cambridge.org/ija

Review Article

Cite this article: Westall F, Hickman-Lewis K, Cavalazzi B, Foucher F, Clodoré L, Vago JL (2021). On biosignatures for Mars. *International Journal of Astrobiology* **20**, 377–393. https:// doi.org/10.1017/S1473550421000264

Received: 4 June 2021 Revised: 1 October 2021 Accepted: 1 October 2021 First published online: 18 November 2021

Keywords:

Mars; ALH84001; biosignatures; carbonates; clays; salts; silica

Author for correspondence: Frances Westall, E-mail: frances.westall@cnrs-orleans.fr

© The Author(s), 2021. Published by Cambridge University Press. This is an Open Access article, distributed under the terms of the Creative Commons Attribution licence (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted re-use, distribution and reproduction, provided the original article is properly cited.



On biosignatures for Mars



Frances Westall¹, Keyron Hickman-Lewis^{1,2,3}, Barbara Cavalazzi^{1,2,4}, Frédéric Foucher¹, Laura Clodoré¹ and Jorge L. Vago⁵

¹CNRS-Centre de Biophysique Moléculaire, Orléans, France; ²University of Bologna, Bologna, Italy; ³Natural History Museum, London, UK; ⁴University of Johannesburgh, Johannesburgh, South Africa and ⁵European Space Agency ESTEC, Noordwijk, The Netherlands

Abstract

In this work, we address the difficulty of reliably identifying traces of life on Mars. Several independent lines of evidence are required to build a compelling body of proof. In particular, we underline the importance of correctly interpreting the geological and mineralogical context of the sites to be explored for the presence of biosignatures. We use as examples to illustrate this, ALH84001 (where knowledge of the geological context was very limited) and other terrestrial deposits, for which this could be properly established. We also discuss promising locations and formations to be explored by ongoing and future rover missions, including Oxia Planum, which, dated at 4.0 Ga, is the most ancient Mars location targeted for investigation yet.

Introduction

Since the seminal McKay et al. (1996) paper on possible traces of life in Martian meteorite ALH84001 (Fig. 1), we have come a long way in understanding the nature of biosignatures and their preservation, through experimentation, high-resolution analysis and Mars analogue studies. McKay et al. based their conclusions on three distinct phenomena that they considered to pertain to life: (1) globular and linear mineral features that were interpreted as nanobacteria (nanobacteria were all the 'rage' at that period following the works of Robert Folk; e.g. Folk and Lynch, 1997; Folk and Lynch, 2001); (2) the presence of polycyclic aromatic hydrocarbons (PAHs) interpreted as the remains of carbonaceous life forms; and (3) the presence of tiny magnetite crystals very similar to those produced by terrestrial magnetotactic bacteria. In addition, all of these features occurred in association with Fe/Mg globules deposited within fractures in the meteorites that they considered to have formed via low-temperature aqueous deposition, i.e. the cumulate volcanic rocks from which the meteorite was ejected had water percolating through their fractures, just as in terrestrial surface rocks. The temperature of deposition was later confirmed by Halevy et al. (2011) to be ~18 °C. Another important factor in the equation is the age of the meteorite: originally dated at \sim 4.5 Ga, subsequent investigations established an age of 4.09 Ga (Lapen et al., 2010), with the age of the carbonate globules dated to ~4 Ga (Borg et al., 1999). The geological and environmental contexts were consistent with the presence of life forms in water-filled crustal fractures. Thus, the McKay group based their interpretation on multiple types of biogenicity criteria - morphology, organic chemistry and biomineralization, a basic prerequisite in this type of study.

The original investigation was, at the time, state of the art. Lacking was sufficient understanding of the nature of primitive microorganisms and their processes of fossilization, which has come with subsequent research. It was later determined that the carbonate 'nanofossils' were mineralogical artefacts (Gibson et al., 2001). Isotopic analysis established that some of the organic molecules in the meteorite that fell on the Antarctic glacier were of terrestrial origin (Jull et al., 1998) but there was still the question of the origin of the PAHs. These kinds of organic molecules are generic and ubiquitous. PAHs occur in carbonaceous meteorites (Sephton, 2002) and they have been documented in Martian surface materials at Gale Crater by the instrument SAM on the NASA Mars Science Laboratory Curiosity rover (Eigenbrode et al., 2018). Steele et al. (2012) found evidence of mantle carbon in the ALH84001 meteorite, as well as in other Martian meteorites. Thus, abiotic processes could explain the presence of the PAHs. There has been more controversy over the nature of the tiny magnetite crystals, with the group of David McKay pushing the frontiers of analytical techniques to document the similarities between these crystals and those of magnetotactic microbes (Thomas-Keprta et al., 2009); however, others have demonstrated the formation of magnetite crystallites with similar 'biosignatures' (e.g. high crystallinity, chemical purity, a single-domain magnetic structure, and well-defined crystal morphology) through abiotic processes (Golden et al., 2004; Till et al., 2017). The salutary conclusion of these investigations is the degree of detail and stringency necessary to determine the all-important criterion of



Fig. 1. Characteristics of the ALH84001 meteorite and its interpreted biosignatures. (a) Pyroxenite, similar to the parent rock of the meteorite (at ARC Centre of Excellence for Core to Crust Fluid Systems). (b) The ALH84001 meteorite (NASA). (c) Rosettes of Fe carbonate (orange), Mg carbonate (white) and magnetite between the two (black) (NASA). (d) Globular-shaped carbonate structures interpreted to be nanobacteria (Gibson *et al.*, 2001; Elsevier, permission). (e) A tiny magnetite nanocrystal from the ALH84001 meteorite (Gibson *et al.*, 2001; Elsevier, permission). (f) Structure of phenanthrene, one of the organic compounds found in the meteorite (Wikipedia).

biogenicity. With the exception of the organic analyses, none of this would have been possible *in situ* on Mars.

Despite subsequent evidence showing that plausible non-biological processes can lead to the formation of each of the individual 'biosignatures' identified by McKay *et al.* (1996), this seminal paper kick-started the field of astrobiology. Today we have a far better understanding of microbes and how their traces can be preserved. We also have a deeper awareness of associated problems, such as the distinction between a microbial biosignature and a similar abiotic signature, or the necessity of demonstrating syngenicity, i.e. formation at the same time as the host rock. Importantly, since the time of the McKay study, many new techniques have been developed, or older ones adapted, to characterize biosignatures.

The discovery of traces of endogenous life in Martian rocks, either *in situ* or in returned samples, would have a profound effect on our view of the distribution of life in the Universe: it would suggest that the processes leading to the emergence of life may be common in the Solar System and, by extension, on rocky planets elsewhere. Importantly, we will want to understand more about the ecology of these life forms. For example, whether they lived at the time of the formation of the rock or whether they were later colonizers, for instance of pores and fractures. We will also want to learn about what kinds of metabolisms they used. In the following, we will define the term 'biosignature', emphasizing the importance of geological and mineralogical context, as well as the fossilization and preservation of traces of life. Taking the example of the Alan Hills 84001 meteorite as a starting point, we will make a brief overview of preserved traces of life in different types of terrestrial sediments and mineral matrices of relevance for Mars. We will consider examples of lithologies and minerals detected on Mars, especially with consideration to the Mars 2020 mission to Jezero Crater on Mars that will take samples to return to Earth (Williford *et al.*, 2018) and the ExoMars 2022 mission that will be searching for traces of life on Oxia Planum (Vago *et al.*, 2015, 2017).

Biosignatures

Biosignature definition

We define biosignatures as morphological, chemical (organic, elemental and/or mineral) and isotopic traces of organisms preserved in minerals, sediments and rocks. They represent the physical presence of the organisms as well as evidence of their metabolic activities and metabolites (Westall and Cavalazzi, 2011 within put from Summons *et al.*, 2011; Mustard *et al.*, 2013; Hays *et al.*, 2017; Vago *et al.*, 2017). Firstly, we aim to recognize the morphological expression of microorganisms, including mineral-replaced cells, colonies or microbial biofilms/ mats, as well as the physical expression of the interaction of microorganisms with their sedimentary environment. Examples of this are three-dimensional microbial constructions, including microbially induced sedimentary structures (MISS, Noffke et al., 2008; Heubeck, 2009; Noffke, 2009, 2010) and stromatolites (Byerly et al., 1986; Hofmann and Davidson, 1998; Allwood et al., 2006). The organic remains of microorganisms and their colonies may still be identifiable as such, if preserved under anoxic conditions, unless they have been subjected to such a high degree of metamorphism that the organic matter becomes graphitized (although carbon isotopic signatures may still be preserved in graphite). During diagenesis and metamorphism, organic molecules break up, the lower molecular weight portions degrading first leaving relics of more resistant components, such as lipids and recalcitrant biopolymers (Brocks and Summons, 2003). Factors influencing the composition and structure of the more recalcitrant molecules during diagenesis include oxidation, reduction, sulphurization, desulphurization and molecular rearrangement reactions. The loss of functional groups results in products with more robust aromatic and aliphatic structures that may still be identifiable as microbial hydrocarbons. Thus, the structure and composition of the degraded organic molecules can be preserved, even up to lower greenschist facies metamorphism (and occasionally even beyond if the signature is encased in a resistant mineral; Hassenkam et al., 2017). This preservation includes their enantiomeric signature, patterns in the odd over even carbon number, δ^{13} C signature, diastereoisomeric preference, structural isomer preference and repeating constitutional subunits (Vago et al., 2017). These characteristics, even for relatively degraded molecules, can aid in the interpretation of biogenicity. Fortunately, given the smaller size and lower gravity of Mars, the degree of alteration of organic molecules due to metamorphism on the planet is expected to be very low, unless the rocks are affected by local impact metamorphism. It may therefore be possible to detect biomarker molecules, i.e. molecules whose structure and composition can be traced back to specific microbial precursors more easily on Mars than on Earth. To date, the oldest documented biomarkers occur in the 1.73 Ga Wollogorang Formation in the southern McArthur Basin, Australia (Vinnichenko et al., 2020). They comprise degradation products, such as C14-C19 2,3,4- and 2,3,6-trimethyl aryl isoprenoids. The latter can be correlated to carotenoids of cyanobacteria and/or green sulphur bacteria (Chlorobiaceae), while the former 2,3,4-AI series may be derived from purple sulphur bacteria (Chromatiaceae). With respect to extraterrestrial life, it is not the molecules themselves, but specific patterns that can be diagnostic, even, without knowing the nature of Martian life (if it exists), i.e. agnostic biosignatures.

Additional biogenicity signatures include those resulting from microbial metabolism, for example, the fractionation of certain isotopes, such as carbon, nitrogen or iron (Schidlowski, 2001; Teng *et al.*, 2017), as well as the concentration of bio-essential elements C, H, N, O, P, S (e.g. Hickman-Lewis *et al.*, 2016; Delarue *et al.*, 2020), and transition metals Fe, Ni, Cu, Mn, Co, Mo, V and As (Wogelius *et al.*, 2011; Hickman-Lewis *et al.*, 2020*a*). Also resulting from metabolic activity are certain biominerals precipitated either passively or actively, including various carbonate minerals, or magnetite. Finally, the presence of microbes and their metabolic products strongly influences mineral composition, habit, dissolution and even size (Banfield *et al.*, 2001).

The Allan Hills meteorite 84001 provides a salutary example of the pitfalls regarding the identification of potential biosignatures. Many of the characteristics listed above can result from abiogenic processes (Westall and Cavalazzi, 2011; McMahon *et al.*, 2018). Particularly with respect to possible extraterrestrial life, a multiscale, multisignature approach is favoured to enhance confidence in interpretations of biogenicity. While the above characteristics are related to life as we know it, the recently developed concept of agnostic biosignatures seeks to address the signatures of life as we do not know it. This concept concerns a broader definition of life based on activity rather than structure and is not reliant on specific biogeochemistry (Johnson *et al.*, 2018*a*, 2018*b*), concerning instead an increase in the degree of complexity of organic molecules.

The importance of geological and mineralogical context

Another aspect of biosignature study demonstrated by the ALH84001 example is that of geological and mineralogical context. The overall geological context of the meteorite is not a traditional one for life. Canonically, we expect to find traces of life associated with abundant water and, more specifically, detrital and/or chemical sediments deposited by water. This is because most primitive life forms (prokaryotes) are found in such environments. Exceptions are more evolved prokaryotes, such as cyanobacteria, that can survive in very dry environments, such as endoliths in hot or cold deserts. Indeed, Imre Friedmann, the renowned microbiologist who first identified endolithic life in the supposedly uninhabited Dry Valleys of Antarctica in the 1980s, was the first to suggest that, before disappearing from the surface of Mars altogether, Martian life, if it existed, would have adopted such endolithic habitats (Friedmann and Koriem, 1989).

