Mesophilic microorganisms build terrestrial mats analogous to Precambrian microbial jungles

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Development of Archean paleosols and patterns of Precambrian rock weathering suggest colonization of continents by subaerial microbial mats long before evolution of land plants in the Phanerozoic Eon. Modern analogues for such mats, however, have not been reported, and possible biogeochemical roles of these mats in the past remain largely conceptual. We show that photosynthetic, subaerial microbial mats from Indonesia grow on mafic bedrocks at ambient temperatures and form distinct layers with features similar to Precambrian mats and paleosols. Such subaerial mats could have supported a substantial aerobic biosphere, including nitrification and methanotrophy, and promoted methane emissions and oxidative weathering under ostensibly anoxic Precambrian atmospheres. High C-turnover rates and cell abundances would have made these mats prime locations for early microbial diversification. Growth of landmass in the late Archean to early Proterozoic Eons could have reorganized biogeochemical cycles between land and sea impacting atmospheric chemistry and climate.
Modern global biogeochemical cycles are distributed across the terrestrial and marine realms. Half of the global primary production, for example, is from algal and cyanobacterial growth in the sunlit surface ocean, while the other half comes from plants inhabiting the continents1. More generally, global nutrient cycles are characterized by extensive links between terrestrial and marine ecosystems, forming multiple feedbacks that regulate biological production and climate over geological time-scales2. Such biogeochemical feedbacks between land and ocean have operated at least since the emergence of terrestrial plants some 400 million years ago3–5, but may have begun much earlier during the Precambrian Eons, when microbial mats developed on land hundreds of millions or even billions of years before the evolution of plants6–8. Aquatic biofilms, hypersaline, and marine microbial mats are relatively abundant today, e.g. 9–13, and these provide insight into the ecology and biogeochemistry of prokaryote-dominated ecosystems. Subaerial examples, however, with the capacity to broadly colonize the early continents9–13, have remained elusive on the modern Earth. Indeed, large-scale colonization of land by microbial mats today is precluded by both animal predation and strong competition with plants for photosynthetic niche-space19,20. The lack of information on the ecology, physiology, and biogeochemical functioning of terrestrial microbial mats leaves models for the possible role of such ecosystems in the Precambrian Earth system almost entirely conceptual. Such information, in principle, could be obtained through the study of modern terrestrial mats with features analogous to those inferred from the remains of terrestrial ecosystems preserved in the rock record.

We discovered a subaerial microbial mat growing on exposed ultramafic rock surfaces at the Balambano Hydroelectric Dam on Sulawesi Island, Indonesia (Supplementary Fig. 1). The mat grows on steep outcrops tens of square metres in size (Fig. 1a), accretes to thicknesses up to 6 cm, and has many chemical and biological features that are comparable to the terrestrial microbial communities found in Precambrian rocks1–3. Geochemical and microbiological analyses reveal that the mat hosts a diverse photosynthetic microbial community with the metabolic capacity to drive large-scale biogeochemical cycling of C, N, and other elements released during rock weathering. Quantitative, model-based reconstructions of the possible role of similar mats during the Precambrian Eons imply that terrestrial ecosystems could have driven fluxes of matter and energy at global scales, promoted biological evolution, and helped regulate climate well before the emergence of land plants in the Phanerozoic Eon.

Results

Microbial community composition. The Balambano microbial mat supports a taxonomically diverse assemblage of microorganisms. It is compositionally layered (Figs. 1b and 2), and is largely comprised of filamentous bacteria and their exopolymer, as well as suspended mineral grains (Figs. 1c, d and 2). Cell abundances range from 10^9 to 10^9 cells cm^-3 and decrease with increasing depth in the mat (Supplementary Table 1). The mat is made up of a diverse assemblage of bacteria including an abundance of oxygenic and anoxygenic phototrophs (Fig. 3). Estimates of species richness are on the order of 3000–6000 taxa based on 97% identity between 165 rRNA genes (Supplementary Fig. 2 and Supplementary Table 2)—comparable to phototrophic communities in the sunlit surface ocean21 and hypersaline microbial mats16, but less than typical soil microbial communities, which can exceed tens of thousands of species22,23.

