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Unusual microbial mat-related structural diversity 2.1 billion years ago and implications for the Francevillian biota

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Abstract

The 2.1-billion-year-old (Ga) Francevillian series in Gabon hosts some of the oldest reported macroscopic fossils of various sizes and shapes, stimulating new debates on the origin, evolution and organization of early complex life. Here we document ten representative types of exceptionally well-preserved mat-related structures, comprising ‘elephant-skin’ textures, putative macro-tufted microbial mats, domal buildups, flat pyritized structures, discoidal microbial colonies, horizontal mat growth patterns, wrinkle structures, ‘kinneyia’ structures, linear patterns and nodule-like structures. A combination of petrographic analyses, scanning electron microscopy, Raman spectroscopy and organic elemental analyses of carbon-rich laminae and microtexture converge on a biological origin for these structures. The observed microtextures encompass oriented grains, floating silt-sized quartz grains, concentrated heavy
minerals, randomly oriented clays, wavy-crinkly laminae and pyritized structures. Based on comparisons with modern analogues, as well as an average $\delta^{13}C$ organic matter ($C_{org}$) composition of $-32.94 \pm 1.17\%$o (1 standard deviation, s.d.) with an outlier of $-41.26\%$o, we argue that the mat-related structures contain relicts of multiple carbon pathways including heterotrophic recycling of photosynthetically derived $C_{org}$. Moreover, the relatively close association of the macroscopic fossil assemblages to the microbial mats may imply that microbial communities acted as potential benthic $O_2$ oases linked to oxyphototrophic cyanobacterial mats and grazing grounds. In addition, the mat’s presence likely improved the preservation of the oldest large colonial organisms, as they are known to strongly biostabilize sediments. Our findings highlight the oldest community assemblage of microscopic and macroscopic biota in the aftermath of the ‘Great Oxidation Event’, widening our understanding of biological organization during Earth’s middle age.

Introduction

The Paleoproterozoic Era hosted one of the most important geochemical events in Earth’s history, marked by a measurable accumulation of atmospheric oxygen, the so-called ‘Great Oxidation Event’ (GOE) between 2.45–2.32 billion years ago (Ga). Evidence for the GOE comes from the disappearance of detrital pyrite, uraninitie and siderite from fluvial and deltaic deposits, an increase in the retention of iron in paleosols, an enrichment of Cr and U in iron formations, and perhaps most importantly, the disappearance of sedimentary sulfur isotope mass-independent (S-MIF) anomalies indicative of atmospheric $SO_2$ processing in the absence of appreciable ozone (Holland, 2002; Bekker et al., 2004; Farquhar et al., 2011; Konhauser et al., 2011; Partin et al., 2013a; Lyons et al., 2014). In the wake of the GOE, large positive excursions in the $\delta^{13}C$ of marine inorganic carbon, during the ‘Lomagundi Event’ ca. 2.22–2.1 Ga (Karhu & Holland, 1996), is believed to reflect large-scale burial of
organic matter in marine sediments (Berner, 2004; Bekker & Holland, 2012). New evidence suggests that the ‘Lomagundi Event’ ended with a drastic drop in Earth’s oxygen content (e.g. Partin et al., 2013b), perhaps related to large-scale oxidation of organic carbon (Canfield et al., 2013) and/or changes in ocean biogeochemical processes that may have inhibited primary productivity (Chi Fru et al., 2015).

The Francevillian sedimentary rocks ca. 2.1 Ga record these biogeochemical fluctuations in Earth’s oxygen accumulation dynamics through extreme excursions in the carbon cycle, in addition to hosting the oldest large colonial macroorganisms (El Albani et al., 2010, 2014). Major elements, trace metals, organic carbon and isotope analyses offer constraints on the basin geochemistry, revealing variations from the base to the top that are linked to sea level changes (Canfield et al., 2013; Bankole et al., 2016). The Lower Francevillian fluviatile rocks were deposited in oxygenated waters (Bankole et al., 2016), whereas the Upper Francevillian marine rocks were deposited in oxic, ferruginous and euxinic waters (Canfield et al., 2013). The large macrofossils are highly variable in terms of size, shape and pyritization process being represented by lobate, elongated and rod-shaped as well as disk-shaped morphotypes. The pyritization process did not fully occur in the circular disks, indicating that their organic composition differs from that of pyritized specimens (El Albani et al., 2010, 2014). In addition, some carbonaceous spheroidal microstructures have also been reported as organic-walled acritarchs that might have a planktonic origin.

Recently, microbial fossils were found in the Francevillian sequence, including putative bacteriomorphous structures (Dubois et al., 2015) and a Gunflint-type assemblage of microfossils in the stromatolitic units (Lekele Baghekema et al., 2017). The presence of colonial macroorganisms and microfossils makes the Francevillian biota unique, but interestingly, evidence of microbial mats has never been demonstrated. In this work, we describe for the first time ever microbial mat-related structures (MRS) from 2.1 Ga
siliciclastic sandstones and black shales from the FB2 Member of the Francevillian basin, Gabon. MRS are physical remains of destroyed mats and structures associated with decay, instead of well-preserved mat growth features (Eriksson et al., 2010). Their sedimentary features are preserved because microbes secrete extracellular polymeric substances (EPS), an adhesive mucilage (Decho, 2000), that provide a coherent and protective coating that is unlikely to arise in sediments without a biological input (Gerdes et al., 1993, 2000; Porada & Bouougri, 2007; Noffke, 2010).

Microbial life in the Paleoproterozoic has been poorly described (Davies et al., 2016) even though existing paleontological and biogeochemical evidence points towards the existence of significant microbial diversity previously in the Archean. Through a combination of macroscopic and microtexture analyses, petrographic, geochemical, organic elemental analyses and stable carbon isotope composition of bulk organic matter, we compare modern and fossil mat-related structures, in order to characterize the marine paleoenvironment in which the putative microbial mats formed. The data reveal the biogenicity of these structures, adding new insights into the origins of the Francevillian biota (El Albani et al., 2010; Ossa Ossa, 2010; Parize et al., 2013; El Albani et al., 2014; Ngombi Pemba, 2014; Reynaud et al., 2017). Importantly, we assessed and described the organisational, ecological and taphonomical processes that led to the development and preservation of the unique Francevillian biota.

**Geological background**

The Francevillian basin, located in the southeastern part of the Republic of Gabon (Figure 1a), is a 35,000 km² depression comprising 2.2-2.0 Ga marine sedimentary rocks that unconformably overly an Archean basement (Weber, 1969). Petrographic and geochemical analyses indicate that the Francevillian sediments were not affected by metamorphic
transformation (maximum temperature 100 °C; Gauthier-Lafaye & Weber, 1989; Ngombi-Pemba et al., 2014), resulting in the preservation of seawater composition at the time when deposition took place. Moreover, carbon isotopic and redox sensitive element compositions suggest that the ‘Lomagundi Event’ was a global event (Canfield et al. 2013).