In the case of the Allan Hills meteorite, the geological context is on a micro-scale, rather than a macro-scale. It is a cumulate rock, i.e. a relatively coarse-grained basaltic rock that is formed at the base of a thick lava flow. When extruded under water, the surfaces of lava flows are rapidly colonized by microorganisms as soon as they have cooled down, since organisms use them as a support for colonization, as well as a substrate from which to obtain energy through redox reactions at the water/rock interface and essential transition metals extracted from the rock (Thorseth et al., 1995; Einen et al., 2006). Pores formed by gas bubbles or amygdules in the surfaces of basalt may also be colonized by microbes (Cavalazzi et al., 2011, 2012). However, it is only in the last couple of decades that we have been made aware of a vast subsurface biosphere in fractures in the terrestrial crust. Deep gold mines in South Africa showed that the subsurface down to more than 3.5 km is an important habitat where surface water trickling through fractures fuels a relatively wide array of microbial life (Hoehler and Jørgensen, 2013; Onstott, 2016). These groundwaters pick up a variety of nutrients depending on the rocks through which they flow, thus producing a distinctive fracture fluid chemistry reflecting that of the surrounding rock. They are sources of energy and nutrients for subsurface microbial life and also transport microorganisms through the subsurface (Casar et al., 2021). Indeed, subsurface experiments show that iron-bearing minerals, such as pyrite and siderite, are prime sites for microbial colonization because of the high amount of energy released during redox reactions (Casar et al., 2020). It is estimated that the continental deep subsurface is probably one of the largest reservoirs of bacteria and archaea on Earth, where



Fig. 2. Potential subsurface habitats on Mars (Onstott et al., 2019; Liebert, permission).

microorganisms survive on minimal organic carbon sources, obtaining their energy instead from mineral sources (McMahon and Parnell, 2014; Casar *et al.*, 2021). If life ever appeared on Mars, and if it still exists, extant life will most likely be found in the subsurface of the planet (Fig. 2; Onstott *et al.*, 2019).

The ALH84001 meteorite is petrographically classified as a coarse-grained orthopyroxenite comprising 97% orthopyroxene, 2% chromite, with interstitial plagioclase, augite, olivine, pyrite, apatite and silica, as well as maskelynite formed during impact shock and later secondary Fe-Mg-Mn-Ca carbonate (Harvey and McSween Jr, 1994; Mittlefehldt, 1994). Orthopyroxenes are Fe- and Mg-rich chain silicates while chromites are Fe- and Cr-rich oxides. Olivine, augite and pyrite are also Fe-rich minerals, as is a fraction of the secondary carbonate deposits. By virtue of this association of minerals, groundwaters flowing through the fractures of the consolidated basaltic flow would have derived sufficient nutrients (energy sources) to support a subsurface microbial biome. Carbon would have been limited, likely brought in from the surface along with the ground waters. Note that although magmatic carbon occurs in this rock as PAH-containing macromolecular carbon (Steele et al., 2012), such materials are generally insoluble organic carbon phases. Nevertheless, microbial utilization of mantle-derived carbon does occur (Lang et al., 2012), but it is not certain whether microbes can directly utilize this carbon or whether it first needs to be oxidized by the microbes into smaller, more assimilable phases. As indicated in the above discussion of subsurface life in the Earth's crust, a priori it is perfectly feasible that cracks in the Martian crust (or lava flows) could have supported microbial activity.

Had McKay *et al.*'s (1996) hypothesis for evidence of life in ALH84001 been confirmed, the geological and mineralogical context would have allowed us to draw further conclusions as to the mode of life and metabolic activity of the microorganisms: they could have been either (1) anaerobic chemotrophs, more precisely lithotrophs possibly cycling iron and sulphur, or (2) other types of organisms (e.g. heterotrophs) flushed into the fractures during surface water flow. Further refinements would not be possible since we know nothing about putative Martian life.

This example demonstrates the importance of geological and mineralogical context for supporting hypothetical biosignature identification and interpretation and for obtaining environmental details that provide some indication as to the lifestyle of the potential organism. Of additional importance is the information that geological and mineralogical context can provide for understanding the potential for fossilization and long-term preservation of biosignatures. Here, again, the ALH84001 meteorite is a salutary example.

Fossilization and preservation of traces of life

We saw above that biosignatures may be organic, organo-mineral, morphological, isotopic or geochemical. However, in order to be preserved over long periods of geological time, any signature of life needs to be encased or cemented within its mineralogical context and to have survived various episodes of chemical and physical destruction, including metamorphism. Destruction could be related to percolating groundwaters or hydrothermal fluids or physical destruction due to mass wasting, erosion or impacts. While the smaller size of Mars and the lack of plate tectonics precludes the kinds of metamorphic alteration associated with rocks buried to similar depths on Earth, the numerous impacts that have marked its surface over the last >4 Ga will have left significant shock effects as well as regions of total destruction. Study of the ALH84001 meteorite shows that it was sourced from the coarse-grained, fractionated base of a consolidated melt (flow or sill) and was affected subsequently by a variety of processes on Mars. After consolidation of the flow, fractures and shock features in the rock indicate that the host formation was impacted. The rock was subjected to a high-temperature metamorphism, possibly related to the original impact (Treiman, 1995). Following this impact and metamorphism, groundwaters penetrated through the fractures, depositing secondary Fe-Mg-Mn carbonates along the walls of the fractures. A later shock event affected the whole rock and ejected a portion into space, which landed on the Antarctic ice cap about 13 000 years ago after >16My in space.

The processes of fossilization of microorganisms have been detailed previously in a number of publications (Westall *et al.*, 2015*a*, 2018; Westall and Hickman-Lewis, 2018). Briefly, microorganisms can be preserved as physical structures, i.e. as cells, colonies and biofilms, if they are rapidly coated and encased in a mineral cement (Fig. 3). This needs to occur during the life time of a microbe or colony otherwise its physical structure very rapidly degrades. Orange *et al.* (2009) noted that the chelation of minerals to a cell surface occurs within 24 h after which purely chemical polymerization takes place creating a crust of various structures and thicknesses, depending on the cell wall

Fig. 3. (a) Calcified bacteria, SEM micrograph (Brunelli *et al.*, 2012; SAGE journals, permission). (b) Silicified bacteria, TEM micrograph of experimental fossilisation (Westall *et al.*, 1995; The Palaeontological Association, permission). (c) Fossilized microorganism with a thick silica crust (black arrow) and the mould of a microorganism outlined by phyllosilicates (white arrow), TEM micrograph of experimental fossilisation (Westall *et al.*, 1995; The Palaeontological Association, permission). (d) 1 Ga old eukaryote cell preserved by external Fe clays (c) and internal K clays (d) (Wacey *et al.*, 2014; Sci. Rep., permission). (e) Embedding of gypsum crystals on the outside and inside of an experimentally fossilized microorganism (Yersinia), TEM micrograph (Gaboyer *et al.*, 2017; Sci. Rep., permission). (f) Algal cells embedded in a modern halite crystal (left) and in a 150 Ka halite crystal (Sankaranarayanan *et al.*, 2011; Open Commons, permission).

structure (Westall, 1997; Orange et al., 2009). The moulds of former cells can also be preserved in a mineral matrix (e.g. Liebig et al., 1996; Homann et al., 2015). Physical microbial constructions, such as those produced by phototrophic microbial mats (e.g. stromatolites, MISS and other microbialites) are preserved by in situ mineralization of organic matter (EPS, cells and trichomes) and early cementation of host sediments (Fig. 4; e.g. Hofmann et al., 1999; Noffke et al., 2008). In reducing conditions, the degraded organic components of fossilized structures are encased by the surrounding mineral matrix. Oxidizing conditions, on the other hand, will destroy them. Diverse organo-geochemical signatures may be associated with either remnant organic matter in the structures or trapped within the embedding mineral matrix. For example, the silicified remains of organically preserved anaerobic phototrophic biofilms and microbial mats in Palaeoarchaean, Mars-analogue sediments from 3.3-3.5 billion-year-old fossiliferous horizons in the Barberton Greenstone Belt in South Africa, reveal organic biosignatures in the form of kerogen distributions (Westall et al., 2006b, 2011), molecular structure (e.g. CH₃/CH₂ ratios) and δ^{13} C ratios (Hickman-Lewis et al., 2020a, 2020c), as well as enhanced transition metal concentrations indicative of microbial origin (Hickman-Lewis et al., 2020a). In general, the physical remains of cells, colonies and mats are rarely wellpreserved; it is more common to find poorly preserved remains or simply generic organic matter coming from the degraded cells that are trapped in a mineral cement that remains (Walsh and Lowe, 1999; van Zuilen *et al.*, 2007).

Subtler indications of microbial presence, such as their effects on mineral growth, corrosion features, or even biominerals, are very difficult to identify without additional evidence in the rock record. Of relevance for Mars, evidence of microbial corrosion tunnels in basaltic pillow lava surfaces from the Palaeoarchaean of the Barberton Greenstone Belt has been hotly contested (Furnes et al., 2004; McLoughlin et al., 2010). Furnes et al. (2004) compared microtextures comprising radiating filaments of microcrystalline titanite in the rims of ancient pillow basalts to similar features occurring during modern microbial corrosion of volcanic glasses (Furnes et al., 2007). Nevertheless, these particular tunnel-like features have also been interpreted as abiogenic (McLoughlin et al., 2010; Lepot et al., 2011). For example, they have been reported to lack the organic lining that would be expected were they of microbial origin (McLoughlin et al., 2012). In contrast, Foucher et al. (2010) document fossilized (silicified) polymeric substances infilling tunnels corroded into the surfaces of volcanic glass particles from the 3.45 Ga Kitty's Gap Chert in the Pilbara Greenstone Belt, Australia. In this



Fig. 4. (a) Cold seeps from the Arctic (Williscroft *et al.*, 2017; GSA, permission). (b) Moulds of rod-shaped microbes in microbially-precipitated carbonate from a limestone cave (Cacchio *et al.*, 2003; Taylor and Francis, permission).

case, the direct association of organic polymer within tunnels at the surface of the volcanic glass, plus associated fossilized cells and colonies at the surfaces of the volcanic clasts (Westall *et al.*, 2006a, 2011), all later entombed by a penecontemporaneous cement of silica, is a strong indication of corrosion due to microbial metabolic processes.

Importantly, especially when considering the ancient terrestrial rock record or the search for life on Mars, several types of biosignature needs to be present to increase the probability of a reliable interpretation. This multi-signature approach was attempted by McKay *et al.* (1996); however, the resulting interpretations were incorrect because of the lack of understanding of biosignatures at that time.

Carbonates

(1) Carbonates in subsurface environments: The carbonates in ALH84001 formed in a subsurface environment, in this case in an igneous rock. The McKay team had identified globular features and lineations of such features in the carbonate deposits of ALH84001 that they interpreted as nanobacteria. Carbonates are a mineral phase whose formation may be mediated by microbial metabolisms (Riding, 1991; Spadafora et al., 2010), often preserving physical biosignatures, such as stromatolites as well as, occasionally, individual microbial cells (e.g. Fig. 3a; Perri et al., 2012). Biomineralization may be a passive process in which a mineral forms as a result of microbial metabolic processes affecting its immediate environment, for example, changing pH, or the production of metabolites that then combine with various ions in solution. It may also be actively induced, for instance, when bacteria control the Ca²⁺ content in their cells by using a crossmembrane proton pump to eliminate excess ions. The expelled Ca^{2+} then combines with CO_3^- in solution to precipitate around the cell walls (Bosak and Newman, 2003). In the subsurface, carbonate deposits can form naturally in certain carbonate cave systems (Fig. 4b; Cacchio et al., 2003), as well as from drip water in mines (García et al., 2016). These environments are aerobic to microaerophilic. Fossilization of the bacterium Acinetobacter gyllenbergii, isolated from Columbian mines, was demonstrated experimentally with the precipitation of carbonate around cell walls (García et al., 2016). Subsurface carbonates also form in non-carbonate rock lithologies. For example, carbonates in fractures in granitic host rock formed as a result of microbial oxidation of methane (Drake et al., 2015). The latter preserves the most extreme ¹³C depletion recorded to date (-125%) as a result of the incorporation of C from biogenic methane in this energy-limited and C-poor environment. The carbonates also host lipid-derived organic components emanating from a microbial consortium comprising anaerobic methane-oxidizing archaea and sulphate-reducing bacteria (SRB). Note, however, that the biosignatures are confined to carbon-bearing mineral phases (clays) occurring at the boundaries of the calcite grains, not within the carbonate grains themselves.

In the case of the ALH84001 meteorite, it would in theory be possible for microorganisms to mediate carbonate precipitation, but it seems more likely that such processes are limited to heterotrophs. As these organisms obtain their energy and nutrients from the degradation of pre-existing carbon, one could hypothesize that such microorganisms were washed into fractures in lava flows at the surface of Mars. However, in a subsurface, oligotrophic environment without a primary carbon source, it is questionable as to whether the microorganisms would have been able to metabolize and produce a carbonate deposit. Nevertheless, similar to the precipitates in the ALH84001 meteorite, Benzerara et al. (2003) described nano-sized bacteriomorph crystallites of carbonate in a terrestrial weathering phase of pyroxenes in the Tatahouine meteorite, in this case associated with terrestrial microbes. They concluded that, while the formation of the crystallites was microbially-induced, the structures in themselves were chemical precipitates, i.e. abiotic growth, not fossilized microbes.

(2) Surface carbonate deposits: The majority of known microbially-mediated carbonate formation occurs in subaerial environments, in particular in relationship with phototrophic microbial mats resulting in tabular or three-dimensional stromatolitic structures (Warthmann et al., 2000; Dupraz et al., 2009; Bosak et al., 2013). While it is unclear whether Martian life could have reached the evolutionary state of phototrophy owing to the phenomenon of punctuated habitability (Westall et al., 2015a), we cannot completely discount this possibility. Therefore, a biosignature of particular relevance for anaerobic early Mars is the microbial biomineralization of carbonate mediated by consortia of anaerobic phototrophs and their accompanying heterotrophic degraders, such as SRB. In this case, SRB activity creates the chemical conditions for the precipitation of carbonate by increasing alkalinity in the environment such that Ca²⁺ ions released from microbial extracellular polymeric substances (EPS), an important component of phototrophic biofilms and mats, react with the CO²⁻ dissolved in the aqueous phase to form CaCO₃. SRBs in such consortia precipitating nanocrystals of aragonite and CaSO₄ within degraded organic matter were documented from Mars-analogue, shallow water volcanoclastic sediments dated to 3.33 Ga from the Barberton Greenstone Belt (Westall et al., 2006a, 2011).

Substantial carbonate precipitations in the form of mounds occur in association with cold methane seeps, as well as white smoker vents (Fig. 4a). Methane seeps are widespread and are found in the terrestrial rock record throughout the Phanerozoic (Barbieri and Cavalazzi, 2008). They host anaerobic chemosynthetic communities that oxidize methane and sulphide to obtain energy, using dissolved CO_2 as their carbon source. An increase of pH value driven by their metabolisms leads to the precipitation of calcium carbonate, as in the phototrophic mat communities described above. Although carbonate is formed by these processes, it is rare for individual cells to be preserved although fossil microbial filaments and mats have been documented within cold seep carbonate deposits of Devonian age from Morocco (Cavalazzi, 2007; Cavalazzi *et al.*, 2007).

(3) Carbonates on Mars: Spectral analysis from orbit of the surface of Mars has detected carbonates associated with aqueous weathering profiles across the planet's Noachian terranes (pre 3.7 Ga), suggesting a warmer, wetter context for their formation (Bultel et al., 2019). These detections are associated with surficial weathering profiles, i.e. environments in which nutrients may have been lost by leaching, thereby reducing habitability (Duddy, 1980; Nesbitt et al., 1980), although such environments on the modern, oxygenated and highly habitable Earth host a diverse array of life. At the ExoMars 2022 Oxia Planum landing site, there is a spectral suggestion of carbonate associated with phyllosilicates (Mandon et al., 2021) although the exact environment of deposition in this case has yet to be elucidated, whether weathering, lacustrine or marine (Quantin-Nataf et al., 2021). On the other hand, Jezero crater, the landing site of the Mars 2020 Perseverance rover, displays carbonates in association with a variety of interpreted paleoenvironments, including weathering profiles of volcanic ash, as well as marginal lakeshore and/or fluvial precipitations of carbonate that are compared to carbonate reefs, microbialites or tufa, having the potential to preserve the kinds of biosignatures described above (Horgan et al., 2020). Similarly, Franz et al. (2020) evoke carbonate in the endogenous carbon cycle of Gale crater (Fig. 5). The main problem with carbonates is their propensity to corrode under the acidic weathering conditions proposed for early Mars, which had a thicker CO₂ atmosphere than today (Viennet et al., 2019). Such conditions would lead to the destruction or significant alteration of potential biosignatures.

Other mineral associations with biosignatures: While not associated with the ALH84001 meteorite, other minerals that are present on Mars, including silica, phyllosilicates and evaporitic phases such as sulphates, could also preserve biosignatures. In addition, the preservation of MISS in clastic sediments should be noted, for example, in the Palaeo-Mid Archaean formations of the Barberton Greenstone Belt in South Africa and the Pilbara Greenstone Belt in Australia (Noffke, 2010; Noffke *et al.*, 2013).

Silica

(1) Siliceous hydrothermal environments, both subaqueous and subaerial, are prime sites for microbial development because of the significant supply of nutrients (Walter and Des Marais, 1993; Cady and Farmer, 1996). With their high concentrations of nutrients (either organic, such as small molecules, gases such as CH₄, H₂), essential transition elements (Fe, Ni, V, Co, As), and variable gradients in pH, Eh and temperature, they provide a habitat for a wide variety of microbial life (Baross, 1998). On present-day Earth, siliceous hydrothermal environments are essentially, but not only, associated with deep sea black smokers. Upon initial formation, these >300 °C environments are sterile but later become colonized by hyperthermophilic archaea transported by aqueous currents to the edifices (Wirth *et al.*, 2018).

We will here consider hydrothermal environments on the early Earth because of their particular relevance for early Mars, the environments of the two planets being similar (Westall et al., 2015b). Marine hydrothermal environments on the early Earth occurred at all water depths including in the shallow water basins on the submerged oceanic plateaux that characterized the early protocontinents. Indeed, they also occur in the littoral and adjacent subaerial environments (Westall et al., 2015a, 2015b; Djokic et al., 2017). Higher heat flow from the early Earth's mantle (Sleep, 2000; Zahnle et al., 2007; Kamber, 2015; Labrosse, 2015; Van Kranendonk et al., 2015) fuelled abundant volcanic and hydrothermal activity. Rare earth element (REE) signatures including Eu and Y anomalies, Y/Ho ratio, as well as the physical evidence of hydrothermal veins and pervasive silicification of all lithologies in contact with seawater, attest to the pervasive influence of hydrothermal fluids in the global ocean (e.g. Van Kranendonk, 2006; Hofmann and Bolhar, 2007; Hofmann and Harris, 2008; Westall et al., 2015b, 2018), as well as in restricted shallow basins on top of the oceanic plateaux where influences from non-marine waters have been additionally detected (Hickman-Lewis et al., 2020b).

Most physical evidence for hydrothermal activity on the early Earth points to subaqueous systems. Due to pervasive silicification (>96% SiO₂) of Palaeoarchaean sediments and the immediately underlying volcanics owing to oversaturation of silica in the early oceans, any physical expression of vents is difficult to detect, although some hydrothermal veins are readily identifiable by virtue of their cross-cutting nature (Hofmann and Bolhar, 2007; Westall et al., 2015a). Veins can also form bedding-parallel intrusions into partially lithified sediment (Westall et al., 2015a). The 3.33 Ga Josefsdal Chert from the Barberton Greenstone Belt in South Africa comprises both volcanic sands and chemically precipitated sedimentary silica gel, now chert. Adjacent to hydrothermal vents, the volcanic sands are heavily encrusted by carbonaceous matter (CM), while silica gel mixed with very fine volcanic dust is characterized by stellate clots of CM identified as probable chemotrophic biomass (Westall et al., 2015a, 2015b; Hickman-Lewis et al., 2020a). The 3D, irregular, stellate formations of CM around the volcanic clasts (up to 30 µm thickness) and in irregular clots within the silica gel (up to 500 µm diameter) are categorically incompatible with abiotic CM aggregation, deposition or diffusion. In addition, the CM has enhanced concentrations of transition metals (Fe, V, Ni, As and Co), and a δ^{13} C signature dominantly between -21.0 and -11.5‰ and as light as -41.3‰. These elemental and isotopic signatures, together with the morphological distribution of the CM and its petrographic context, suggest that this kerogen represents the remnants of lithotrophic or organotrophic consortia, possibly cycling methane or nitrogen (Hickman-Lewis et al., 2020a). When observed in field exposures, however, the heavy silicification makes direct detection of the clotted colonies in these sediments difficult, although sometimes a faint mottled texture reveals the presence of the carbonaceous clots.

It is rare for individual cells to be preserved morphologically in these ancient sediments, rapid silicification prior to heterotrophic degradation being a prerequisite (e.g. Fig. 3b). The rarely preserved examples show that individual cells are generally too small (<1 μ m) to be analysed individually by *in situ* methods on Mars, although, as we have seen above, it may be possible to identify the carbonaceous remains of colonies of several 100 s μ m size.

Not all Palaeoarchaean hydrothermal environments are subaqueous. Recently, subaerial sinter springs, similar to those on the modern Earth, have been described from the 3.48 Ga



Fig. 5. Possible carbon (and carbonate) cycle at Gale crater (Franz et al., 2020; Nature, permission).

Dresser Formation, Pilbara, Australia. Djokic *et al.* (2017, 2021) used textures and REE and yttrium geochemistry to demonstrate similarities between the Dresser geyserites and Phanerozoic hot spring sinters, noting the temporal association of a diverse suite of textural biosignatures that indicates a thriving microbial community. The biosignatures are primarily physical in nature, including stromatolites and microbial palisade fabric preserved in inferred mineralized exopolymeric substance. It is suggested that such geyserites and their textural features could be an explanation for the silica deposits identified around Home Plate in Gusev Crater (Ruff and Farmer, 2016; Djokic *et al.*, 2021).

Silica as a chemical sediment was ubiquitous on the early Earth both because of generally high silica concentrations in the sea water (Siever, 1992) due to high rates of weathering of exposed landmasses and lack of removal by organisms using silica as a framework for their cells (e.g. diatoms that evolved later), and because of elevated hydrothermal influence. Another significant silica contribution comes from the devitrification of subaqueous volcanic sediments. This high concentration of silica in the early oceans was the prime reason for the silicification of early sediments and volcanic rocks exposed to seawater. It is also the reason for the rapid preservation of microbial biosignatures, even in non-hydrothermal, oligotrophic environments, as described by Westall *et al.* (2011, 2015*a*).

Silica can also be concentrated in evaporitic settings, contributing to the entombment of microorganisms inhabiting the environment. Examples include the ~800 Ma Draken Formation in Svalbard where enhanced silica concentrations in an evaporitic environment resulted in the silicification of cyanobacterial mats (Foucher and Westall, 2013). While we do not expect to find cyanobacterial ecosystems on Mars, this example demonstrates the high fidelity of preservation by evaporitic silica and, in addition, documents a different type of biosignature, metastable minerals. In this example, metastable opal occurs within the organically preserved cell envelopes of the cyanobacteria while the enclosing silica cement is recrystallized to quartz, the organic phase enhancing the stability of the metastable mineral. It is further noteworthy that a beautifully preserved sequence of delicate microbial biofilms in the 3.33 Ga Josefsdal Chert formed in an evaporitic environment as attested to by the pseudomorph evaporite minerals embedded in their surfaces (Westall *et al.*, 2011). NanoSIMS scans across the thicknesses of the biofilms document that they had been thoroughly silicified.

Continuing with the theme of weathering profiles on Mars evoked above, Rutledge *et al.* (2018) documented high rates of glacial alteration of basaltic volcanic materials and the subsequent precipitation of amorphous (opaline) silica on rock surfaces in subglacial conditions on Iceland, analogous to those on early Mars; however, no biosignatures occur in these phases. Interestingly, a significant amount of poorly crystalline silica phases is mixed in with the detrital products of glacial weathering of the volcanic rock. X-ray amorphous silica is an important constituent of the sediments in Gale Crater where it can constitute up to 90% of the rock (Rapin *et al.*, 2018).

(2) *Silica on Mars*: Hydrated silica deposits have been detected from orbit in many locations at the surface of Mars. Sun and Milliken (2018) determined that the mineral phases detected fall into two categories: amorphous and/or dehydrated silica-bearing bedrock deposits, and more crystalline and/or hydrated silica-bearing aeolian deposits. The latter provide evidence of impact-related, hydrothermally produced opaline silica and the former altered volcanic phases or dehydrated silica (Pineau *et al.*, 2020).

The most significant deposit of silica on Mars found to date is that around Home Plate, where amorphous SiO₂.nH₂O occurs in what may be an ancient volcanic hydrothermal setting in Gusev crater (Fig. 6(a) and (b); Squyres *et al.*, 2008; Ruff *et al.*, 2011). This deposit has been attributed to either fumarole-related, acid-



Fig. 6. (a) High-resolution image of Home Plate from orbit (Tao *et al.*, 2021). (b) Comparison of the Home Plate silica deposits in Gusev Crater, Mars (left), with the El Tatio hot spring deposits in the Atacama (right); field views above, details of the silica spicules below (Ruff and Farmer, 2016).

sulphate leaching or precipitation from hot spring fluids with the latter being a privileged hypothesis based on analogue textural observations (Ruff and Farmer, 2016) from hot springs at El Tatio in Chile. These authors suggest that the nodular and digitate silica structures at El Tatio, produced by a combination of biotic and abiotic processes, are very similar to those observed at Home Plate. Djokic *et al.* (2021) also suggest similarities between textural biosignatures in Palaeoarchaean geyserites and those of Home Plate.

Hydrated (opaline) silica deposits detected in Jezero crater are of interest because they are a good mineral phase for biosignature preservation (Tarnas *et al.*, 2019). Associated with specific rock units, such as the opal-magnesite-smectite series detected in the olivine-rich unit and the opal-Al-phyllosilicate phase series co-located with the basement unit, the formation processes of the hydrated silica deposits are not yet understood. Possible candidates are: primary volcanism, diagenesis, authigenic formation in a lacustrine environment, detrital transport of material formed authigenically in the Jezero watershed, or transport to Jezero crater *via* aeolian processes. Hydrated silica has been detected from orbit at the edges of deltaic fan deposits formed in relationship with a standing body of water in Oxia Planum, the chosen landing site for the ExoMars 2022 mission (Quantin-Nataf *et al.*, 2021). Here, the hydrated silica deposits overlie clay-rich facies and indicate a changing hydrological regime possibly related to marine regression, or at least the drying of a water-filled basin, concurrent with the global hydrological trajectory of Mars through the Noachian and Hesperian periods.

Phyllosilicates

Direct morphological evidence of life will be difficult to detect on Mars because of (1) the very small size of the individual microorganisms, (2) the difficulty of distinguishing eventual microbial 'clotted' textures from abiotic ones, and (3) the possibility that distinctive macroscopic features, such as complex stromatolites, may not occur on Mars. Phyllosilicate minerals, however, have a propensity to accumulate and preserve organic matter, which chelates onto and in between the silicate sheets. As such, they are favoured mineral phases for the conservation of potential organic biosignatures (Bishop *et al.*, 2013; Vago *et al.*, 2017), although environmental parameters on Mars, such as UV radiation and acid leaching, can negatively influence the long-term preservation of organic molecules, even associated with phyllosilicates (Gil-Lozano *et al.*, 2020).

The hypothesis is that phyllosilicates on Mars may have chelated the organic remnants of microbial life. These minerals can also physically fossilize microorganisms and their EPS. Wacey *et al.* (2014) noted the role of authigenic clay precipitation in organism preservation based on experimental investigations and studies of phenomena in the natural environment, ranging from marine settings, lakes, rivers and hydrothermal systems. A variety of clay mineral types, including Fe-rich phyllosilicates (e.g. nontronite), as well as Al clays (kaolinite and halloysite), and Mg or K-rich clays, are implicated in this process (Fig. 3(d)). The composition of the authigenic minerals that precipitate under the influence of microbial mediation depends on factors, such as pH, Eh, sulphate concentration, ionic availability and the nature of the biological substrate, which will affect its affinity for specific ions.

Authigenic precipitation or chelation of phyllosilicates may occur on the surfaces of individual organisms in multispecies communities. Under specific physico-chemical conditions, this leads to the replacement of whole microbial mats by phyllosilicates. Phyllosilicates chelating to the surfaces of microorganisms form a 'halo' around them. Such phenomena have been observed in Oligocene (\sim 30 Ma) deep sea sediments (Monty *et al.*, 1991; Westall and Rincé, 1994), as well as in artificial fossilization of microbes in their deep sea sedimentary matrices (Fig. 3(c); Westall *et al.*, 1995). In all cases, there is an intimate intergrowth of phyllosilicates with microbial polymer (EPS) such that fibrillike strands of polymer stretch between packets of phyllosilicates (Westall *et al.*, 2015*b*).

The importance of phyllosilicates in the preservation of softbodied organisms was highlighted in ~500 My Burgess Shale-type deposits hosting soft-bodied fauna, where kaolinite played an important role in the preservation process (Gaines et al., 2008; LaFlamme et al., 2011). Experimental studies showed how this process worked. Working with cyanobacteria, well known for their formation of significant mats that covered the Ediacaran seafloors, thus entombing the soft-bodied macrobiota, Newman et al. (2016) exposed the microorganisms to elevated concentrations of clay minerals and silica, showing that clays coated the sheaths of the microorganisms through two mechanisms: trapping and indirect precipitation. Extending the importance of phyllosilicate precipitation to the cell walls of microorganisms, Wacey et al. (2014) studied 1 Ga lake sediments of the Torridon Group, Scotland, and described a microbiallymediated Fe-rich phyllosilicate phase that occurs in direct contact with cellular material; the Fe-rich phyllosilicates replace the most labile biological material, while a K-rich clay occurs within and exterior to cell envelopes, forming where the supply of Fe was exhausted (Fig. 3(d)).

Phyllosilicates on Mars: Orbital spectral measurements of the surface of Mars have identified large areas covered by diverse phyllosilicate phases. Their distribution and associations with other mineral phases suggest major compositional changes through time related to changing global environmental conditions on Mars (e.g. Poulet *et al.*, 2005; Bishop *et al.*, 2018). There is much debate as to the formation mechanisms and environmental constraints for the clays observed at Mars' surface. Their relative abundance in the Noachian suggests that, in its early history, Mars had a long-lived warmer and wetter climate, although this

scenario is at odds with climate models, which suggest that Noachian Mars was cold. Nevertheless, geomorphological features unambiguously indicate water activity at the surface of the planet. Noting that the time necessary for clay minerals to form in cold environments is extremely long, and taking into account climate models, geomorphological signatures and mineral phases, Bishop et al. (2018) hypothesized phyllosilicate formation during sporadic, short-term warm and wet periods on a generally cold early Mars. They concluded that associations of Mg-rich claybearing rocks with mixed Fe/Mg smectite, chlorite, talc, serpentine and zeolite likely formed in subsurface hydrothermal environments. On the other hand, other assemblages exhibiting vertical zonation of Al/Fe³⁺-rich and Fe/Mg smectites, as well as sulphates are interpreted to have formed in different physicochemical environments with different rock/water ratios on the surface of Mars. Another possibility is that the phyllosilicates could have formed preferentially in the water region in contact with a 'hot' ground - that is, heated from the interior, but not necessarily in the subsurface. In this scenario, most of Mars would have been cold (consistent with the then reduced insolation and with climate models), but with topical warm environments, where water flow and alteration could proceed mediated by internal heat. So, for example, some of the ancient hydrous features we observe today could have flowed under a top-frozen laver of water.

Clays at Gale crater: Gale crater is a ~150 km diameter impact crater of Mars located at the limit of the cratered highlands of the south and the low plains of the north. Its age is about 3.6-3.8 Ga (Noachian-Hesperian boundary) according to the crater counts (Thomson et al., 2011). Orbital observations by the visibleinfrared spectrometer, Compact Reconnaissance Imaging Spectrometer for Mars (CRISM) and the spectro-imager, Thermal Emission Imaging System (THEMIS) characterized the mineral phases in the crater, while the Mars Science Laboratory mission's rover Curiosity has been assessing past habitability in the region. Mount Sharp (formally named Aeolis Mons), the large central mound inside the crater, has relatively flat strata that may have preserved evidence of the evolution of the Mars environment (Grotzinger et al., 2012). The older, lower part of the mound contains clays minerals mixed with hydrated sulphates, indicating aqueous activity, while the younger layers higher on the mound are covered by a dust mantle (Milliken et al., 2010; Thomson et al., 2011; Wray, 2013).

Fluvial valley walls and crater rims and floors contain olivinebearing bedrock associated with Fe/Mg phyllosilicates, kaolin family minerals, hydrated silica and other hydrated/hydroxylated phases (Wray, 2013; Carter et al., 2015; Ehlmann and Buz, 2015). CRISM spectra of the walls and rims show absorption bands that can be associated to Fe/Mg phyllosilicates, such as nontronite and saponite, while the Mount Sharp spectra exhibit different spectral properties characteristic of Al phyllosilicates, including montmorillonite and kaolinite (Buz et al., 2017). The detection of different types of phyllosilicates thus suggests protoliths of a different nature or the formation of clays by fluids of different composition. There are several suggested scenarios to explain the origin of the clays on the crater floor: either the clays were transported to the crater floor from the already partially altered crater rim; or they were formed in situ by alteration of the mafic bedrock by water and subsequently altered as a result of burial diagenesis (Buz et al., 2017).

Clay minerals at Jezero Crater: The 3.7 Ga old Jezero Crater at the edge of Isidis Planitia hosts a well-formed delta and deposits

indicating the presence of lakes in the past. At the landing site of the Mars 2020 mission and the Perseverance rover, there is a diversity of phyllosilicate-bearing units that may have high biosignature preservation potential, including Al-clays and potential Fe-Mg-bearing smectites (Goudge et al., 2015; Brown et al., 2020; Horgan et al., 2020). These units are predominantly associated with the western delta, although zones of phyllosilicate-rich mineralogy have also been identified dispersed throughout the crater (Horgan et al., 2020), including the region close to the rover landing site. Additionally, localized zones of phyllosilicate richness have been noted within the marginal olivine carbonate unit (Goudge et al., 2015; Horgan et al., 2020). Systematic variations in CRISM spectral parameters have been used to infer variability in Fe/Mg-phyllosilicates and Mg-carbonates between the different (stratigraphic) units of Jezero crater; however, these interpretations await validation by surface observations and analyses. The regional-scale origin of phyllosilicate-rich phases and their relationships with the surrounding crater floor and margin units have been addressed in detail by Brown et al. (2020); these scenarios imply a complex geological history at Jezero crater with aqueous periods that may have been conducive to habitability throughout the Noachian and earliest Hesperian. Of particular interest for habitability and palaeoenvironment is the unresolved origin of Fe/Mg-phyllosilicates, which may have formed during near-surface weathering and diagenesis, through post-impact hydrothermal activity, or both (Ehlmann et al., 2021).

Clay minerals at Oxia Planum: Oxia Planum is a ~4 Ga depression located near the equator of the planet, on the eastern border of Chryse Planitia. It is the landing site chosen for the ESA (European Space Agency)-Roscosmos ExoMars 2022 mission. The Rosalind Franklin rover will touch down on 10 June 2023 to begin its search for traces of past and present life on Mars and to study the aqueous environment (Vago et al., 2017). Oxia Planum is particularly flat and contains clay-rich lithofacies that suggest past water activity in the region. The clay deposits show absorption bands centred at 1.9 and 2.3 µm that are characteristic of Fe/Mg clays, probably a vermiculite-smectite type of clay (Carter et al., 2016). In some places, an additional spectral absorption at 2.5 µm likely suggests the presence of a carbonate or other type of clay mineral. If carbonate phases are present, the site may have been preserved from any acidic environment, which enhances its potential for preserving traces of life (Mandon et al., 2021).

The layered clay-bearing unit has a total thickness of 100 m, with each layer varying from 0.7 to 3 m in thickness. According to HiRISE images, the clay-rich facies corresponds to a fractured material of different tonalities that is widely exposed and can be divided into two sub-units: the first one has a reddish tone with smaller fractures and the second one overlaps the first one and has a bluish tone with wider fractures (Quantin-Nataf et al., 2021). The colour difference is explained by mineralogical composition, rather than variable surficial dust or sand cover (Mandon et al., 2021). The clay-rich deposits cover terrains of mid- and late-Noachian age, draping the paleotopography, therefore, the unit is mid-Noachian or younger. These strata are systematically overlain by a thin, mid-toned, mantling unit that appears to be indurated and layered with a thickness of $\sim 5 \text{ m}$ (Quantin-Nataf et al., 2021). The clays may have a marine or lacustrine sedimentary origin. Other units present in the region include deltaic and fluvial deposits, remnant rounded buttes and a dark resistant unit. The latter is recognizable by its rough, dark and massive appearance and it notably overlies the

other units. Its thickness is ~ 20 m, it appears to be composed of mafic minerals, and is probably of volcanic origin, possibly Amazonian in age (Carter *et al.*, 2016; Quantin-Nataf *et al.*, 2021). Thus, volcanic rocks covered the clays and other aqueous deposits, preserving potential biosignatures from the severe radiation and oxidation of the surface of the planet. The clay-bearing facies have been exposed by erosion relatively recently.

Evaporite minerals

Inorganic precipitation from supersaturated solutions forms evaporitic deposits in a variety of marine and terrestrial environments. Evaporite deposits - soluble salts precipitating from brines (Warren, 2016) - and minerals have good biosignature preservation potential because they can embed and preserve sediments containing organic matter, as well as trapping microorganisms including endoliths and fluid-gas-solid inclusions. Such deposits can be preserved over long periods of geological time (e.g. Shkolyar and Farmer, 2018; Benison, 2019). Rapid growth of halite and gypsum from brines may easily entrap primary fluid inclusions that could serve as proxies for environmental conditions, including life. A number of taphonomic processes in evaporites and postdepositional changes, such as sediment phase solubility, porosity, permeability and postdepositional contamination, which can negatively affect the preservation of biosignatures (e.g. Barbieri, 2013). Nevertheless, there are several factors that can enhance biosignature preservation even though the window for preservation occurs under a narrow range of conditions that exist immediately after deposition (e.g. Shkolvar and Farmer, 2018; Benison, 2019). For instance, in hypersaline sabka environments, the dense-stratified brines can increase bottom-water anoxia and promote organic preservation (e.g. Westall and Cavalazzi, 2011). On Earth, many kinds of halophiles have been isolated from a wide range of evaporitic environments, including sabka, salt marshes, undersea salt domes and subterranean halite deposits from evaporated ancient seas (e.g. Warren, 2016; Benison, 2019). Microorganisms have been isolated from salt crystals, evaporite deposits and associated fluid inclusions, in the rock record (Stan-Lotter et al., 2002; Vreeland et al., 2007; Schubert et al., 2010; Fendrihan et al., 2012; Carnevale et al., 2019). For example, great palaeobiodiversity is described for the Miocenic Messinian evaporitic deposits, Italy (Carnevale et al., 2019). Recently, the unique extreme environments at the Dallol geothermal area, Ethiopia (Cavalazzi et al., 2019; Cavalazzi and Filippidou, 2021) have provided the opportunity to study life at its biophysical limits in a hypersaline poly-extreme environment (Belilla et al., 2019; Gómez et al., 2019). Importantly, MISS are common in such environments where evaporate minerals may form on and within the microbial mats (Noffke, 2021).

Evaporites on Mars: Evaporite minerals and deposits, such as hydrated Mg or Fe sulphates, have been observed on Mars both from orbit and *in situ* (Squyres *et al.*, 2006; Rapin *et al.*, 2016), suggesting formation under evaporitic conditions within the context of past shallow lake, mudflat/sandflat and eolian environments, as well as past shallow acid saline groundwaters (Grotzinger *et al.*, 2005; Hynek *et al.*, 2015; Lynch *et al.*, 2015; Pondrelli *et al.*, 2017; Benison, 2019). Sulphate- and other saltbearing deposits occur in several places on Mars, including Meridiani Planum, Gusev Crater and Gale Crater (Grotzinger *et al.*, 2005; Ehlmann *et al.*, 2016; Pondrelli *et al.*, 2017). The thermochemical stability of such deposits under Martian conditions would enhance their potential for preserving possible halophilic microorganisms.

At Gale crater, for example, muddy to sandy sediments deposited in a lacustrine environment are characterized by a network of fractures filled in with light-toned deposits of sulphate evaporites, as well as nodules and raised ridges of similar materials. The fractures formed in the consolidated sediments owing to subsequent compression and extension related to fluid overpressure and cracking (De Toffoli *et al.*, 2020). They were then infilled by briny solutions that deposited the evaporite mineral suites. Rapin *et al.* (2019) documented bulk enrichments within the bedrock of significant amounts of calcium sulphate intercalated with a stratum enriched in hydrated magnesium sulphate resulting from early diagenetic, pre-compaction salt precipitation from brines concentrated by evaporation. Such environments would be habitable for halophilic-type microorganisms.

Experimental studies document the ability of halophiles to survive when exposed to surface Mars conditions (e.g. UV radiation, cold temperatures and desiccation; Oren, 2014). For all these reasons, evaporite deposits are a key exploration target for the ongoing astrobiological exploration of Mars and for future Mars Sample Return programs (Hays *et al.*, 2017).

Concluding remarks and significance for Mars missions

The McKay *et al.* (1996) paper was a seminal benchmark that stimulated the opening of the field of astrobiology and the search for life on Mars. Despite the fact that the weight of subsequent experiments and understanding of the microbial world and its preservation in the rock record shows that the original conclusions by McKay *et al.* were incorrect, their multiscale and multi-technique approach to documenting a variety of types of complementary, potential biosignatures paved the way for future work. A key parameter that was missing was the *in situ* geological context (which could, however be surmised).

Most important over the last quarter century is the crossdisciplinary understanding between experts in specific domains that is the only path to providing a holistic understanding of traces of past microbial life, both on Earth and elsewhere in the Solar System. It is the complementary expertise that will enable deciphering the rock record of Mars with the missions Mars Science Laboratory, Mars 2020 and ExoMars 2022 to determine habitability and to identify any potential evidence of past life (without excluding the minute possibility of extant life).

In this mini-review, we have discussed the nature of biosignatures and their formation and preservation through the lens of the ALH84001 study. We have documented a non-exhaustive but pertinent variety of mineral phases that can preserve biosignatures, including carbonates, silica, phyllosilicates and evaporite phases, both in surface environments and in the subsurface. This latter point is of particular interest for the ExoMars 2022 mission because of the ability of the rover to drill to depths of up to 2 m, a first in the history of Mars exploration. The importance of this feat is that organic biosignatures (or even abiotic/prebiotic organic signatures) will be better protected from the deleterious effects of surface oxidation and UV radiation at such depths. Secondly, the targeted site, Oxia Planum, comprises clay-enriched lithologies. As we have seen above, phyllosilicates have high propensity for trapping, chelating and protecting organic molecules. Thirdly, Oxia Planum at ~4-3.9 Ga is the oldest location yet to be investigated, having formed during a period when Mars had wetter and more habitable conditions. Thus, if there are preserved traces of Martian life at the landing site, the ExoMars 2022 rover, Rosalind Franklin, should be able to uncover them.

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/S1473550421000264.

Conflict of interest. None.

References

- Allwood AC, Walter MR, Kamber BS, Marshall CP and Burch IW (2006) Stromatolite reef from the Early Archaean era of Australia. *Nature* **441**, 714–718.
- Banfield JF, Moreau JW, Chan CS, Welch SA and Little B (2001) Mineralogical biosignatures and the search for life on Mars. *Astrobiology* **1**, 447–465.
- Barbieri R (2013) The role of terrestrial analogs in the exploration of the habitability of Martian evaporitic environments. In de Vera J-P and Seckbach J (eds), *Habitability of Other Planets and Satellites*. Dordrecht: Springer, pp. 163–180.
- Barbieri R and Cavalazzi B (2008) Fossil microorganisms at methane seeps: an astrobiological perspective. In Seckbach J and Walsh M (eds), From Fossils to Astrobiology: Records of Life on Earth and Search for Extraterrestrial Biosignatures. Dordrecht: Springer, pp. 297–317.

Baross JA (1998) Living on the edge. Nature 395, 136.

- Belilla J, Moreira D, Jardillier L, Reboul G, Benzerara K, López-García JM, Bertolino P, López-Archilla AI and López-García P (2019) Hyperdiverse archaea near life limits at the polyextreme geothermal Dallol area. *Nature Ecology & Evolution* 3, 1552–1561.
- **Benison KC** (2019) How to search for life in martian chemical sediments and their fluid and solid inclusions using petrographic and spectroscopic methods. *Frontiers in Environmental Science* **7**, 108.
- Benzerara K, Menguy N, Guyot F, Dominici C and Gillet P (2003) Nanobacteria-like calcite single crystals at the surface of the Tataouine meteorite. *Proceedings of the National Academy of Sciences* 100, 7438.
- Bishop JL, Loizeau D, McKeown NK, Saper L, Dyar MD, Des Marais DJ, Parente M and Murchie SL (2013) What the ancient phyllosilicates at Mawrth Vallis can tell us about possible habitability on early Mars. *Planetary and Space Science* 86, 130–149.
- Bishop JL, Fairén AG, Michalski JR, Gago-Duport L, Baker LL, Velbel MA, Gross C and Rampe EB (2018) Surface clay formation during short-term warmer and wetter conditions on a largely cold ancient Mars. *Nature Astronomy* 2, 206–213.
- Borg LE, Connelly JN, Nyquist LE, Shih C-Y, Wiesmann H and Reese Y (1999) The age of the carbonates in martian meteorite ALH84001. *Science (New York, N.Y.)* 286, 90.
- **Bosak T and Newman DK** (2003) Microbial nucleation of calcium carbonate in the Precambrian. *Geology* **31**, 577–580.
- Bosak T, Knoll AH and Petroff AP (2013) The meaning of Stromatolites. Annual Review of Earth and Planetary Sciences 41, 21–44.
- Brocks JJ and Summons RE (2003) Sedimentary hydrocarbons, biomarkers for early life. *Treatise on Geochemistry* **8**, 63–115.
- Brown AJ, Viviano CE and Goudge TA (2020) Olivine-carbonate mineralogy of Jezero Crater. Journal of Geophysical Research Planets 125, e2019JE006011.
- Brunelli G, Carinci F, Zollino I, Candotto V, Scarano A and Lauritano D (2012) SEM evaluation of 10 infected implants retrieved from man. *European Journal of Inflammation* **10**, 7–12.
- Bultel B, Viennet J-C, Poulet F, Carter J and Werner SC (2019) Detection of carbonates in Martian weathering profiles. *Journal of Geophysical Research: Planets* 124, 989–1007.
- Buz J, Ehlmann BL, Pan L and Grotzinger JP (2017) Mineralogy and stratigraphy of the Gale crater rim, wall, and floor units. *Journal of Geophysical Research: Planets* **122**, 1090–1118.
- Byerly GR, Lowe DR and Walsh MM (1986) Stromatolites from the 3,300– 3,500-Myr Swaziland supergroup, Barberton Mountain Land, South Africa. *Nature* **319**, 489–491.
- Cacchio P, Ercole C, Cappuccio G and Lepidi A (2003) Calcium carbonate precipitation by bacterial strains isolated from a limestone cave and from a loamy soil. *Geomicrobiology Journal* 20, 85–98.
- Cady SL and Farmer JD (1996) Fossilization processes in siliceous thermal springs: trends in preservation along thermal gradients. *Ciba Foundation Symposium*. Chichester: Wiley, pp. 150–173.

- Carnevale G, Gennari R, Lozar F, Natalicchio M, Pellegrino L and Dela Pierre F (2019) Living in a deep desiccated Mediterranean Sea: an overview of the Italian fossil record of the Messinian salinity crisis. *Bollettino della Società Paleontologica Italiana* 58, 109–140.
- Carter J, Viviano-Beck C, Loizeau D, Bishop J and Le Deit L (2015) Orbital detection and implications of akaganéite on Mars. *Icarus* 253, 296–310.
- **Carter J, Quantin C, Thollot P, Loizeau D, Ody A and Lozach L** (2016) Oxia planum: a clay-laden landing site proposed for the ExoMars rover mission: aqueous mineralogy and alteration scenarios. Paper presented at *47th Lunar and Planetary Science Conference*, The Woodlands, Texas: LPI Contributions, 2064.
- Casar CP, Kruger BR, Flynn TM, Masterson AL, Momper LM and Osburn MR (2020) Mineral-hosted biofilm communities in the continental deep subsurface, Deep Mine Microbial Observatory, SD, USA. *Geobiology* 18, 508–522.
- Casar CP, Kruger BR and Osburn MR (2021) Rock-hosted subsurface biofilms: mineral selectivity drives hotspots for intraterrestrial life. *Frontiers in Microbiology* **12**, 702.
- Cavalazzi B (2007) Chemotrophic filamentous microfossils from the Hollard Mound (Devonian, Morocco) as investigated by focused ion beam. *Astrobiology* 7, 402–415.
- **Cavalazzi B and Filippidou S** (2021) Microbial survival and adaptation in extreme terrestrial environments the case of the Dallol Geothermal area in Ethiopia. In Branislav V, Joseph S and Roger G (eds), *Planet Formation and Panspermia: New Prospects for the Movement of Life through Space.* Beverly, MA: Scrivener Publishing LLC, pp. 93–118.
- Cavalazzi B, Barbieri R and Ori GG (2007) Chemosynthetic microbialites in the Devonian carbonate mounds of Hamar Laghdad (Anti-Atlas, Morocco). *Sedimentary Geology* 200, 73–88.
- **Cavalazzi B, Westall F, Cady SL, Barbieri R and Foucher F** (2011) Potential fossil endoliths in vesicular pillow basalt, coral patch seamount, Eastern North Atlantic Ocean. *Astrobiology* **11**, 619–632.
- Cavalazzi B, Westall F and Cady SL (2012) Vesicular basalts as a niche for microbial life. In Hanslmeier A, Kempe S and Seckbach J (eds), *Life on Earth and other Planetary Bodies. Cellular Origin, Life in Extreme Habitats and Astrobiology.* Dordrecht: Springer, pp. 27–43.
- Cavalazzi B, Barbieri R, Gómez F, Capaccioni B, Olsson-Francis K, Pondrelli M, Rossi AP, Hickman-Lewis K, Agangi A, Gasparotto G, Glamoclija M, Ori GG, Rodriguez N and Hagos M (2019) The Dallol geothermal area, Northern Afar (Ethiopia) – an exceptional planetary field analog on Earth. Astrobiology 19, 553–578.
- Delarue F, Robert F, Derenne S, Tartèse R, Jauvion C, Bernard S, Pont S, Gonzalez-Cano A, Duhamel R and Sugitani K (2020) Out of rock: a new look at the morphological and geochemical preservation of microfossils from the 3.46 Gyr-old Strelley Pool Formation. *Precambrian Research* **336**, 105472.
- De Toffoli B, Mangold N, Massironi M, Zanella A, Pozzobon R, Le Mouélic S, L'haridon J and Cremonese G (2020) Structural analysis of sulfate vein networks in Gale crater (Mars). *Journal of Structural Geology* 137, 104083.
- Djokic T, Van Kranendonk MJ, Campbell KA, Walter MR and Ward CR (2017) Earliest signs of life on land preserved in ca. 3.5 Ga hot spring deposits. *Nature Communications* **8**, 15263.
- Djokic T, Van Kranendonk MJ, Campbell KA, Havig JR, Walter MR and Guido DM (2021) A reconstructed subaerial hot spring field in the ~3.5 billion-year-old dresser formation, North Pole Dome, Pilbara Craton, Western Australia. *Astrobiology* **21**, 1–38.
- Drake H, Åström ME, Heim C, Broman C, Åström J, Whitehouse M, Ivarsson M, Siljeström S and Sjövall P (2015) Extreme 13 C depletion of carbonates formed during oxidation of biogenic methane in fractured granite. *Nature Communications* 6, 1–9.
- **Duddy LR** (1980) Redistribution and fractionation of rare-earth and other elements in a weathering profile. *Chemical Geology* **30**, 363–381.
- Dupraz C, Reid RP, Braissant O, Decho AW, Norman RS and Visscher PT (2009) Processes of carbonate precipitation in modern microbial mats. *Microbial Mats in Earth's Fossil Record of Life: Geobiology* **96**, 141–162.
- Ehlmann BL and Buz J (2015) Mineralogy and fluvial history of the watersheds of Gale, Knobel, and Sharp craters: a regional context for the Mars Science Laboratory Curiosity's exploration. *Geophysical Research Letters* 42, 264–273.

- Ehlmann BL, Anderson FS, Andrews-Hanna J, Catling DC, Christensen PR, Cohen BA, Dressing CD, Edwards CS, Elkins-Tanton LT, Farley KA, Fassett CI, Fischer WW, Fraeman AA, Golombek MP, Hamilton VE, Hayes AG, Herd CDK, Horgan B, Hu R, Jakosky BM, Johnson JR, Kasting JF, Kerber L, Kinch KM, Kite ES, Knutson HA, Lunine JI, Mahaffy PR, Mangold N, McCubbin FM, Mustard JF, Niles PB, Quantin-Nataf C, Rice MS, Stack KM, Stevenson DJ, Stewart ST, Toplis MJ, Usui T, Weiss BP, Werner SC, Wordsworth RD, Wray JJ, Yingst RA, Yung YL and Zahnle KJ (2016) The sustainability of habitability on terrestrial planets: insights, questions, and needed measurements from Mars for understanding the evolution of Earth-like worlds. *Journal of Geophysical Research: Planets* 121, 1927–1961.
- Ehlmann BL, Bell J, Brown A, Horgan B, Hurowitz JA, Kelemen P, Mangold N, Mayhew L, Quantin C, Rapin W, Razzell Hollis J, Scheller E, Shuster D, Stack K, Sun V, Tarnas J and Treiman A (2021) Mineralogy from Mars-2020: updates to the regional geologic history of Jezero Crater, its watershed, and a framework for perseverance exploration. Paper presented at *Lunar and Planetary Science Conference*, 1721.
- Eigenbrode JL, Summons RE, Steele A, Freissinet C, Millan M, Navarro-González R, Sutter B, McAdam AC, Franz HB, Glavin DP, Archer PD, Mahaffy PR, Conrad PG, Hurowitz JA, Grotzinger JP, Gupta S, Ming DW, Sumner DY, Szopa C, Malespin C, Buch A and Coll P (2018) Organic matter preserved in 3-billion-year-old mudstones at Gale crater, Mars. Science (New York, N.Y.) 360, 1096–1101.
- Einen J, Kruber C, Øvreås L, Thorseth IH and Torsvik T (2006) Microbial colonization and alteration of basaltic glass. *Biogeosciences Discussions* 3, 273–307.
- Fendrihan S, Dornmayr-Pfaffenhuemer M, Gerbl FW, Holzinger A, Grösbacher M, Briza P, Erler A, Gruber C, Plätzer K and Stan-Lotter H (2012) Spherical particles of halophilic archaea correlate with exposure to low water activity – implications for microbial survival in fluid inclusions of ancient halite. *Geobiology* 10, 424–433.
- Folk RL and Lynch FL (1997) The possible role of nannobacteria (dwarf bacteria) in clay-mineral diagenesis and the importance of careful sample preparation in high-magnification SEM study. *Journal of Sedimentary Research* 67, 583–589.
- Folk RL and Lynch FL (2001) Organic matter, putative nannobacteria and the formation of ooids and hardgrounds. *Sedimentology* **48**, 215–229.
- Foucher F and Westall F (2013) Raman imaging of metastable opal in carbonaceous microfossils of the 700–800 Ma old Draken formation. *Astrobiology* 13, 57–67.
- Foucher F, Westall F, Brandstätter F, Demets R, Parnell J, Cockell CS, Edwards HG, Bény J-M and Brack A (2010) Testing the survival of microfossils in artificial Martian sedimentary meteorites during entry into Earth's atmosphere: the STONE 6 experiment. *Icarus* 207, 616–630.
- Franz HB, Mahaffy PR, Webster CR, Flesch GJ, Raaen E, Freissinet C, Atreya SK, House CH, McAdam AC, Knudson CA, Archer PD, Stern JC, Steele A, Sutter B, Eigenbrode JL, Glavin DP, Lewis JMT, Malespin CA, Millan M, Ming DW, Navarro-González R and Summons RE (2020) Indigenous and exogenous organics and surface–atmosphere cycling inferred from carbon and oxygen isotopes at Gale crater. *Nature Astronomy* 4, 526–532.
- Friedmann EI and Koriem AM (1989) Life on Mars: how it disappeared (if it was ever there). Advances in Space Research 9, 167–172.
- Furnes H, Banerjee NR, Muehlenbachs K, Staudigel H and de Wit M (2004) Early life recorded in Archean pillow lavas. *Science (New York, N.Y.)* **304**, 578.
- Furnes H, Banerjee NR, Staudigel H, Muehlenbachs K, McLoughlin N, de Wit M and Van Kranendonk M (2007) Comparing petrographic signatures of bioalteration in recent to Mesoarchean pillow lavas: tracing subsurface life in oceanic igneous rocks. *Earliest Evidence of Life on Earth* **158**, 156–176.
- Gaboyer F, Le Milbeau C, Bohmeier M, Schwendner P, Vannier P, Beblo-Vranesevic K, Rabbow E, Foucher F, Gautret P and Guégan R (2017) Mineralization and preservation of an extremotolerant bacterium isolated from an early Mars analog environment. *Scientific Reports* 7, 1–14.
- Gaines RR, Briggs DEG and Yuanlong Z (2008) Cambrian Burgess Shale-type deposits share a common mode of fossilization. *Geology* 36, 755–758.
- García GM, Márquez GMA and Moreno HCX (2016) Characterization of bacterial diversity associated with calcareous deposits and drip-waters,

and isolation of calcifying bacteria from two Colombian mines. *Microbiological Research* **182**, 21–30.

- Gibson EK, McKay DS, Thomas-Keprta KL, Wentworth SJ, Westall F, Steele A, Romanek CS, Bell MS and Toporski J (2001) Life on Mars: evaluation of the evidence within Martian meteorites ALH84001, Nakhla, and Shergotty. *Precambrian Research* **106**, 15–34.
- Gil-Lozano C, Fairén AG, Muñoz-Iglesias V, Fernández-Sampedro M, Prieto-Ballesteros O, Gago-Duport L, Losa-Adams E, Carrizo D, Bishop JL, Fornaro T and Mateo-Martí E (2020) Constraining the preservation of organic compounds in Mars analog nontronites after exposure to acid and alkaline fluids. *Scientific Reports* 10, 15097.
- Golden DC, Ming DW, Morris RV, Brearley AJ, Lauer Jr HV, Treiman AH, Zolensky ME, Schwandt CS, Lofgren GE and McKay GA (2004) Evidence for exclusively inorganic formation of magnetite in Martian meteorite ALH84001. American Mineralogist 89, 681–695.
- Gómez F, Cavalazzi B, Rodríguez N, Amils R, Ori GG, Olsson-Francis K, Escudero C, Martínez JM and Miruts H (2019) Ultra-small microorganisms in the polyextreme conditions of the Dallol volcano, Northern Afar, Ethiopia. Scientific Reports 9, 1–9.
- Goudge TA, Mustard JF, Head JW, Fassett CI and Wiseman SM (2015) Assessing the mineralogy of the watershed and fan deposits of the Jezero crater paleolake system, Mars. *Journal of Geophysical Research: Planets* 120, 775–808.
- Grotzinger JP, Arvidson RE, Bell Iii JF, Calvin W, Clark BC, Fike DA, Golombek M, Greeley R, Haldemann A and Herkenhoff KE (2005) Stratigraphy and sedimentology of a dry to wet eolian depositional system, Burns formation, Meridiani Planum, Mars. Earth and Planetary Science Letters 240, 11–72.
- Grotzinger JP, Crisp J, Vasavada AR, Anderson RC, Baker CJ, Barry R, Blake DF, Conrad P, Edgett KS, Ferdowski B, Gellert R, Gilbert JB, Golombek M, Gómez-Elvira J, Hassler DM, Jandura L, Litvak M, Mahaffy P, Maki J, Meyer M, Malin MC, Mitrofanov I, Simmonds JJ, Vaniman D, Welch RV and Wiens RC (2012) Mars science laboratory mission and science investigation. Space Science Reviews 170, 5–56.
- Halevy I, Fischer WW and Eiler JM (2011) Carbonates in the Martian meteorite Allan Hills 84001 formed at 18 ± 4 °C in a near-surface aqueous environment. *Proceedings of the National Academy of Sciences* 108, 16895.
- Harvey RP and McSween Jr HY (1994) Ancestor's bones and palimpsests: olivine in ALH 84001 and orthopyroxene in Chassigny. *Meteoritics* 29, 472.
- Hassenkam T, Andersson MP, Dalby KN, Mackenzie DMA and Rosing MT (2017) Elements of Eoarchean life trapped in mineral inclusions. *Nature* **548**, 78–81.
- Hays LE, Graham HV, Des Marais DJ, Hausrath EM, Horgan B, McCollom TM, Parenteau MN, Potter-McIntyre SL, Williams AJ and Lynch KL (2017) Biosignature preservation and detection in Mars analog environments. Astrobiology 17, 363–400.
- Heubeck C (2009) An early ecosystem of Archean tidal microbial mats (Moodies Group, South Africa, ca. 3.2 Ga). Geology 37, 931–934.
- Hickman-Lewis K, Garwood RJ, Brasier MD, Goral T, Jiang H, McLoughlin N and Wacey D (2016) Carbonaceous microstructures from sedimentary laminated chert within the 3.46 Ga Apex Basalt, Chinaman Creek locality, Pilbara, Western Australia. *Precambrian Research* 278, 161–178.
- Hickman-Lewis K, Cavalazzi B, Sorieul S, Gautret P, Foucher F, Whitehouse MJ, Jeon H, Georgelin T, Cockell CS and Westall F (2020*a*) Metallomics in deep time and the influence of ocean chemistry on the metabolic landscapes of Earth's earliest ecosystems. *Scientific Reports* **10**, 1–16.
- Hickman-Lewis K, Gourcerol B, Westall F, Manzini D and Cavalazzi B (2020b) Reconstructing Palaeoarchaean microbial biomes flourishing in the presence of emergent landmasses using trace and rare earth element systematics. *Precambrian Research* **342**, 105689.
- Hickman-Lewis K, Westall F and Cavalazzi B (2020c) Diverse communities of Bacteria and Archaea flourished in Palaeoarchaean (3.5–3.3 Ga) microbial mats. *Palaeontology* **63**, 1007–1033.
- Hoehler TM and Jørgensen BB (2013) Microbial life under extreme energy limitation. Nature Reviews Microbiology 11, 83–94.
- Hofmann A and Bolhar R (2007) Carbonaceous cherts in the Barberton greenstone belt and their significance for the study of early life in the Archean record. *Astrobiology* 7, 355–388.

- Hofmann HJ and Davidson A (1998) Paleoproterozoic stromatolites, Hurwitz Group, Quartzite Lake area, Northwest Territories, Canada. Canadian Journal of Earth Sciences 35, 280–289.
- Hofmann A and Harris C (2008) Silica alteration zones in the Barberton greenstone belt: a window into subseafloor processes 3.5–3.3 Ga ago. *Chemical Geology* 257, 221–239.
- Hofmann HJ, Grey K, Hickman AH and Thorpe RI (1999) Origin of 3.45 Ga coniform stromatolites in Warrawoona group, Western Australia. *Geological Society of America Bulletin* 111, 1256–1262.
- Homann M, Heubeck C, Airo A and Tice MM (2015) Morphological adaptations of 3.22 Ga-old tufted microbial mats to Archean coastal habitats (Moodies Group, Barberton Greenstone Belt, South Africa). *Precambrian Research* 266, 47–64.
- Horgan BHN, Anderson RB, Dromart G, Amador ES and Rice MS (2020) The mineral diversity of Jezero crater: evidence for possible lacustrine carbonates on Mars. *Icarus* 339, 113526.
- Hynek BM, Osterloo MK and Kierein-Young KS (2015) Late-stage formation of Martian chloride salts through ponding and evaporation. *Geology* 43, 787–790.
- Johnson SS, Anslyn EV, Graham HV, Mahaffy PR and Ellington AD (2018a) Fingerprinting non-terran biosignatures. Astrobiology 18, 915–922.
- Johnson SS, Zaikova E, Millan M, Williams AJ, Craft K, Wagner NY, Bevilacqua JG, Bai A, Fuqua S and Kobs-Nawotniak SE (2018b) Searching for life in the Martian dark: an investigation of biosignatures in analog lava tubes. Paper presented at *AGU Fall Meeting Abstracts*, P24D-08.
- Jull AJT, Courtney C, Jeffrey DA and Beck JW (1998) Isotopic evidence for a terrestrial source of organic compounds found in Martian meteorites Allan Hills 84001 and Elephant Moraine 79001. Science (New York, N.Y.) 279, 366–369.
- Kamber BS (2015) The evolving nature of terrestrial crust from the Hadean, through the Archaean, into the Proterozoic. *Precambrian Research* 258, 48–82.
- Labrosse S (2015) Thermal evolution of the core with a high thermal conductivity. *Transport Properties of the Earth's Core* 247, 36–55.
- Laflamme M, Schiffbauer JD, Narbonne GM and Briggs DEG (2011) Microbial biofilms and the preservation of the Ediacara biota. *Lethaia* 44, 203–213.
- Lang SQ, Früh-Green GL, Bernasconi SM, Lilley MD, Proskurowski G, Méhay S and Butterfield DA (2012) Microbial utilization of abiogenic carbon and hydrogen in a serpentinite-hosted system. *Geochimica et Cosmochimica Acta* 92, 82–99.
- Lapen TJ, Righter M, Brandon AD, Debaille V, Beard BL, Shafer JT and Peslier AH (2010) A younger age for ALH84001 and its geochemical link to shergottite sources in Mars. Science (New York, N.Y.) 328, 347–351.
- Lepot K, Benzerara K and Philippot P (2011) Biogenic versus metamorphic origins of diverse microtubes in 2.7 Gyr old volcanic ashes: multi-scale investigations. *Earth and Planetary Science Letters* **312**, 37–47.
- Liebig K, Westall F and Schmitz M (1996) A study of fossil microstructures from the Eocene Messel Formation using transmission electron microscopy. *Neues Jahrbuch für Geologie und Paläontologie-Monatshefte* 4, 218–231.
- Lynch KL, Horgan BH, Munakata-Marr J, Hanley J, Schneider RJ, Rey KA, Spear JR, Jackson WA and Ritter SM (2015) Near-infrared spectroscopy of lacustrine sediments in the Great Salt Lake Desert: an analog study for Martian paleolake basins. *Journal of Geophysical Research: Planets* **120**, 599–623.
- Mandon L, Parkes Bowen A, Quantin-Nataf C, Bridges JC, Carter J, Pan L, Beck P, Dehouck E, Volat M, Thomas N, Cremonese G, Tornabene LL and Thollot P (2021) Morphological and spectral diversity of the claybearing unit at the ExoMars landing site Oxia Planum. Astrobiology 21, 464–480.
- McKay DS, Gibson EK, Thomas-Keprta KL, Vali H, Romanek CS, Clemett SJ, Chillier XD, Maechling CR and Zare RN (1996) Search for past life on Mars: possible relic biogenic activity in Martian meteorite ALH84001. *Science (New York, N.Y.)* 273, 924–930.
- McLoughlin N, Staudigel H, Furnes H, Eickmann B and Ivarsson M (2010) Mechanisms of microtunneling in rock substrates: distinguishing endolithic biosignatures from abiotic microtunnels. *Geobiology* **8**, 245–255.

- McLoughlin N, Grosch EG, Kilburn MR and Wacey D (2012) Sulfur isotope evidence for a Paleoarchean subseafloor biosphere, Barberton, South Africa. *Geology* 40, 1031–1034.
- McMahon S and Parnell J (2014) Weighing the deep continental biosphere. *FEMS Microbiology Ecology* 87, 113–120.
- McMahon S, Bosak T, Grotzinger JP, Milliken RE, Summons RE, Daye M, Newman SA, Fraeman A, Williford KH and Briggs DEG (2018) A field guide to finding fossils on Mars. *Journal of Geophysical Research: Planets* 123, 1012–1040.
- Milliken RE, Grotzinger JP and Thomson BJ (2010) Paleoclimate of Mars as captured by the stratigraphic record in Gale Crater. *Geophysical Research Letters* 37, L04201.
- Mittlefehldt DW (1994) ALH84001, a cumulate orthopyroxenite member of the Martian meteorite clan. *Meteoritics* 29, 214–221.
- Monty CLV, Westall F and van der Gaast S (1991) 37. Diagenesis of siliceous particles in subantartic sediments, Hole 699A: possible microbial mediation. *Proceedings of the Ocean Drilling Program, Scientific Results* **114**, 685–710.
- Mustard JF, Adler M, Allwood A, Bass DS, Beaty DW, Bell JF, Brinckerhoff W, Carr M, Des Marais DJ and Brake B (2013) Report of the Mars 2020 science definition team. Mars Exploration Program Analysis Group (MEPAG) 150, 155–205.
- Nesbitt HW, Markovics G and Price RC (1980) Chemical processes affecting alkalis and alkaline earths during continental weathering. *Geochimica et Cosmochimica Acta* 44, 1659–1666.
- Newman SA, Mariotti G, Pruss S and Bosak T (2016) Insights into cyanobacterial fossilization in Ediacaran siliciclastic environments. *Geology* 44, 579–582.
- Noffke N (2009) The criteria for the biogenicity of microbially induced sedimentary structures (MISS) in Archean and younger, sandy deposits. *Microbial Mats in Earth's Fossil Record of Life: Geobiology* 96, 173–180.
- **Noffke N** (2010) Microbial Mats in Sandy Deposits from the Archean Era to Today. Heidelberg, Germany: Springer Verlag.
- Noffke N (2021) Microbially induced sedimentary structures in clastic deposits: implication for the prospection for fossil life on Mars. *Astrobiology* 2021, 1–27.
- Noffke N, Beukes N and Bower D, Hazen RM and Swift DJP (2008) An actualistic perspective into Archean worlds (cyano-)bacterially induced sedimentary structures in the siliciclastic Nhlazatse Section, 2.9 Ga Pongola Supergroup, South Africa. *Geobiology* **6**, 5–20.
- Noffke N, Christian D and Wacey D and Hazen RM (2013) Microbially induced sedimentary structures recording an ancient ecosystem in the ca. 3.48 billion-year-old dresser formation, Pilbara, Western Australia. *Astrobiology* 13, 1103–1124.
- **Onstott TC** (2016) *Deep Life: The Hunt for the Hidden Biology of Earth, Mars, and Beyond.* New Jersey, US: Princeton University Press.
- Onstott TC, Ehlmann BL, Sapers H, Coleman M, Ivarsson M, Marlow JJ, Neubeck A and Niles P (2019) Paleo-rock-hosted life on Earth and the search on Mars: a review and strategy for exploration. *Astrobiology* **19**, 1230–1262.
- Orange F, Westall F, Disnar J-R, Prieur D, Bienvenu N, Le Romancer M and Défarge CH (2009) Experimental silicification of the extremophilic Archaea *Pyrococcus abyssi* and *Methanocaldococcus jannaschii*: applications in the search for evidence of life in early Earth and extraterrestrial rocks. *Geobiology* 7, 403–418.
- **Oren A** (2014) Halophilic archaea on Earth and in space: growth and survival under extreme conditions. *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences* **372**, 20140194.
- Perri E, Tucker ME and Spadafora A (2012) Carbonate organo-mineral micro- and ultrastructures in sub-fossil stromatolites: Marion lake, South Australia. *Geobiology* 10, 105–117.
- Pineau M, Le Deit L, Chauviré B, Carter J, Rondeau B and Mangold N (2020) Toward the geological significance of hydrated silica detected by near infrared spectroscopy on Mars based on terrestrial reference samples. *Icarus* 347, 113706.
- Pondrelli M, Rossi AP, Le Deit L, Fueten F, Schmidt G, van Gasselt S, Hauber E and Pozzobon R (2017) Process variability and depositional architecture of the ELDs in the Firsoff south crater (Arabia Terra, Mars). Paper presented at the *International Meeting of Sedimentology*. Toulouse, France.

- Poulet F, Bibring J-P, Mustard JF, Gendrin A, Mangold N, Langevin Y, Arvidson RE, Gondet B, Gomez C, Berthé M, Bibring J-P, Langevin Y, Erard S, Forni O, Gendrin A, Gondet B, Manaud N, Poulet F, Poulleau G, Soufflot A, Combes M, Drossart P, Encrenaz T, Fouchet T, Melchiorri R, Bellucci G, Altieri F, Formisano V, Fonti S, Capaccioni F, Cerroni P, Coradini A, Korablev O, Kottsov V, Ignatiev N, Titov D, Zasova L, Mangold N, Pinet P, Schmitt B, Sotin C, Hauber E, Hoffmann H, Jaumann R, Keller U, Arvidson R, Mustard J, Forget F and The Omega Team (2005) Phyllosilicates on Mars and implications for early Martian climate. Nature 438, 623–627.
- Quantin-Nataf C, Carter J, Mandon L, Thollot P, Balme M, Volat M, Pan L, Loizeau D, Millot C, Breton S, Dehouck E, Fawdon P, Gupta S, Davis J, Grindrod PM, Pacifici A, Bultel B, Allemand P, Ody A, Lozach L and Broyer J (2021) Oxia Planum: the landing site for the ExoMars 'Rosalind Franklin' Rover Mission: geological context and prelanding interpretation. *Astrobiology* **21**, 345–366.
- Rapin W, Meslin P-Y, Maurice S, Vaniman D, Nachon M, Mangold N, Schröder S, Gasnault O, Forni O and Wiens RC (2016) Hydration state of calcium sulfates in Gale crater, Mars: identification of bassanite veins. *Earth and Planetary Science Letters* 452, 197–205.
- Rapin W, Chauviré B, Gabriel TSJ, McAdam AC, Ehlmann BL, Hardgrove C, Meslin P-Y, Rondeau B, Dehouck E, Franz HB, Mangold N, Chipera SJ, Wiens RC, Frydenvang J and Schröder S (2018) *In situ* analysis of opal in Gale Crater, Mars. *Journal of Geophysical Research: Planets* 123, 1955–1972.
- Rapin W, Ehlmann BL, Dromart G, Schieber J, Thomas NH, Fischer WW, Fox VK, Stein NT, Nachon M and Clark BC (2019) An interval of high salinity in ancient Gale crater lake on Mars. *Nature Geoscience* 12, 889–895.
- Riding R (1991) Calcareous Algae and Stromatolites. Berlin, Heidelberg: Springer.
 Ruff SW and Farmer JD (2016) Silica deposits on Mars with features resembling hot spring biosignatures at El Tatio in Chile. Nature Communications 7, 13554.
- Ruff SW, Farmer JD, Calvin WM, Herkenhoff KE, Johnson JR, Morris RV, Rice MS, Arvidson RE, Bell III JF, Christensen PR and Squyres SW (2011) Characteristics, distribution, origin, and significance of opaline silica observed by the Spirit rover in Gusev crater, Mars. *Journal of Geophysical Research: Planets* **116**, E00F23.
- Rutledge AM, Horgan BHN, Havig JR, Rampe EB, Scudder NA and Hamilton TL (2018) Silica dissolution and precipitation in glaciated volcanic environments and implications for Mars. *Geophysical Research Letters* 45, 7371–7381.
- Sankaranarayanan K, Timofeeff MN, Spathis R, Lowenstein TK and Lum JK (2011) Ancient microbes from halite fluid inclusions: optimized surface sterilization and DNA extraction. *PLoS ONE* 6, e20683.
- Schidlowski M (2001) Carbon isotopes as biogeochemical recorders of life over 3.8 Ga of Earth history: evolution of a concept. *Precambrian Research* 106, 117–134.
- Schubert BA, Lowenstein TK, Timofeeff MN and Parker MA (2010) Halophilic archaea cultured from ancient halite, Death Valley, California. *Environmental Microbiology* **12**, 440–454.
- Sephton MA (2002) Organic compounds in carbonaceous meteorites. *Natural Product Reports* 19, 292–311.
- Shkolyar S and Farmer JD (2018) Biosignature preservation potential in Playa evaporites: impacts of diagenesis and implications for Mars exploration. *Astrobiology* **18**, 1460–1478.
- Siever R (1992) The silica cycle in the Precambrian. Geochimica et Cosmochimica Acta 56, 3265–3272.
- Sleep NH (2000) Evolution of the mode of convection within terrestrial planets. Journal of Geophysical Research: Planets 105, 17563–17578.
- Spadafora A, Perri E, McKenzie JA and Vasconcelos C (2010) Microbial biomineralization processes forming modern Ca: Mg carbonate stromatolites. *Sedimentology* 57, 27–40.
- Squyres SW, Knoll AH, Arvidson RE, Clark BC, Grotzinger JP, Jolliff BL, McLennan SM, Tosca N, Bell JF and Calvin WM (2006) Two years at Meridiani Planum: results from the Opportunity Rover. Science 313, 1403–1407.
- Squyres SW, Arvidson RE, Ruff S, Gellert R, Morris RV, Ming DW, Crumpler L, Farmer JD, Marais DJD, Yen A, McLennan SM, Calvin W, Bell JF, Clark BC, Wang A, McCoy TJ, Schmidt ME and de Souza

PA (2008) Detection of Silica-rich deposits on Mars. Science (New York, N.Y.) 320, 1063.