Beyond information on microbial community diversity, we also identify a number of taxa that can be linked to specific ecological roles. These taxa are reported at the genus level where possible, based on sequence resolution and database matching, and at the family level otherwise. In the upper reaches of the mat, the phototrophic community is dominated by non-heterocystous, filamentous Cyanobacteria (Leptolyngbya and Haloptylyngbya) and filamentous, photosynthetic Chlororolixi (Roseiflexus, Chloronema, and Candidatus Chloroploca) (Figs. 1d, 3 and Supplementary Table 3). Heterocystous filaments can also be observed (Fig. 1d) in the upper mat and known heterocystous cyanobacteria including Anabaena and Nostoc genera (Fig. 3c), which belong to family I cyanobacterial groups, comprise up to 5.5% of the photosynthetic community based on 16S rRNA gene sequences (Fig. 3; Supplementary Table 3). Cyanobacteria in the mat derive from some of the deepest branching cyanobacterial lineages including Gloeobacter, Thermosynechococcus and the filamentous Pseudanabaena and Leptolyngbya that likely radiated in the Mesozoic–Nearochean Era24,25. The base of the mat is enriched in firmicutes, including organisms related to anaerobic Clostridia genera and facultative aerobic Bacilli genera. Throughout the mat, Proteobacteria dominate (Figs. 3a and 4 and Supplementary Fig. 3), with each layer of the mat having taxonomically distinct populations. At the family level, Rhodobacteraceae, Aeromonadaeae, and Sphingomonadaeae are most abundant in the upper mat, Rhodobacteraceae, Acetobacteraceae, and Hyphomicrobiaeae dominate in the middle mat, and Pseudomonadaeae, Comamonadaeae, Moraxellaceae, and Burkholoderaceae are most abundant in the bottom layer (Figs. 3a and 4, Supplementary Fig. 3, and Supplementary Table 3). Lineages of the Alpha- and Gammaproteobacteria with metabolic potential for N2 fixation (genera Rhizobium, Azospirillum, and Azonumon) are also present (Supplementary Table 3). Nitrification, furthermore, is implied by the presence of bacteria from the family Nitrosomonadaeae (genus Nitrospira (Supplementary Table 3)), which are known from lab cultures as obligate nitrifiers26. Also identified were members of the phyla Verrucomicrobia (genus Candidatus Methylocylidophilum) and Alphaproteobacteria (family Methylobacteriaceae) (Supplementary Table 3), both known to grow through methylotrophy or methanotrophy27. The archaeal phyla Thaumarcheaetae, Batharchaeota, and Euryarchaeota were also present (Fig. 3b and Supplementary Table 3), suggesting metabolic potential for archaeal nitrification (genus Candidatus Nitrososphaera)28,29 and methanogenesis or C1 compound metabolism (genera Methanomethylovarans and phylum Batharchaeota)30,31, respectively. These subaerial mats thus support taxonomically and metabolically diverse communities with the capacity for biogeochemical cycling of carbon and nitrogen distributed across multiple microbial taxa.

Chemical composition. The Balambano mat microbial community retains bedrock components in the form of mineral grains and amorphous precipitates (Figs. 1c and 2, and Supplementary Fig. 4), which are trapped in the organic matrix of filamentous cells and extracellular polymeric substances (EPS). Such multicellular filaments of cyanobacteria and chloroflexi often stabilize microbial mats. When viewed in cross-section (Fig. 1b, c), the mat exhibits millimeter-scale laminations as well as finer-scale structural features, which include vertically aligned filaments and clusters of opaque inorganic material (Fig. 1c), presumably iron (oxyhydr)oxides. The inorganic chemical composition of the mat (Supplementary Table 4) is mostly Fe, Si, and Mg, with a mineralogy dominated by X-ray amorphous phases (Supplementary Fig. 4), but also sheet silicates such as nontronite [Ca_{10}Fe_{2}(Si,Al)_{4}O_{10}(OH)_{2}]·2H_2O and pimelite [Ni_{2}Si_{4}O_{10}(OH)_{2}]·5H_2O. The presence of clay minerals in the mat confirms its capacity to trap and retain material to form protosol. Organic carbon concentrations are highest (12 wt.% in
the upper 2 mm of the mat, where the isotopic composition of organic C ($\delta^{13}C$) is $-33\%$ (Supplementary Fig. 5), indicative of photosynthetic C fixation. Deeper in the mat, organic carbon concentrations decline to 1 wt.% with isotopic compositions as light as $-41\%$ (Supplementary Fig. 5), indicating incorporation of $^{13}$C-depleted carbon, likely through methanotrophy. With $10^8$–$10^9$ cells $cm^{-3}$, cellular biomass makes up from 0.001 to 20% of the total organic C in the mat (see Methods below). Organic N reaches nearly 1 wt.% in the surface of the mat with a C:N ratio of 15—higher than the Redfield ratio of around 6 that typifies marine phytoplankton today and in the past (Supplementary Fig. 5). The N isotopic composition of organic matter (OM) ranges from $-3.4$ to $-0.6\%$ (Supplementary Fig. 5), indicative of biological fixation of atmospheric $N_2$ (ref. 35).

**Biological production and growth rate.** Physiological measurements and mass balance considerations constrain the metabolic potential and overall biogeochemical activity of the Balambano mat. Measurements of photosynthesis revealed net $O_2$ production of 23 mmol m$^{-2}$ d$^{-1}$ (Fig. 5 and Table 1), while mass balance calculations of mat growth suggest in situ organic carbon accretion rates of 20 mmol m$^{-2}$ d$^{-1}$ (Supplementary Fig. 6 and Table 1). Measured CH$_4$ production rates were up to 4.1 mmol m$^{-2}$ d$^{-1}$ (Supplementary Fig. 6 and Table 1). Together, these rates indicate that both retention of OM and CH$_4$ loss are important sinks of reduced carbon in the mat that balance net $O_2$ production (Supplementary Fig. 6 and Table 1).

**Discussion**

Many of the Balambano mat features are similar to the geochemical, mineralogical, fossil, and structural evidence for the presence of microbial life on the continents and early paleosols during the Archean Eon. The filamentous morphology of the Balambano mat may be essential for the stabilization of subaerial mats and is amenable to preservation as readily identifiable microfossils, such as those found in Precambrian terrestrial rocks. In fact, a number of microbial lineages that have been interpreted from Archean paleosols, such as Actinobacteria, purple sulphur bacteria, and methanogenic Archaea, are found in the Balambano mat (Fig. 3). At a larger scale, mineral trapping (Fig. 1c), and growth through the accretion of compositionally distinct layers (Figs. 1b and 2), can leave characteristic laminated features in the rock record akin to structures recognized as stromatolites in Archean terrestrial and marine rocks. Taken together, the structural support provided by filaments to the Balambano mat and the putative filamentous nature of early terrestrial stromatolite communities suggest that the evolution of multicellularity in the form of filamentous bacteria may have been requisite for the proliferation of thick, subaerial microbial mats and their preservation in the rock record. Photosynthetic and methanogenic organisms in the mat impart characteristic.
Fig. 2 Stylized illustration of Balambano microbial mat. The illustration depicts the positions of major microbial taxa, weathered rock fragments, and underlying bedrock. Cells not drawn to scale.

Fig. 3 Composition of the Balambano mat microbial community. Distribution of Illumina 16S rRNA gene sequences are shown per bacterial phylum (a), per archaeal phylum (b) and per cyanobacterial genera (c), in three sections of the microbial mat (top = 0–2 mm; middle = 2–5 mm; bottom = 5–20 mm). The number of sequence reads per phylum is calculated as a percentage of the total sequence reads for each sample. *Other represents reads that contributed <1 and 0.05% of the total number of reads for A and C, respectively.
geochemical signals such as $^{13}$C-depleted OM, as well as elemental stoichiometries, that can ultimately be recorded as fossil OM in paleosols and fossilized microbial mats. Notably, mat retention of inorganic material (Fig. 1b, c) provides a weathering environment conducive to clay mineral formation and soil accretion—processes critical to early soil development, mineral evolution, and the preservation of weathering products as paleosols. By extension, microbial mats would also have influenced the delivery of weathering products to oceans, and the burial of OM in marine sediments. For example, terrestrial microbial activity has been invoked to explain increased nutrient, sulphate, and trace element fluxes to the oceans in the Archean and Neoproterozoic Eras. Nevertheless, most metabolic potential hosted within these taxa emerged early in Earth’s history and is likely retained at the community level such that the capacity to drive biogeochemical cycling is conserved through time despite taxonomic reshuffling of metabolic potential through horizontal gene transfer. We thus argue that the metabolic potential and biogeochemical functioning of Balambano mat makes a good analogue for similar mats in the Precambrian Eons, but recognize that the taxonomic composition of the modern mat is unlikely to reflect the taxonomic composition of Precambrian mats. Relationships between taxonomy and function remain controversial, but when better known, may prove important considerations for establishing and interpreting analogue systems.

Rates of photosynthetic productivity and OM accretion in Balambano mat are similar, on an area-specific basis, to moderately productive coastal marine waters or perennial grasslands today, implying that if subaerial mats were widespread on the Precambrian continents, then they would have played a key role in global carbon and oxygen budgets (Fig. 6). Upper estimates for gross marine productivity in the Precambrian Eons are between 170 and 500 Tmol yr$^{-1}$ (ref. 49). Respiration in Balambano mat consumes 66% of the O$_2$ produced giving a gross production of 38 mmol m$^{-2}$ d$^{-1}$ (Table 1) or 14 Mmol C km$^{-2}$ yr$^{-1}$, similar to values estimated for Archean benthic freshwater mats and at the lower end of rates (range 6–1200 mmol m$^{-2}$ d$^{-1}$, average 240 mmol m$^{-2}$ d$^{-1}$, Supplementary Table 5) measured in hypersaline, marine, and aquatic environments (Supplementary Table 5). