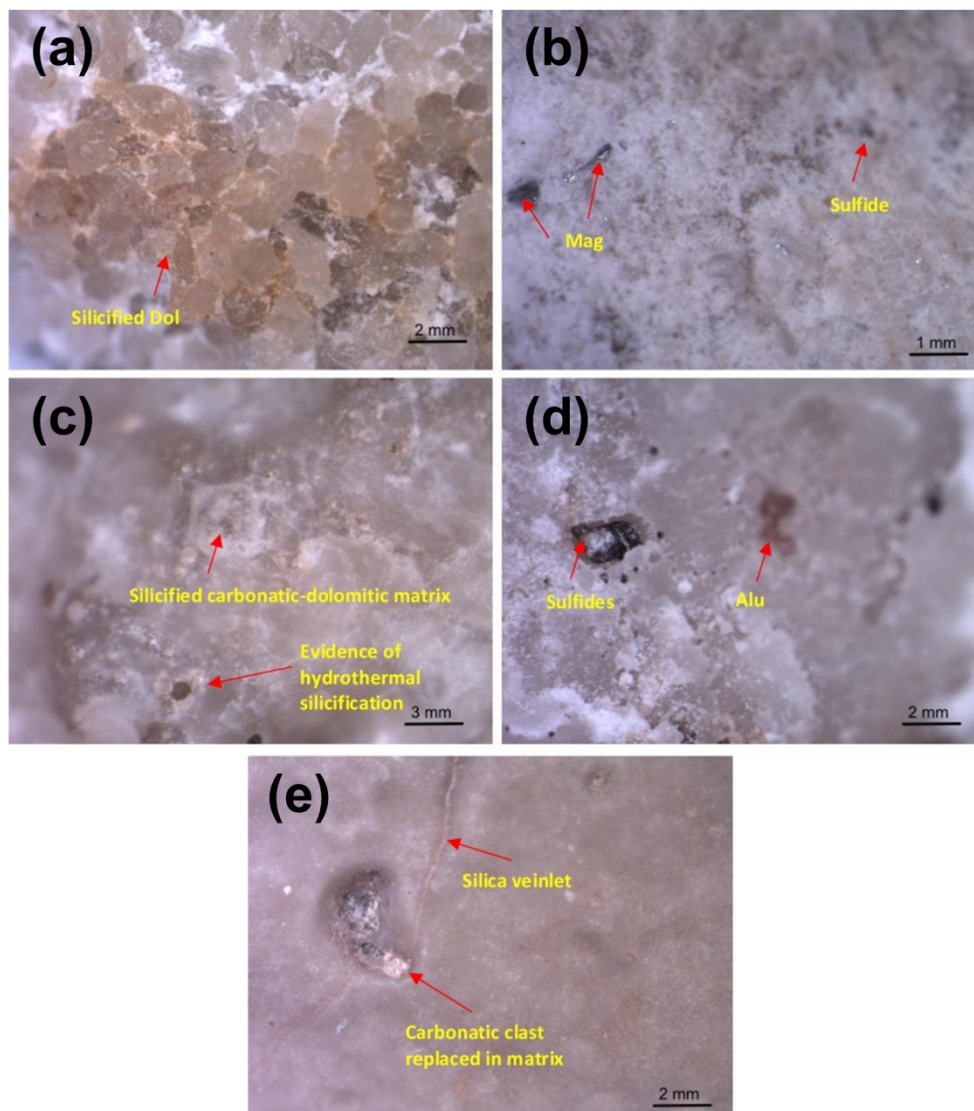


Figure 6

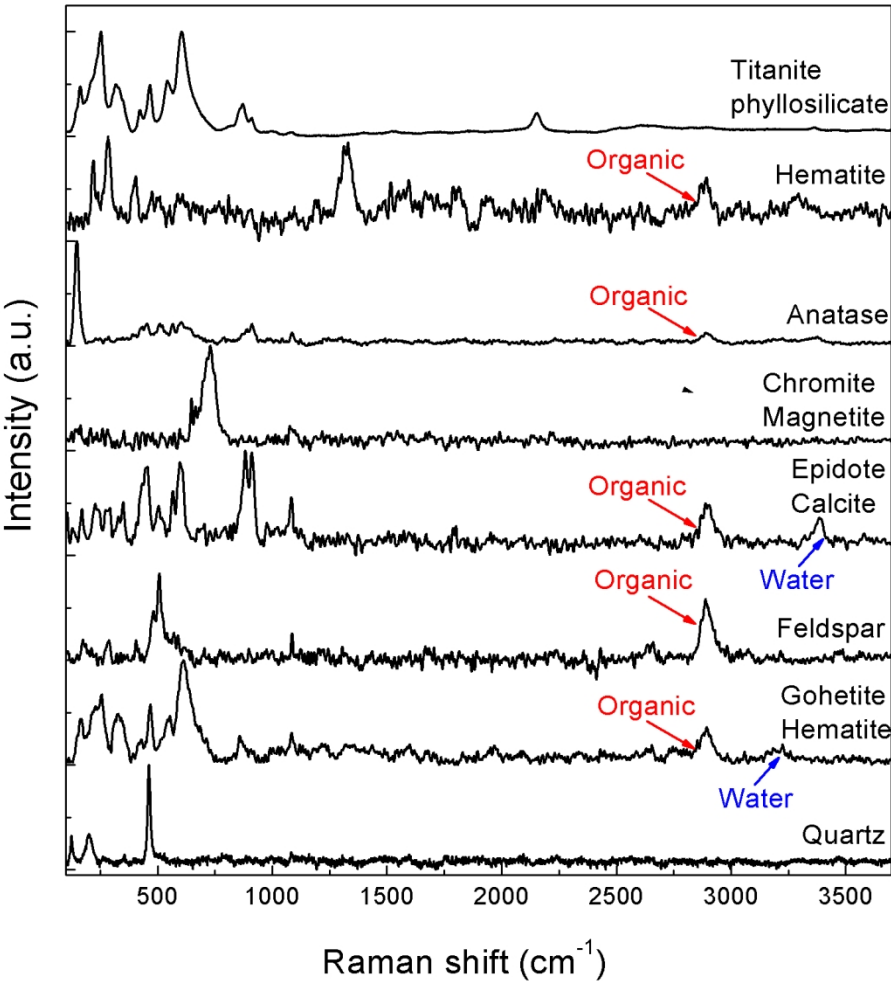


Figure 7

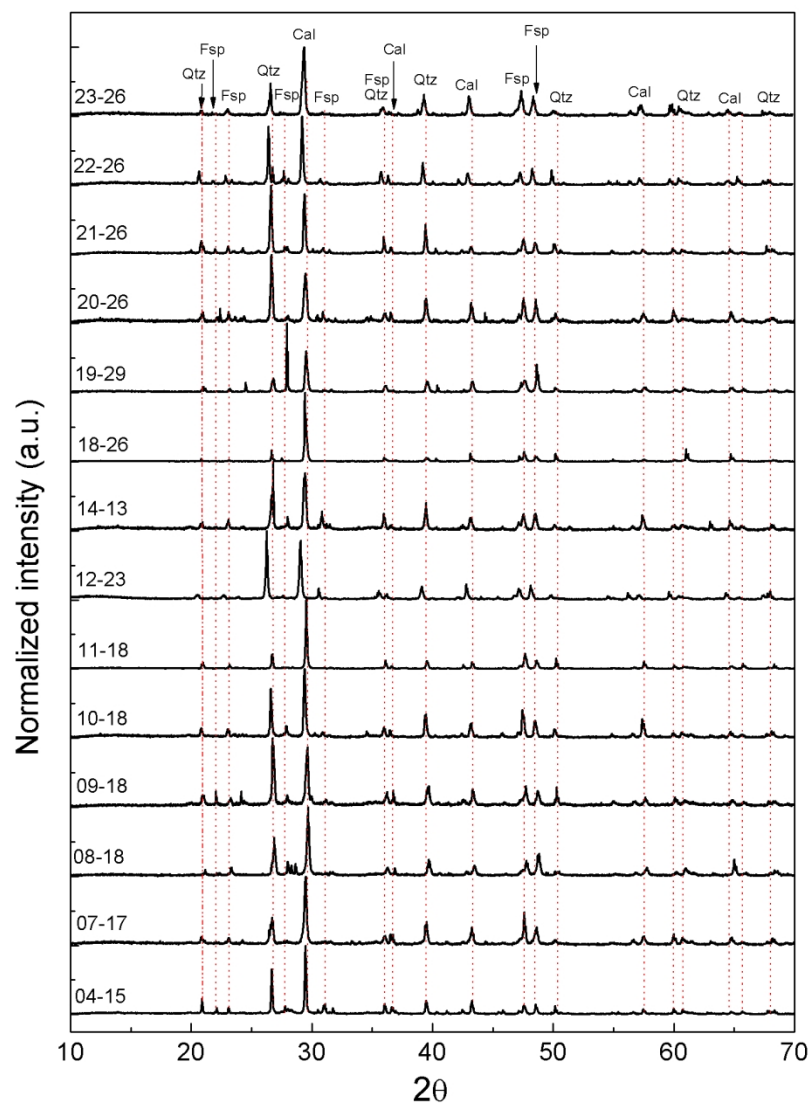


Figure 8

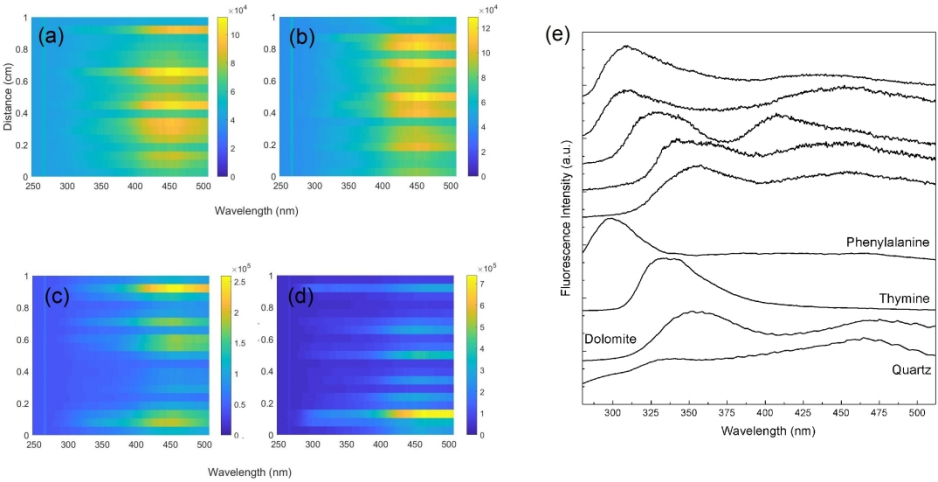


Figure 9

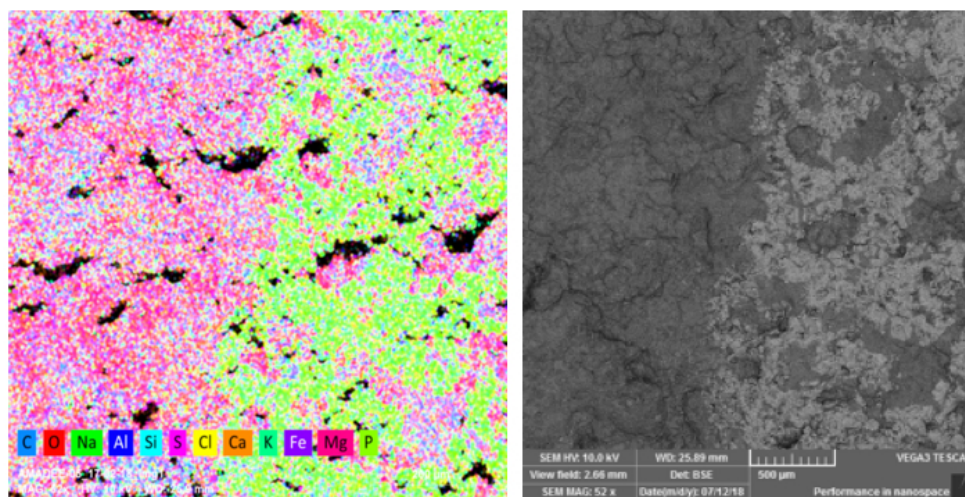


Figure 10

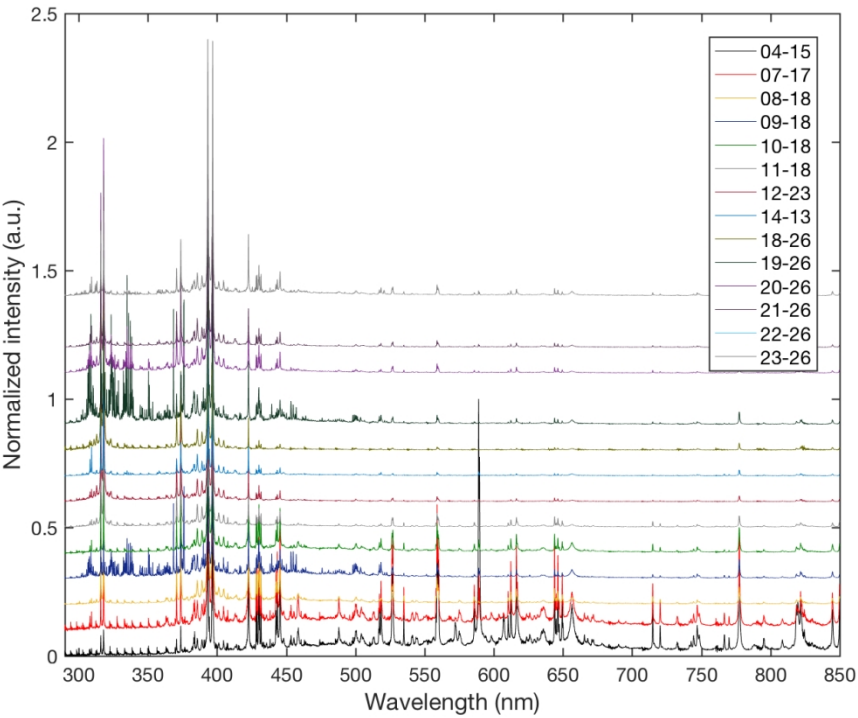


Figure 11

510x402mm (72 x 72 DPI)