- Stan-Lotter H, Pfaffenhuemer M, Legat A, Busse H-J, Radax C and Gruber C (2002) Halococcus dombrowskii sp. nov., an archaeal isolate from a Permian alpine salt deposit. International Journal of Systematic and Evolutionary Microbiology 52, 1807–1814.
- Steele A, McCubbin FM, Fries M, Kater L, Boctor NZ, Fogel ML, Conrad PG, Glamoclija M, Spencer M, Morrow AL, Hammond MR, Zare RN, Vicenzi EP, Siljeström S, Bowden R, Herd CDK, Mysen BO, Shirey SB, Amundsen HEF, Treiman AH, Bullock ES and Jull AJT (2012) A reduced organic carbon component in martian basalts. *Science* (*New York, N.Y.*) 337, 212.
- Summons RE, Amend JP, Bish D, Buick R, Cody GD, Des Marais DJ, Dromart G, Eigenbrode JL, Knoll AH and Sumner DY (2011) Preservation of martian organic and environmental records: final report of the Mars biosignature working group. Astrobiology 11, 157–181.
- Sun VZ and Milliken RE (2018) Distinct geologic settings of Opal-A and more crystalline Hydrated Silica on Mars. *Geophysical Research Letters* 45, 10,221–10,228.
- Tao Y, Conway SJ, Muller J-P, Putri AR, Thomas N and Cremonese G (2021) Single image super-resolution restoration of TGO CaSSIS colour images: demonstration with perseverance rover landing site and Mars science targets. *Remote Sensing* 13, 1777.
- Tarnas JD, Mustard JF, Lin H, Goudge TA, Amador ES, Bramble MS, Kremer CH, Zhang X, Itoh Y and Parente M (2019) Orbital identification of hydrated silica in Jezero Crater, Mars. *Geophysical Research Letters* 46, 12771–12782.
- Teng W, Kuang J, Luo Z and Shu W (2017) Microbial diversity and community assembly across environmental gradients in acid mine drainage. *Minerals* 7, 106.
- Thomas-Keprta KL, Clemett SJ, McKay DS, Gibson EK and Wentworth SJ (2009) Origins of magnetite nanocrystals in Martian meteorite ALH84001. *Geochimica et Cosmochimica Acta* 73, 6631–6677.
- Thomson BJ, Bridges NT, Milliken R, Baldridge A, Hook SJ, Crowley JK, Marion GM, de Souza Filho CR, Brown AJ and Weitz CM (2011) Constraints on the origin and evolution of the layered mound in Gale Crater, Mars using Mars Reconnaissance Orbiter data. *Icarus* 214, 413–432.
- Thorseth IH, Torsvik T, Furnes H and Muehlenbachs K (1995) Microbes play an important role in the alteration of oceanic crust. *The Mantle-Ocean connection* **126**, 137–146.
- Till JL, Guyodo Y, Lagroix F, Morin G, Menguy N and Ona-Nguema G (2017) Presumed magnetic biosignatures observed in magnetite derived from abiotic reductive alteration of nanogoethite. *Comptes Rendus Geoscience* 349, 63–70.
- Treiman AH (1995) A petrographic history of martian meteorite ALH84001: two shocks and an ancient age. *Meteoritics* **30**, 294–302.
- Vago J, Witasse O, Svedhem H, Baglioni P, Haldemann A, Gianfiglio G, Blancquaert T, McCoy D and de Groot R (2015) ESA ExoMars program: the next step in exploring Mars. *Solar System Research* 49, 518–528.
- Vago JL, Westall F, Pasteur Instrument Teams LSSWG and Other Contributors, Coates AJ, Jaumann R, Korablev O, Ciarletti V, Mitrofanov I, Josset J-L, De Sanctis MC, Bibring J-P, Rull F, Goesmann F, Steininger H, Goetz W, Brinckerhoff W, Szopa C, Raulin F, Westall F, Edwards HGM, Whyte LG, Fairén AG, Bibring J-P, Bridges J, Hauber E, Ori GG, Werner S, Loizeau D, Kuzmin RO, Williams RME, Flahaut J, Forget F, Vago JL, Rodionov D, Korablev O, Svedhem H, Sefton-Nash E, Kminek G, Lorenzoni L, Joudrier L, Mikhailov V, Zashchirinskiy A, Alexashkin S, Calantropio F, Merlo A, Poulakis P, Witasse O, Bayle O, Bayón S, Meierhenrich U, Carter J, García-Ruiz JM, Baglioni P, Haldemann A, Ball AJ, Debus A, Lindner R, Haessig F, Monteiro D, Trautner R, Voland C, Rebeyre P, Goulty D, Didot F, Durrant S, Zekri E, Koschny D, Toni A, Visentin G, Zwick M, van Winnendael M, Azkarate M, Carreau C and the ExoMars Project Team (2017) Habitability on early Mars and the search for biosignatures with the ExoMars rover. Astrobiology 17, 471-510.
- Van Kranendonk MJ (2006) Volcanic degassing, hydrothermal circulation and the flourishing of early life on Earth: a review of the evidence from c. 3490–3240 Ma rocks of the Pilbara Supergroup, Pilbara Craton, Western Australia. *Earth-Science Reviews* 74, 197–240.

- Van Kranendonk MJ, Smithies RH, Griffin WL, Huston DL, Hickman AH, Champion DC, Anhaeusser CR and Pirajno F (2015) Making it thick: a volcanic plateau origin of Palaeoarchean continental lithosphere of the Pilbara and Kaapvaal cratons. *Geological Society, London, Special Publications* 389, 83.
- van Zuilen MA, Chaussidon M, Rollion-Bard C and Marty B (2007) Carbonaceous cherts of the Barberton Greenstone Belt, South Africa: isotopic, chemical and structural characteristics of individual microstructures. *Geochimica et Cosmochimica Acta* 71, 655–669.
- Viennet J-C, Bultel B and Werner SC (2019) Experimental reproduction of the martian weathering profiles argues for a dense Noachian CO2 atmosphere. *Chemical Geology* 525, 82–95.
- Vinnichenko G, Jarrett AJM, Hope JM and Brocks JJ (2020) Discovery of the oldest known biomarkers provides evidence for phototrophic bacteria in the 1.73 Ga Wollogorang Formation, Australia. *Geobiology* 18, 544–559.
- Vreeland RH, Jones J, Monson A, Rosenzweig WD, Lowenstein TK, Timofeeff M, Satterfield C, Cho BC, Park JS and Wallace A (2007) Isolation of live Cretaceous (121–112 million years old) halophilic Archaea from primary salt crystals. *Geomicrobiology Journal* 24, 275–282.
- Wacey D, Saunders M, Roberts M, Menon S, Green L, Kong C, Culwick T, Strother P and Brasier MD (2014) Enhanced cellular preservation by clay minerals in 1 billion-year-old lakes. *Scientific Reports* 4, 5841.
- Walsh MM and Lowe DR (1999) Modes of accumulation of carbonaceous matter in the early Archean: a petrographic and geochemical study of the carbonaceous cherts of the Swaziland Supergroup. Special Papers-Geological Society of America 329, 115–132.
- Walter MR and Des Marais DJ (1993) Preservation of biological information in thermal spring deposits: developing a strategy for the search for fossil life on Mars. *Icarus* 101, 129–143.

Warren JK (2016) Evaporites: A Geological Compendium. New York, US: Springer.

- Warthmann R, van Lith Y, Vasconcelos C, McKenzie JA and Karpoff AM (2000) Bacterially induced dolomite precipitation in anoxic culture experiments. *Geology* 28, 1091–1094.
- Westall F (1997) Influence of cell wall composition on the fossilisation of bacteria and the implications for the search for early life forms. In Cosmovici C, Bower S and Wertheimer D (eds), Astronomical and Biochemical Origins and the Search for Life in the Universe. Bologna, Italy: Editrici Compositori, pp. 491.
- Westall F and Cavalazzi B (2011) Biosignatures in rocks. In Thiel V and Reitner J (eds), *Encyclopedia of Geobiology*. Dordrecht: Springer, p. 482.
- Westall F and Hickman-Lewis K (2018) Fossilization of bacteria and implications for the search for early life on earth and astrobiology missions to Mars. In Kolb VM (ed). *Handbook of Astrobiology*. Boca Raton, Florida, US: CRC Press, pp. 609–631.
- Westall F and Rincé Y (1994) Biofilms, microbial mats and microbe-particle interactions: electron microscope observations from diatomaceous sediments. *Sedimentology* 41, 147–162.
- Westall F, Boni L and Guerzoni E (1995) The experimental silicification of microorganisms. *Palaeontology* 38, 495–528.
- Westall F, de Vries ST, Nijman W, Rouchon V, Orberger B, Pearson V, Watson J, Verchovsky A, Wright I, Rouzaud J-N, Marchesini D and Severine A (2006a) The 3.466 Ga 'Kitty's Gap Chert', an early Archean microbial ecosystem. In Reimold WU and Gibson RL (eds). Processes on the Early Earth, vol. Special Papers 405. Boulder, Colorado, US: Geological Society of America, pp. 105–131.
- Westall F, de Ronde CEJ, Southam G, Grassineau N, Colas M, Cockell C and Lammer H (2006b) Implications of a 3.472–3.333 Gyr-old subaerial microbial mat from the Barberton greenstone belt, South Africa for the UV environmental conditions on the early Earth. *Philosophical Transactions of the Royal Society B: Biological Sciences* 361, 1857–1876.
- Westall F, Cavalazzi B, Lemelle L, Marrocchi Y, Rouzaud J-N, Simionovici A, Salomé M, Mostefaoui S, Andreazza C, Foucher F, Toporski J, Jauss A, Thiel V, Southam G, MacLean L, Wirick S, Hofmann A, Meibom A, Robert F and Défarge C (2011) Implications of *in situ* calcification for photosynthesis in a 3.3 Ga-old microbial biofilm from the Barberton greenstone belt, South Africa. *Earth and Planetary Science Letters* 310, 468–479.
- Westall F, Campbell KA, Bréhéret JG, Foucher F, Gautret P, Hubert A, Sorieul S, Grassineau N and Guido DM (2015*a*) Archean (3.33 Ga) microbe-sediment systems were diverse and flourished in a hydrothermal context. *Geology* **43**, 615–618.

- Westall F, Foucher F, Bost N, Bertrand M, Loizeau D, Vago JL, Kminek G, Gaboyer F, Campbell KA, Bréhéret J-G, Gautret P and Cockell CS (2015b) Biosignatures on Mars: what, where, and how? Implications for the search for Martian life. *Astrobiology* 15, 998–1029.
- Westall F, Hickman-Lewis K, Hinman N, Gautret P, Campbell KA, Bréhéret JG, Foucher F, Hubert A, Sorieul S, Dass AV, Kee TP, Georgelin T and Brack A (2018) A hydrothermal-sedimentary context for the origin of life. *Astrobiology* 18, 259–293.
- Williford KH, Farley KA, Stack KM, Allwood AC, Beaty D, Beegle L, Bhartia R, Brown AJ, de la Torre Juarez M, Hamran S-E, Hecht MH, Hurowitz J, Rodriguez-Manfredi JA, Maurice S, Milkovich S and Wiens RC (2018) The NASA Mars 2020 rover mission and the search for extraterrestrial life. In Cabrol NA and Grin EA (eds). From Habitability to Life on Mars. Amsterdam, The Netherlands: Elsevier, pp. 275–308. https://doi.org/10.1016/B978-0-12-809935-3.00010-4.
- Williscroft K, Grasby SE, Beauchamp B, Little CTS, Dewing K, Birgel D, Poulton T and Hryniewicz K (2017) Extensive Early Cretaceous (Albian) methane seepage on Ellef Ringnes Island, Canadian High Arctic. GSA Bulletin 129, 788–805.
- Wirth R, Luckner M and Wanner G (2018) Validation of a hypothesis: colonization of black smokers by hyperthermophilic microorganisms. *Frontiers in Microbiology* 9, 524.
- Wogelius RA, Manning PL, Barden HE, Edwards NP, Webb SM, Sellers WI, Taylor KG, Larson PL, Dodson P and You H (2011) Trace metals as biomarkers for eumelanin pigment in the fossil record. Science (New York, N.Y.) 333, 1622–1626.
- Wray JJ (2013) Gale crater: the Mars science laboratory/Curiosity rover landing site. International Journal of Astrobiology 12, 25–38.
- Zahnle K, Ardnt N, Cockell C, Halliday A, Nisbet E, Selsis F and Sleep NH (2007) Emergence of a habitable planet. Space Science Reviews 129, 35–78.