The basin fill, composed of siliciclastic fluvial and marine deposits that are 1000–2500 m thick, is subdivided into four major lithostratigraphic formations, FA to FD (Figure 1b; Weber, 1969; Gauthier-Lafaye & Weber, 1989; Gauthier-Lafaye, 2006; El Albani et al., 2010, 2014). Fluvialite to deltaic conglomeratic sandstones, overlain by marine sandstones deposited in a tidal environment (i.e., syntectonic filling), make up the FA Formation, with the overlying sediments hosting diagenetic uranium ore deposits and the natural nuclear fission reactors of Oklo (Gauthier-Lafaye & Weber, 1989, 2003). The marine-dominated FB sequence rests unconformably on FA. The former was deposited below storm wave base during basin deepening, and is subdivided into the FB1 (a, b and c) and FB2 (a and b) subunits. The FB1a and FB1b subunits are composed of interbedded shales, sandstones and conglomerate, stacked into fining upwards packages, while the overlying FB1c subunit mainly consists of black shales with a thin iron formation, likely corresponding to a maximum flooding surface. This is then overlain by thick Mn-rich carbonates. Massive sandstone beds of the FB2a subunit, probably deposited in channels near the fair-weather wave base, are capped by thinly laminated black shales that are 5 m thick, and interbedded with thin siltstone layers (FB2b), presumably deposited by waning storm surges. Outcrops of the FB2b subunit are scarce but host the well-known colonial macroorganisms described in El Albani et al. (2010, 2014). The overlying FC Formation consists of dolomite and thickly-banded stromatolitic cherts that were deposited under shallow-water conditions (Bertrand-Sarfati & Potin, 1994). The uppermost FD Formation consists mainly of black shales, with pyroclastic material at the top deposited in a deep marine environment (Gauthier-Lafaye & Weber, 2003).
The Francevillian basin has been intensively studied because of economic interests in their uranium and manganese ore content (Gauthier-Lafaye & Weber, 1989, 2003). Consequently, various ages have been obtained for the Francevillian sediments. For example, monazites contained in Archean plutonic rocks from the Chaillu massif close to the FA Formation transition provided U-Pb ages of $2998 \pm 25$ Ma to $2621 \pm 30$ Ma (Mouélé et al., 2014). A U-Pb age of $2050 \pm 30$ Ma reported for uranium mineralization, delineates early diagenesis at the FA-FB boundary (Gancarz, 1978). Coarse-grained syenites of the N’Goutou volcanic complex, in the northern part of the Republic of Gabon, that are interlayered with rocks at the base of the FB1 sequence, yield an Rb-Sr age of $2143 \pm 143$ Ma (Bonhomme et al., 1982). It is considered that these volcanic rocks were formed simultaneously with the sedimentary rocks. Diagenetic illites from the top of the FB1b subunit have been dated at $2099 \pm 115$ Ma by using the Sm-Nd method (Bros et al., 1992). Finally, recent dating of zircons from welded tuffs near the top of the FD Formation produced a U-Pb age of $2083 \pm 6$ Ma (Horie et al., 2005).

The Paleoproterozoic Francevillian MRS were discovered in the Moulendé quarry, the same fossiliferous quarry where large colonial organisms have been reported (El Albani et al., 2010, 2014). MRS are heterogeneously distributed in the Upper FB2a subunit, as well as in the overlying laminated black shale. The latter is $5$ m thick and forms the FB2b subunit (Figure 2), covering a $\sim 20$ m thick host interval.

Methods

Sampling and sample preparation

About one hundred samples were collected in order to describe large-scale variability in morphology, structural organization and diversity from outcrops localized in one locality over several years (Figure 1). All samples were logged to provide information on their spatial and
time repartitions of the studied area (Figure 2), and then photographed at the University of Poitiers using a Nikon Europe D610 digital single-lens reflex camera equipped with a Nikon AF-S 24-120 mm f/4G ED VR lens. Polished slabs were observed using a ZEISS Discovery.V8 stereoscope combined with Axio Cam ERc 5s microscope camera. Based on this pre-screening analysis, representative samples were selected and powdered in an agate mortar for elemental and carbon isotope systematics.

Petrographic and mineralogical analysis

Petrographic and mineralogical examinations were performed by transmitted and reflected light on eight polished thin sections using a Nikon ECLIPSE E600 POL microscope equipped with a Nikon Digital Sight DS-U1 camera and NIS-Element D software for scanning observations. Seven thin sections were carbon coated and examined for textural and compositional analyses using a FEI Quanta 200 scanning electron microscope (SEM) at the University of Lille 1. The SEM, coupled to Rontec energy-dispersive spectra (EDS) for semi-quantitative mineral analysis in backscattered electron mode (BSE), was operated at 15 kV accelerating volts and a 1 nA beam current at a working distance of 10.5 mm.

Carbon analysis

Raman spectroscopy was used to determine the composition of carbon preserved within the MRS on two representative samples. Analyses were carried out at the University of Poitiers with a HORIBA JOBIN YVON Labram HR800UV, an integrated Olympus confocal microscope coupled to a Peltier-cooled CCD detector. All analyses were performed by means of 514.5 nm Ar⁺ laser of 1 mW, 200 μm confocal hole, 1800 grooves/mm grating. Data recording and treatment were done with LabSpec 5 software.

Ten representative samples, composed of five mat samples and five associated
surrounding sediments, were crushed for the determination of the elemental content of associated organic matter, as well as carbon and sulfur composition, at the University of Poitiers. Analyses were performed using a CHNS analyser, model FlashEA 1112 (Thermofisher Scientific) by flash dynamic combustion at 970 °C under a constant flow of helium. Data were recorded using Eager 300 software. A calibration curve was obtained with sulfanilamide and BBOT – 2,5-Bis(5-tert-butyl-benzoxazol-2-yl)thiophene – before each analysis. The results of carbon contents include both inorganic and organic carbon, but petrographic analyses showed that calcium carbonate and dolomite concentrations were low.

Stable carbon isotope measurements ($^{12}$C, $^{13}$C) of organic matter from 14 mat samples (from both FB2a and FB2b subunits) were conducted at the Stable Isotopes Laboratory of UMR Pegase (INRA Rennes, France), using an elemental analyser coupled with an isotope ratio mass spectrometer (IRMS) (VG Isoprime; UMR PEGASE – INRA Rennes). More than 500 mg of fine-grained powders were analysed and loaded in tin capsules for each sample. IAEA-C6 (sucrose ANU, $\delta^{13}$C = -10.63‰), USGS-24 (graphite, $\delta^{13}$C = -15.98‰), USGS-40 (glutamic acid, $\delta^{13}$C = -26.32‰) and Urea Isotopic Working Standard ($\delta^{13}$C = -38.3‰) were used as reference standards. Data are expressed as delta values (‰) relative to Vienna Pee Dee Belemnite (V-PDB). No correction was added to the measured values, and the analytical uncertainty is estimated to be lower than 0.2‰.

Facies analysis of the Moulendé Quarry

The Moulendé quarry has been examined over an area of ~ 5 ha, for a total of nine studied outcrops (Figure 3a). Among them, the F8 outcrop (Figure 3a-c) was logged in detail because of its well-exposed strata and sedimentary structures. It is the only outcrop where the FB2a-FB2b transition is observed with accompanying black shale deposits. Lithological composition, vertical facies relationships and sedimentary features allow recognition of two
depositional facies from the MRS-bearing strata.

Facies 1

Thick-bedded, coarse-grained sandstones represent the main facies of the Moulené quarry. These deposits are commonly referred to the Poubara sandstones (FB2a subunit; Figures 2, 3a-c). This facies is 15 m thick in the quarry, as well as in the closest drill core (GR5; Azzile Azzibrouck, 1986), but could be up to 100 m thick elsewhere in the basin (Weber, 1969).

Interbedded thin-bedded, coarse-grained sandstones and laminated black shales form heterolithic beds. The uppermost massive sandstone bed is laterally heterogeneous in thickness and its bedding surface is undulatory. At the top of several massive sandstone beds, sedimentary features, including load structures, sandstone clasts and water escape structures, such as dishes, are present. These beds occasionally contain dm-scale hummocky cross stratification (HCS), restricted to the upper part of the succession (Figures 2, 3d; Reynaud et al., 2017).

Facies 2

Numerous thin-bedded, parallel-laminated black shales with interbeds of siltstones lie conformably upon Facies 1. They form stacks of variable thickness in the upper part of the studied succession and are about 5 m thick. Connected to the underlying facies and approximately 0.4 m in height, a sandstone dyke cross-cuts these multilayered black shales (Figures 2, 3e). There is no evidence of fracturing along the dyke, indicating that the sediments were not consolidated prior to injection. The main component of facies 2 is characterized by rhythmic successions of mm-scale light and dark laminae of siltstones and shales. Small-scale cross-laminae are found within mm-thick interbedded siltstones. Load-shaped and flame-shaped structures forming convolute bedding (Figures 2, 3f) are observed in
interbedded medium-to-coarse-grained siltstone beds ranging from 2 to 5 cm in thickness. Interference ripples are also observed, indicating paleowaves coming from two directions (Figures 2, 3g). Lastly, a dm-scale bed containing dark-coloured convex laminae associated with cm-scale current ripples comprise the thickest interbedded coarse-grained siltstones (Figures 2, 3h).

Interpretations

The variation in thickness of the uppermost massive sandstone level (at ~15 m in the stratigraphy) may be related to detachment folds and troughs later infilled by overlying sediments (Reynaud et al., 2017). Sudden and rapid deposition of coarse-grained sediments produce load and dish structures by immediate dewatering (Reineck & Singh, 1980). Rapid sedimentation also results in sandstone dykes formed by elevated pore pressures leading to fluidization of the unconsolidated sediment. In addition, conditions that are favourable for the formation of HCS are commonly found between the fair-weather and storm wave base (Immenhauser, 2009). Various interpretations have been proposed for the paleoenvironmental setting of the FB2a subunit. They range from a deltaic paleoenvironment subjected to tidal influence (El Albani et al., 2010; Ossa Ossa, 2010), to a shoreface paleoenvironmental setting (Ngombi Pemba, 2014), to a turbidite lobe setting in waters deeper than 200 m (Parize et al., 2013). Reynaud et al. (2017) showed that the Poubara sandstones are composed of high-density sediment gravity flow deposits, emplaced during a forced regressive system tract. These authors underline that the Francevillian basin was likely isolated because of the absence of wave deposits. They also noticed the presence of linguoid and interference ripples that is consistent with shallow water settings.

Thinly laminated black shales are deposited in a relatively quiet water setting but interrupted by quickly deposited supplies of millimetre-to-centimetre-scale siltstones. These
laminae host many sedimentary structures that indicate a strong current. Reynaud et al. (2017) suggest that these sediments were deposited close to the maximum regressive surface whereas Parize et al. (2013) argue for the deposition of turbidites on an upper slope setting.

Mat-related structures (MRS) in the FB2 Member

The MRS documented here come from a 20 m thick interval of the FB2 sequence (Figure 2). Some specimens are entirely or partially pyritized, while others display a few isolated pyrite grains within the mat layers. Following the classification scheme by Sarkar et al. (2008), the Francevillian MRS are divided into two major categories: (1) mat-layer structures, representing potential in situ growth of a microbial mat (Figures 4-5, Figure S1) and (2) mat-protected structures (Figure 6), possibly formed by abiotic processes but requiring the protection of a microbial mat (Davies et al., 2016).

1. Mat-layer structures

This group of MRS represents the “mat-layer itself that may be intact, deformed or rafted” (Sarkar et al., 2008). In this work, we interpret well-preserved, intact mat growth according to the following criteria.

1a. ‘Elephant-skin’ textures

Description

‘Elephant-skin’ textures (Runnegar & Fedonkin, 1992) encompass quasi-polygonal, geometrically distinct reticulate patterns, ranging from the millimetre to centimetre scale, and bearing bulges or ridges (Figure 4a-c). They are both identified at the top of the massive Poubara sandstones and the interbedded siltstones in the black shales (Figure 2). Individual polygons are 0.3–1.3 cm in diameter, with a maximum relief of 0.2 cm.
Microscopic analyses show reticulate patterns preserved as non-homogenous dark laminae above a light, coarser material 200 to 600 µm thick (Figure 7a, f). The overlying layer consists of abundant wavy-crinkly laminae entangling 20 to 50 µm quartz grains and heavy minerals (e.g., titanium oxides, apatites and monazites); the latter suggested by the relative proportions of the elements found through SEM-EDS analysis (Figure S2), as well as transmitted and reflected light microscopy. Most of the grains in the wavy-crinkly laminae are randomly arranged within a clayey matrix, and are dominated by O, Al, Si, Mg, K and Fe (Figure S2; Table S1). Reticulate patterns are also C-rich (up to 8.52%) compared to the underlying sandstone bed, as evident from the Raman (Figure 8a-c) and organic elemental analyses (Table S2). The carbonaceous layer is punctuated by tufted microstructures, either on bulge rims or within the bulges, with similar striking morphologies to each other and where clay layers are almost vertically oriented (Figures 7f, 9a-b). Based upon the description by Noffke et al. (2013), the height/base index is 30/100 µm to 120/220 µm long, separated at regular distances of 150 – 230 µm from each other.

Interpretation

‘Elephant-skin’ textures are formed by a specific arrangement of tufts, pinnacles, bulges and smaller-scale bulges arranged into ridges. Tufts, often arising in ridge junctions or at the intersection of microbial bulges, are proposed as a particular type-feature of cyanobacterial mat filaments growing vertically towards a light source (phototaxis) (Gerdes et al., 2000; Porada & Bouougri, 2007; Bose & Chafetz, 2009). These textures are known in modern supratidal environments (Gerdes et al., 2000; Gerdes, 2007; Bose & Chafetz, 2009; Taj et al., 2014), but have also been recorded and described in some ancient shallow water environments from the terminal Proterozoic (Runnegar & Fedonkin, 1992; Gehling, 1999), and perhaps even the Archean (Flannery & Walter, 2012). However, Shepard & Sumner
(2010) debated the role of phototaxis in the formation of reticulate patterns and tufted structures. They investigated whether filamentous cyanobacteria were influenced by light to form polygonal structures or whether they were related to their own morphotype. Thus, reticulate patterns may result from an undirected motility of filamentous bacteria without the influence of light.

1b. *Putative macro-tufted microbial mat structures*

**Description**

The putative macro-tufted microbial mats are preserved as positive epireliefs that display patches of vertical bump-like structures protruding 0.05–0.2 cm on the upper surface of interbedded, coarse-grained siltstones (Figures 2, 4d).

Microscopic features seen in longitudinal thin sections reveal that the coarse-grained siltstone is draped by a 50–500 µm thin, dark, clay-rich layer with floating quartz grains (Figure 7b, g-h). These grains are clearly different in size compared to the coarser grains in the underlying sediment. Some clay particles are micaceous and oriented more or less parallel to the bedding plane. Furthermore, it can be noted that the thickness of this overlying dark layer varies along the longitudinal section. The thinner dark-coloured layer is found at the top of the microtopography, whereas the thicker one overlies the troughs.

SEM imaging reveals that the dark lamina encompasses heterogeneously-sized, 5–40 µm quartz grains and heavy minerals (Figure 9c), while clay minerals are positioned sub-vertically. In addition, EDS analyses suggest that these laminated minerals are mainly made of O, Si, Al, K elements, with minor Mg, Fe and Na (Table S1), consistent with the elemental composition of mica and illite minerals (*e.g.* Velde & Meunier, 2008).

**Interpretation**
A similar macro-tufted microbial mat was recorded from a 2.0 Ga paleodesert environment where 0.2 cm high mound-like structures are randomly arranged on a fine-grained sandstone (Simpson et al., 2013). The mechanism behind the development of this feature in the sedimentary rock record remains unclear. However, by changing the polarity of their filamentous structure, the growth of a microbial biofilm is able to produce similar bump-like structures in modern sedimentary environment (Gerdes et al., 2000; Gerdes, 2007; Bose & Chafetz, 2009; Taj et al., 2014). This pattern may also result from the direct filling of the pre-existing microtopography by mat-building organisms as they grow and expand (Simpson et al., 2013). The smooth but bumpy morphology is likely enhanced by synsedimentary loading and post-depositional compaction, and so this bio-sedimentary expression could be viewed as a combined microbial and compaction feature.

1c. Domal buildups

**Description**

Domal buildups are preserved as positive bed-top features on bedding surfaces of black shales (Figure 2). They are characterized by 0.1–0.4 cm convex, domal structures projecting upwards from the bedding plane. Distinctive features include a cluster of low mound-shaped structures, 0.4 cm in diameter, nearly equal in all instances and without any central depressions (Figure 4e-f). Alternatively, they can be isolated structures with a diameter ranging from 0.7 cm to 1.5 cm that seem to reveal a cauliflower-like texture associated with extensive positive-relief, wrinkled structures (Figure 4g). Circular disks belonging to the oldest large colonial organisms (El Albani et al., 2014) seem closely associated with these specific structures. Both isolated and clustered domal structures have been shown to be pyritized (Table S2). Bulk sulfur level of up 22.16% within the domal structures is associated with a carbon content <0.53%. There are also flat, pyritized mat-related structures without any
particular distinctive features that tend to develop close to the clustered domes (Figure 4f, Figure S1).

Thin sections reveal a well-defined alternation of clay and silt layers beneath the domal structures (Figure 7c). Most of the quartz particles from the underlying sediment are laminated and/or have a length of grain-to-grain contact of mostly 20–80 μm. None of these grains were found in the topmost dark, clay layer. Furthermore, the domal structures typically show an indication of internal convex lamination between all pyrite grains (Figure 7i), which reflects the position of the domes in epirelief.

Interpretation

Domal buildups have been reported from fluvial and fluviodeltaic settings in Mid-Proterozoic (Garlick, 1988; Schieber, 1998, 1999; Sheldon, 2012; Wilmeth et al., 2014) to shallow water settings inLate Cambrian times (Bottjer & Hagadorn, 2007). The latter authors proposed that they “may represent different life histories, biological affinities, and/or histories of interaction with the physical environment”. Previous studies have also invoked a distinct mat growth pattern (Wilmeth et al., 2014) and inferred that domal structures are built by an upward growth of microbial mats. This might reflects competition for light (Garlick, 1988; Schieber, 1999). The δ¹³C₉₉ isotopic signatures of Mesoproterozoic domal-like structures in Copper Harbor Conglomerate (Upper Peninsula of Michigan) are attributed to photosynthesis (Sheldon, 2012).

EPS possess chemical properties that protect cells by creating sharp geochemical gradients (Decho, 2000), while the anaerobic decay of the buried mat material facilitates the precipitation of diagenetic minerals such as pyrite, depending on sulfate and iron bioavailability. If this diagenetic mineralization follows individual layers within the EPS demarcated boundaries, or if it is associated with specific features, the former presence of
microbial mats are overprinted and preserved as a mineralized signature (Schieber, 1999). Thus, sulfate-reducing microorganisms probably used the carbonaceous material from the domal buildups as their electron donor, thereby releasing hydrogen sulfide to react with available iron. This mechanism is supported by the recurrent low carbon to high sulfur contents in the materials. Analyses of the sulfur isotope contained in pyrite from the Francevillian series measured by Secondary Ion Mass Spectrometry (SIMS) further revealed negative $\delta^{34}$S values associated with widespread microbial decay, linked to the activities of the sulfate-reducing microorganisms (El Albani et al., 2014).

1d. Discoidal microbial colonies

Description

A discoidal microbial colony (Grazhdankin & Gerdes, 2007) is a term for a mat-layer that includes disk-shaped, internal, centimetre-scale features (Figure 5a-g). Two major distinct discoidal microbial colony-like morphologies occur on bedding planes in the black shale unit (Figure 2). The first is characterized by centimetre to the decimetre scale clusters of outward-convex, rod-shaped structures with internal pyritized concentric bands, possessing a series of ~0.4 cm wide light and dark zones (Figure 5a-b). This specific feature does not exhibit any relief. The second variety is made up of well-marked circular bodies represented by sharp outlines separating the specimens from the surrounding black shale (Figure 5c-g). The disk-shaped colonies display slight internal concentric zonations, wrinkle features or a lack of any specific morphological characteristics. A variety of the specimens have a 1.1–1.7 cm high disk-shaped relief with concentric rims restricted to the edges, which may present themselves as cauliflower-like patterns varying between 3.3 and 8.2 cm in diameter (Figure 5c-e). There are numerous distinct large euhedral pyrite crystals on these structures. Less common are disk-shaped bodies of 2.3 cm in diameter and represented by a minor relief, internal
millimetre-scale wrinkles, and devoid of sharp outlines (Figure 5f). Other observable
discoidal structures are relatively small (~ 0.5 cm diameter) that do not have particular
features (Figure 5g). However, these are the only structures that are entirely pyritized by tiny
pyrite crystals.

Interpretation

Previous studies have shown that a wide range of variability exists in the morphology and
organization of modern discoidal microbial colonies in natural environments (Gerdes et al.,
1993; Banerjee et al., 2014; Sarkar et al., 2014) as well as under laboratory conditions
(Matsushita et al., 2004). Concentric zonations with a series of light and dark bands represent
a particular variety of discoidal microbial colony known as ‘fairy rings’. Modern ‘fairy ring’
structures are ubiquitous in salt marshes (Gerdes et al., 1993; Gerdes, 2007) and in shallow
water environments (Banerjee et al., 2014). The concentric ring-shaped structures of the
Francevillian series discoidal structures clearly display the same striking morphologies. All
modern examples indicate that filamentous cyanobacteria and diatoms, which are embedded
in EPS and crowned by micro-pinnacles, form these concentric ring-shaped structures. This
arrangement is likely not random but represents an oriented behaviour that might be triggered
by external environmental factors (Gerdes, 2007). So far, the generally accepted mechanism
of formation involves escaping gas bubbles from the substrate, which may cause concentric
wave propagation (Gerdes et al., 1994). Subsequently, chemotactic responses initiated by
adapted filamentous cyanobacteria may form the concentric organizational patterns. Presently,
the oldest discoidal microbial colonies that have been described are from Mesoproterozoic
and Neoproterozoic shallow submarine environments (Grazhdankin & Gerdes, 2007;
Banerjee et al., 2010, 2014; Sarkar et al., 2014), where pyritized filamentous structures are
sometimes preserved (Grazhdankin & Gerdes, 2007).
Horizontal mat growth pattern

Description

A unique, outward-convex, spindle-shaped structure extending over 34 cm is developed on the bedding plane surface of the black shale unit (Figures 2, 5h). The structure appears similar to the spindle-shaped ‘fairy ring’ structures described above. However, this structure is represented by tiny, 0.1 cm tall, arched ridges, rising horizontally, following a well-defined geometric pattern. These sharp-crested ridges of 0.15 cm in width are regularly spaced, commonly dissected and slightly undulated. It appears that the laminae convexity of the whole structure is unevenly distributed, ranging from 55 to 86 degrees.

Interpretation

The horizontal mat growth pattern could be related to ‘fairy ring’ structures with which the pattern shares some similarities (*e.g.* outward-convex, spindle-shaped, sharp-crested ridges). Nevertheless, there are no alternations of light and dark zones. Processes implicated in the formation of ‘fairy ring’ structures may be involved here, but topography is added. The orientation of the wave propagation and subsequent millimetre-sized ripples may be induced by gravity. Then, pinnacle-forming organisms would interact with the nutrient front by chemotactic responses to colonize the substratum, producing the distinct morphology preserved in the sedimentary record. Alternatively, localized mat failure and strain induced by gravity, waves or currents could also explain the morphology as a mechanical deformation of biogenically stabilized sediment. Although it has not been suggested in other studies, morphological features including a point of origin, persistent “growth” in one vector, and sharp boundaries could also be explained by an underwater seep that is confined by a biomat and promotes mat growth in a downslope direction. In any case, the parallel arrangement of
ridges, as well as the regularity of components in the horizontal mat growth pattern, suggests a biotic origin.

2. Mat-protected structures

This category of mat-related structures arises from structures that probably require mats for their preservation, the excepting being wrinkle marks.

2a. Wrinkle structures

Description

Based on morphology, three types of wrinkle structures are categorized here: wrinkle marks, parallel wavy wrinkle structures and cross-cutting wrinkle structures. Wrinkle marks are represented as laterally continuous or discontinuous, slight, outward-convex positive relief on bedding plane of the black shales (Figures 2, 4h).

Parallel wavy wrinkle structures are preserved as sinuous, continuous ridges separated by parallel, narrow valleys extending for less than 10 cm on bedding surfaces of sandstones and interbedded thin-to-medium-bedded siltstones (Figures 2, 6a-b). The ridges have rounded tops and steep sides, with an individual ridge elevation of 0.1 cm and a spacing of 0.4 cm.

Organic elemental analysis of one sample from the black shale unit does not reveal significant differences in carbon content between the parallel wavy wrinkle structures and the surrounding sediment (Table S2). However, microscopic analysis of thin sections, cut perpendicularly through the wrinkles, indicates that a dark-coloured layer caps the coarse, silt-sized rippled structures. Further down, the sediment displays quasi-planar laminations (Figure 7d). For the length of the section, the topmost carbonaceous layer is similar in thickness and does not onlap the rippled bed, but instead reproduces its topography (Figure 7j). Oriented quartz grains trapped within the carbonaceous layer (Figure 7j) appear to have the same grain
size as those found in the rippled structure. Tiny floating grains are also embedded in the clay-sized fractions. Sheets of various clay particles entangle several heavy minerals, mainly titanium oxides (Table S1).

Cross-cutting wrinkle structures (Banerjee & Jeevankumar, 2005) are characterized by bifurcating and frequently interconnected asymmetric ridges with intervening sub-parallel troughs on the bedding surface of black shales (Figures 2, 6c). The rounded ridges are 0.1 to 0.2 cm high, 0.4 to 0.5 cm wide, and their spacings vary from 0.4 to 1 cm. A few dome-like structures of 0.5 to 0.6 cm are close to these cross-cutting wrinkle structures (Figure 6c).

Interpretation

Wrinkle structures are one of the most common sedimentary features related to microbial mats. They are found over a large paleogeographic range, including the deep-sea (Buatois & Mángano, 2003; Flood et al., 2014), shallow marine (Hagadorn & Bottjer, 1997; Banerjee & Jeevankumar, 2005; Sarkar et al., 2006; Porada & Bouougri, 2007; Mata & Bottjer, 2009; Banerjee et al., 2014; Buatois et al., 2014; Sarkar et al., 2014, 2016; Yang et al., 2017), fluvial and lacustrine (Chu et al., 2015, 2017), and even continental environments (Simpson et al., 2013). Some researchers claimed that relative microbial community growth correlates to with irregular surface of wrinkle structures (Hagadorn & Bottjer, 1997; Banerjee & Jeevankumar, 2005; Sarkar et al., 2006, 2014).

Loading and dewatering processes have also been proposed for wrinkle generation (Noffke et al., 2002). A recent study has proposed that abiotic mechanisms are unable to create millimetre-scale ripples (3–15 mm wavelengths) because waves with small orbital amplitudes are not strong enough to mobilize sand and silt grains (Mariotti et al., 2014). In wave tank experiments, millimetre-scale ripples were perfectly reproduced by means of microbial aggregates rolling along the substrate and transporting grains. The preservation of
such structures requires burial by fine material that may be represented by the biostabilization
of a bacterial community. The dark-coloured layer overlapping the silt-sized rippled structures
(Figure 7d, j) is probably induced by EPS that acts as a trap for particles from the surrounding
environment (Hagadorn & Bottjer, 1997). Oriented grains are result of a growing biofilm that
envelops, lifts, rotates and orientates the grains (Noffke et al., 1997, 2001). By combining
morphological description and petrographic study, it appears that wrinkle structures are
polygenic. Both mat-layer and mat-protected structures are therefore most likely represented
in this case.

2b. ‘Kinneyia’ structures

Description

‘Kinneyia’ structures are characterized by clearly distinct, short, sinuous ridges and troughs of
0.2 cm in width and less than 0.1 cm in height on the bedding surface of the black shale unit
(Figures 2, 6d). These structures fit with the definition of ‘Kinneyia’ as proposed by (Porada
et al., 2008), i.e., they are “comparatively short, curved, frequently bifurcating, flat-topped
crests, 0.5–1 mm high and 1–2 mm wide, which are separated by parallel, round-bottomed
depressions. The crests are usually steep sided and may run parallel”.

Interpretation

The origins of “Kinneyia” structures have long been amongst the most problematic to
understand. From their initial description as fossil algae (Walcott, 1914) to the present day
where the term is now used to describe MRS, they have been interpreted in various ways
(Hagadorn & Bottjer, 1997; Noffke et al., 2002; Porada & Bouougri, 2007; Porada et al.,
2008; Thomas et al., 2013; Mariotti et al., 2014). However, few hypotheses properly explain
how the underlying sediment is affected (Davies et al., 2016). Recently, ‘Kinneyia’ structures
were successfully reproduced in wave tank experiments using microbial aggregates (Mariotti
et al., 2014), although the sharply defined morphology of the depressions preserved might
suggest that gas bubbles formed beneath the biomat could also be a possible mechanism
(Pfluger, 1999). It has also been demonstrated that ‘Kinneyia’ is a polygenetic texture (Davies
et al., 2016), but the Francevillian ‘Kinneyia’ structures do not rule out formation by biotic
mediators.

2c. Linear patterns

Description

Linear patterns are characterized by 0.1–0.3 cm high, parallel, linear ridges, slightly undulated
on the bedding surfaces of massive sandstones and interbedded siltstones (Figures 2, 6e, g).
They are of varying size, ranging from 2 to 8 cm in length and 0.3 to 0.7 cm in width. Around
these linear ridges, the surface morphology is not uniformly flat but bears micrometric spots
that seem to be embedded in the matrix (Figure 6f). Furthermore, Raman spectrometry
indicates this dark-coloured layer is carbon-rich compared to the underlying sediments
(Figure 8d-e).

Microscopic observations reveal that the 100 to 300 μm thick, carbon-rich layer lies on
lighter, sand-sized quartz grains and consists of multiple, well-defined sheets of clay minerals
(Figure 7e, k). SEM imaging shows that numerous small quartz grains and heavy minerals,
such as apatite, titanium oxide and zircon, are embedded in the clay matrix (Figure 9d-e). This
matrix is formed by O, Si, Al, Mg, Fe and K elements, i.e. chlorite and illite, as expressed by
EDS analyses (Table S1). Furthermore, sand-sized quartz grains are observed inside the
ridges as well as the surrounding sediment but are separated by a clayish undulated layer
(Figure 9e). The quartz grains inside the ridges and those from the host rock appear randomly
oriented.
Interpretation

According to the identification of linear patterns in modern tidal flats by (Porada & Bouougri, 2007) “straight to irregular ridges are developed at distance of 1–2 cm and are locally interrupted or reduced to faint lines on the flat mat surface”. Towards the water line, oriented bacterial filaments dominate these structures. In experimental studies, Shepard & Sumner (2010) observed linear ridges of up to 15 cm long made of cyanobacterial filaments running parallel to the incident light. Moreover, it has been proposed that a faint rippled surface may imply a linear pattern because of the preferential microbial growth following slight ridges (Bouougri & Porada, 2007; Gerdes, 2007; Porada & Bouougri, 2007).

The presence of carbonaceous material within the dark-coloured layer is congruent with a biotic origin. Microtextures, such as tiny quartz particles and heavy minerals embedded in the matrix, are caused by the trapping and binding processes related to microbial activity (Gerdes, 2007; Noffke, 2010). Individual grains bound in the organic matrix can be compared to those observed with laboratory-grown cyanobacteria. These are described as oriented grains that are pushed upwards during their growths (Noffke et al., 2001; Noffke, 2010).

On a macroscopic level, the Francevillian MRS strongly resemble the linear structures described above, commonly equivalent in shape but slightly smaller in size. However, no microscopic features were described until now. Taking microtextures into account, it seems unlikely that bacterial orientation is capable of mobilizing large amount of sediments to form ridges. Microbial shrinkage caused by a period of subaerial exposition, is also a possibility (e.g. Chu et al., 2017; Kovalchuk et al., 2017), although the surface morphologies and microtextures are not consistent with this process. Also, dilational strain on a mildly dipping sediment surface could explain these features, but their orientations are irregular and their distributions are sporadic. Liquefaction underneath the microbial mat, due to a rise in pressure
(e.g. Porada et al., 2007), could have been the cause of these particular ridges but neither sediment rising nor upward-facing microbial laminae beneath the ridges have been observed. Finally, linear patterns may result from two growth periods interrupted by small and rippled sedimentary structures (Figure 9e). This condition might be the best explanation for microbial laminae throughout the linear ridges, and the second mat growth period may have followed the ridge orientation as previously thought (Bouougri & Porada, 2007; Gerdes, 2007; Porada & Bouougri, 2007). Indirectly, linear patterns are considered to be mat-protected structures.

2d. Nodule-like structures

Description

Nodule-like structures are characterized by an elongated shape approximately 10 cm long and an irregular surface topography on the bedding surfaces of black shales (Figures 2, 6h). The surfaces of nodule-like structures comprise several millimetre-scale, tiny, crinkled ridges that are randomly distributed. Composed of silt-sized grains and capped by a dark-coloured wrinkle layer, the nodule-like structure was observed in a polished slab.

Interpretation

Similar nodular to biscuit-like surface structures have been observed in laboratory-cultured bacteria and in modern supratidal settings (Gerdes, 2007). This atypical morphology is thought to be produced by a relative abundance of coccoid cyanobacteria. However, the Francevillian nodular structures possibly reflect mat-protected structures instead of mat growth structures since the internal part of the nodules is not composed of organic matter but rather of quartz particles.

3. Isotopic analyses
The $\delta^{13}$C values (V-PDB) of the organic fraction, measured on different mat morphologies in both sandstone and black shale facies range from -30.67‰ to -41.26‰ (Table 1). These values are similar to previous determinations on the bulk $\delta^{13}$C of the organic matter within the FB2 sequence, varying between ca. -35‰ to -30‰ (Gauthier-Lafaye & Weber, 2003; Canfield et al., 2013).

**Discussion**

**Biogenicity**

The biogenicity criteria reviewed in Noffke (2009) and Wacey (2009) establishes the investigated mat-like structures to be of a microbial mat origin, having formed on the shallow marginal self-environment of the 2.1 Ga Francevillian basin (Reynaud et al., 2017). The sedimentary facies on which the structures are identified must not have undergone metamorphism beyond greenschist grade (Noffke, 2009). This is in agreement with the absence of metamorphic overprint in the Francevillian facies (maximum temperature 100 °C; Gauthier-Lafaye & Weber, 1989; Ngombi-Pemba et al., 2014). In terms of shape and size, the Francevillian MRS are a perfect match to those described in the Precambrian and the Phanerozoic (Figures S3-S4) in accordance with the size distribution between modern and ancient MRS, which should be comparable (Noffke, 2009). Typically, the taphonomic preservation window of MRS in the sedimentary record is valid, but not restricted by ecological setting (Davies et al., 2016). Noffke (2009) suggested that most MRS are related to photoautotrophic mats formed in well-lit shallow marine environments, preferentially on fine sand deposits, even though recent studies argued that such features can also arise on deep marine sediments that do not receive light (Davies et al., 2016). Regardless, our samples originated from a shallow marine setting associated with rapid sand flow deposits within a well-lit portion of the 2.1 Ga Francevillian continental shelf (Reynaud et al., 2017).
Microtextural composition of the analysed fabric further support the biological trapping, binding (or ‘flypaper’ effect) and orientation of grains induced by the growth and development of microbial mats and their hydrologically-controlled interaction with sediments. The Francevillian MRS express this biological process by containing clay particles, floating grains of silt-sized quartz and concentrated heavy minerals, as well as by wavy-crinkly laminae (Figures 7f-k, 9, Figure S2; Table S1). These form well-defined organo-sedimentary structures caused by microbial baffling and trapping. The randomly oriented clay minerals in mat laminae suggest that they were trapped on the mat surface whereas laminated clay particles suggest a probable alignment by compaction (Schieber, 1998). Oriented grains reflect a particular microtexture (Figures 6f, 7j), while grain size matches that of the underlying substratum. It is commonly thought that these particles were dragged upwards by cyanobacterial mat growth (Noffke et al., 1997, 2001).

In addition, geochemical evidence suggests bacterially induced biological processes characterized by carbonaceous material enriched in light carbon (Figure 8; Table 1) and pyritized structures (Figures 4e-g, 5a-e, g, 7c, i) that are depleted in heavy sulfur. The latter points to diagenetic mat destruction through burial decay (Noffke, 2009; Noffke et al., 2013) and anaerobic respiration of that organic carbon by sulfate-reducing microorganisms (El Albani et al., 2014; Hill et al., 2016).

Paleoenvironmental interpretations and implications

Modern mat-related structures are mostly described in carbonate and siliciclastic environments but few have been described from shale deposits. The presence of black shales with large amounts of organic matter deposited in the photic zone may arise in restricted to isolated basins (Schwark & Frimmel, 2004). Some anoxygenic photosynthetic bacteria even prospered in these paleoecological conditions and their high productivity may be related to the
availability of essential nutrients. For any MRS-bearing rocks, the hydraulic pattern must be moderated with a low sedimentation rate to promote the development of microbial communities on a substrate (Schieber, 1999; Gerdes et al., 2000; Gerdes, 2007; Noffke, 2009, 2010). As an indicator of paleo-hydrological conditions, clay minerals and silt-sized grains within mat layers, are taken to represent currents strong enough to move thin particles but insufficient to transport sand-sized grains (Schieber, 1998; Noffke, 2009). Silt-sized sedimentary particles and heavy minerals can also be baffled and trapped by cyanobacterial filaments moving vertically upwards in order to escape being buried by the small-sized particles, as well as their need to reach optimal light conditions for growth (Noffke, 2009, 2010).

In this study, some of the mat-like structures, including the ‘elephant-skin’ morphotypes, the tufted structures, and the linear patterns, tend to be associated with silty lenses (Figure 2), suggesting that the baffling and grain trapping processes were operating in potential microbial mat-dominated environments at 2.1 Ga. Further, the comparable association of several MRS to the distribution of modern and fossil analogues have been used to improve the reliability of paleoenvironmental marine interpretations, including physical processes such as desiccation and erosion and biological activity (Bose & Chafetz, 2009; Noffke et al., 2013; Banerjee et al., 2014; Sarkar et al., 2014). Importantly, the absence of desiccation and erosion-induced MRS in the Francevillian rocks suggest a quiet underwater environment from the time of deposition to when the sediments were lithified and buried.

The Paleoproterozoic Francevillian MRS possess a wide spectrum of morphologies that can be related to photoautotrophic microbial assemblages. Indeed, discoidal microbial colonies in the modern environments and domal buildups in ancient settings are results of cyanobacterial chemotaxis or phototaxis behaviours (Gerdes, 2007). The biological mechanisms leading to oriented grains has been reproduced with cultures of cyanobacteria
Reticulate patterns and tufted microbial mats have striking morphologies with modern analogues built by filamentous cyanobacteria (Figure S3a; Gerdes et al., 2000; Gerdes, 2007; Bose & Chafetz, 2009; Taj et al., 2014). It is thought that their formation implies a phototactic behaviour (Gerdes et al., 2000; Reyes et al., 2013), whereas laboratory-cultured filamentous cyanobacteria show a dependency on oxygen concentration (Sim et al., 2012). It has also been proposed that the undirected gliding motility of filamentous bacteria species may form these structures (Shepard & Sumner, 2010). Consequently, all bacteria with highly motile filaments are believed to be able to display these morphologies. Filamentous sulfur-oxidizing bacteria may also have the ability to produce ‘elephant-skin’-like and tuft-like structures (Flood et al., 2014). However, research by genomic comparisons reveals that these phenotypic traits were most likely inherited by horizontal gene transfers from the cyanobacteria (Flood et al., 2014). In other words, those aerobic chemolithoautotrophs may not have been in existence at 2.1 Ga. In addition, recent studies in a perennially ice-covered Antarctic lake have shown the specific assemblage of a photosynthetic microbial ecosystem (Sumner et al., 2015; Jungblut et al., 2016), with cyanobacteria forming tuft-related structures exclusively found in the oxic zone where irradiance is at its highest, whereas flat bacterial mats of anoxygenic photoautotrophs are restricted to the deeper euxinic zone experiencing lower irradiance. Thus, ‘elephant-skin’ and tufted structures may possibly have a link with oxygenic photoautotrophic microorganisms. Importantly, Flannery & Walter (2012) and Homann et al. (2015) thought that cyanobacteria are the only microorganisms capable of producing vertical structures or tufts.

Geomicrobiological implications

It is well established that carbon fixation by autotrophic organisms preferentially incorporates light $^{12}$C over heavy $^{13}$C isotopes in biomass (Schidlowski, 1988, 2001). In this regard, the
\[ \delta^{13}C_{\text{org}} \] values ranging from -30.67 to -41.26‰ (average -32.94 ± 1.17‰) reported here are within the expected range for autotrophic carbon fixation (Schidlowski, 1988, 2001; Berg et al., 2010). The typical \( \delta^{13}C_{\text{org}} \) values generated by the widespread activity of ribulose 1,5-bisphosphate carboxylase/oxygenase (RuBisCo), common in oxyphototrophic cyanobacteria have average values from -20 to -30‰. These values are less negatively fractionated than anoxygenic phototrophs (Quandt et al., 1977; McNevin et al., 2007; Berg et al., 2010), while more negative values < -30‰ are related to autotrophic carbon fixation in the reductive acetyl CoA pathway (i.e., methanogenesis). In the case of the latter, acetoclastic methanogenesis diagenetically supplies isotopically light \({}^{12}\text{C}\) methane to anaerobic oxidation of methane (AOM), the latter often comprising a consortia of sulfate reducers and methanotrophs (Conrad et al., 2010). Consequently, autotrophic carbon fixation via acetyl CoA pathway, combined with anaerobic oxidation of fixed \( C_{\text{org}} \) would effectively lead to deposition of residual \( C_{\text{org}} \) enriched in light \( {}^{12}\text{C} \) in the range found in this study. Such fixation of CO\(_2\) coupled with diagenetic recycling of phototrophically derived \( C_{\text{org}} \) would have inevitably resulted in the sequential overprinting of light \( \delta^{13}C \) in buried biomats. The bulk \( \delta^{13}C_{\text{org}} \) signatures < -30‰ are, therefore, most parsimoniously interpreted to represent a mixed isotopic signal resulting from the activities of various primary producers and heterotrophs.

The FB2b subunit hosts the first known multicellular organisms closely associated with biomats (Figures 2, 10). Burrows, trails and resting traces of metazoans are often closely associated with biomats in the past, being interpreted as sophisticated feeding behaviours (e.g. Buatois & Mángano, 2012; Pecoits et al., 2012; Buatois et al., 2014; Meyer et al., 2014; Chu et al., 2015). In modern environments, photosynthetic bacterial mats create thin O\(_2\)-rich layers, thus providing benthic O\(_2\) oases for macroorganisms that may mine mat layers for unexploited nutrients and O\(_2\) (Gingras et al., 2011). It is perhaps not a coincidence then that the biomats and macroorganisms belong to the same strata. Moreover, the generation of
microenvironments due to the chemical properties of EPS (Decho, 2000) may permit soft-
tissue mineralization, providing further protection against degradation (Sagemann et al.,
1999). Thus, the biomats may have stabilized the depositional surfaces and sheltered the
macroorganisms, allowing them to become imprinted into the rock record. Interestingly, the
large colonial organisms associated with bacterial communities are only known from the
FB2b rocks. Although other black shale facies are recorded in the Francevillian basin (e.g.,
the ~ 2.08 Ga FD black shale formation; Figure 1b), they were deposited in deeper
environments beyond the euphotic zone (Canfield et al., 2013).

Conclusion
Mat-related structures of the 2.1 Ga Francevillian series exhibit ten types of surface
morphologies, providing a new window into the highly diversified Paleoproterozoic microbial
life at that time. Mats are preserved in excellent conservation conditions from a 20 m thick
interval of sandstone and black shale facies. Microtexture analyses provide strong evidence in favour of mat-colonized sediment. Oriented
grains, floating grains, heavy mineral concentrations, randomly oriented clays, pyritized
structures and wavy-crinkly laminae all reflect the growth of microbial communities.
Comparisons with ancient and modern analogues, as well as stable carbon isotope analyses,
suggest growth within a palaeoenvironmental settings corresponding to the euphotic zone,
likely less than 100m deep. Associations between Paleoproterozoic, large colonial organisms
and mats may be similar to interactions of Ediacaran early metazoans and microbial carpets
where O₂-producing cyanobacterial mats may explain this specific pattern. In addition,
microbial mats may have played a major role in sediment biostabilization, fostering the
preservation of complex macroorganisms that represent the first ecosystem comprising
microbial biofilms and large colonial life forms.
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Figure captions

**FIGURE 1** Geological map and lithostratigraphic column. (a) Geological map of the Francevillian basin. The studied quarry is Moulendé (green star). Geological map adopted from (Bouton et al., 2009). (b) Synthetic lithostratigraphy of the Francevillian series. Four sedimentary units rest unconformably on Archean rocks. The red star indicates the detailed lithostratigraphic column observed in the Moulendé quarry (Figure 2).

**FIGURE 2** Detailed lithostratigraphic column. Composite columnar section of the Moulendé quarry in the FB2 unit showing the vertical distribution of ten representative types of mat-related structures (MRS) and sedimentary structures (SS).

**FIGURE 3** Plane view and outcrop pictures of sedimentary facies in the Moulendé quarry. (a) Representation of the quarry from plane view. Red box indicates the main studied outcrop in B (F8). F = outcrops. (b) Details of the bedding geometry at the transition between massive sandstone beds and thinly laminated black shales. (c) Closer view of B. (d) Cross-section view of decimetre-scale hummocky-cross stratifications (HCS), FB2a subunit. (e) Sandstone dyke, FB2a - FB2b transition. Coin diameter: ~ 2 cm. (f) Cross-section view of convolute structures, FB2b subunit. (g) Bedding plane view of interference ripples, FB2b subunit. (h) Longitudinal view of dark-coloured convex laminae associated with cm-scale foreset beds, FB2b subunit.
FIGURE 4 Microbial mat structures in the Francevillian B Formation (FB2): Mat-layer structures. (a-c) ‘Elephant-skin’ textures. (d) Putative macro-tufted microbial mat. (e-f) Clustered domal buildups and flat pyritized microbial structure (red arrow). Macrofossil specimens (white arrows). (g) Isolated domal buildups. (h) Wrinkle marks.

FIGURE 5 Microbial mat structures in the Francevillian B Formation (FB2): Mat-layer structures. (a-b) Discoidal mats likely representing ‘fairy ring’ structures. (c-e) Disc-shaped mats that display a cauliflower-like pattern. (f) Disc-shaped mat with internal wrinkle structures. (g) Small pyritized circular bodies. (h) Horizontal mat growth pattern.

FIGURE 6 Mat-related structures in the Francevillian B Formation (FB2): Mat-protected structures. (a-b) Parallel wavy wrinkle structures. (c) Cross-cutting wrinkle structures. (d) ‘Kinneyia’ structure. (e) Linear pattern. Dashed red box indicates the position of the magnification in f. Red arrow shows the location where the spectrometry Raman was performed. The Raman spectra is visible in Figure 8d. (f) Micrometric spots interpreting as oriented grains. (g) Linear patterns with several parallel ridges. (h) Nodular-like structure.

FIGURE 7 Optical photomicrographs of mat-related structures. (a) Transmitted thin section of ‘elephant-skin’ texture. Dashed red box denoting area magnified in f. (b) Transmitted thin section of putative macro-tufted microbial mat. Dashed red boxes denoting areas magnified in g and h. (c) Transmitted thin section of an isolated domal buildup. Dashed red box denoting area magnified in i. (d) Transmitted thin section of parallel wavy wrinkle structures. Dashed red box denoting area magnified in j. (e) Transmitted thin section of a linear pattern. Dashed red boxes denoting areas magnified in k and Figure 9e. (f) Tufted microstructures and wavy-crinkly laminae. Dashed red box denoting area magnified in Figure 9a. (g-h) Thickness variation across the mat layer with floating grains embedded by clays (red arrows). Mica (white arrows). Dashed red box denoting area magnified in Figure 9c. (i) Reflected magnified thin section of an entirely pyritized dome. An internal convex lamination is indicated by dashed red lines. (j) Clay laminae do not onlap the rippled siltstone bed but rather well follow its topography. Oriented grains (arrows). (k) High amount of quartz particles (arrows as example) within dark-coloured laminae. Dashed red box denoting area magnified in Figure 9d.

FIGURE 8 Polished slab of ‘elephant-skin’ texture and Raman spectra of both ‘elephant-skin’ texture and linear pattern. (a) Polished slab in cross-section perpendicular to bedding plane. Non-homogenous dark layer preserved above a pronounced boundary. Red arrow and white arrow indicate Raman spectra in b and c, respectively. (b) Representative Raman spectra of the microbial mat within bulges. It shows the presence of two Carbon peaks (“C”) at 1336 cm⁻¹ (the “D1” disordered peak) and 1603 cm⁻¹ (the “G” graphite peak). (c) Typical Raman spectra of sandstone with quartz (“Q”) peaks. (d) Representative Raman spectra of mat layers of linear pattern indicated in Figure 6e. It shows the presence of three carbon peaks (“C”) at ~1170 cm⁻¹ (“D4” disordered peak), 1344 cm⁻¹ (the “D1” disordered peak) and 1603 cm⁻¹ (the “G” graphite peak). (e) Typical Raman spectra of host sediment of linear pattern, with quartz (Q) peaks and very small intensities of “C” peaks.

FIGURE 9 SEM imaging of mat-related structures. (a) Magnified view of box area in Figure 7f. Upward clay laminae within tufted microstructures and wavy-crinkly layers. (b) Tufted microstructures and heavy minerals constitute bulges of the ‘elephant-skin’ texture. (c) Magnified view of box area in Figure 7h. Quartz grains, heavy minerals and randomly oriented clays constitute the dark-coloured mat layer. (d) Magnified view of box...
area in Figure 7k. Detrital particles wrapped by sheet clays. (e) Magnified view of box area in Figure 7e. Clay minerals above and throughout the ridge (arrow). No significant clue of liquefaction nor microbial shrinkage.

**FIGURE 10 Examples of fossil macroorganisms associated with microbial mats.** (a) Pyritized lobate form just beneath ‘fairy ring’ structures. (b) Disk with radially striated core (arrow) lies on domal buildups. (c-d) Disk or lobate form and flat pyritized microbial structures on the same strata are closely associated. (e) Circular disks (arrows) rest on wrinkle marks. (f) Disk and lobate form are close to wrinkle marks on the same level or not.

Captions for the supplementary information

**FIGURE S1 Additional flat pyritized microbial mats.**

**FIGURE S2 EDS elemental maps of bulges of reticulate patterns in cross-section perpendicular to bedding plane.** BSE and composite elemental maps. Note the wavy-crinkly laminae with a large amount of embedded heavy minerals.

the Gulf of Cambay, India. Modified after photo published in Banerjee et al. (2014). Lens cap diameter: 6 cm.
(h) Analogue discoidal microbial colony on bedding plane of sandstones from the Precambrian Vindhyan Supergroup, India. Modified after photo published in Banerjee et al. (2014).

**FIGURE S4 Photographs of mat-related structures found in literature.**

Figure 1

157x130mm (300 x 300 DPI)
Figure 2

180x197mm (300 x 300 DPI)
Figure 3

127x170mm (300 x 300 DPI)
Figure 4

151x211mm (300 x 300 DPI)
Figure 5

149x201mm (300 x 300 DPI)
Figure 6

142x185mm (300 x 300 DPI)
Figure 7

152x176mm (300 x 300 DPI)
Figure 8

121x81mm (300 x 300 DPI)
Figure 9

151x132mm (300 x 300 DPI)
Figure 10

152x151mm (300 x 300 DPI)
TABLE 1 δ\textsuperscript{13}C values of organic matter in mat-related structures (MRS).

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Supplementary information

FIGURE S1

FIGURE S1 Additional flat pyritized microbial mats.
FIGURE S2 EDS elemental maps of bulges of reticulate patterns in cross-section perpendicular to bedding plane. BSE and composite elemental maps. Note the wavy-crinkly laminae with a large amount of embedded heavy minerals.
FIGURE S4

Photographs of mat-related structures found in literature. (a) Modern example of submerged wrinkle marks from Redfish Bay, Texas. Modified after photo published in Hagadorn & Bottjer (1997). (b) Patches of wrinkle marks on bedding surface of fine-grained sandstones from the Early Cambrian, Chapel Island Formation, Canada. Modified after photo published in Buatois et al. (2014). (c) Parallel wavy wrinkle structures reproduced in wave tank.
### TABLE S1

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TABLE S1 Representative structural formulae of clays from microbial mats studied with EDS. EST, ‘elephant-skin’ texture; PTMM, putative tufted microbial mat; WS, wrinkle structure; LP, linear pattern.

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**TABLE S2** Organic elemental analyses (carbon and sulfur) on five microbial mats and their host sediments from the FB2 Formation. EST, ‘elephant-skin’ texture; DB, domal buildup; PWWS, parallel wavy wrinkle structure.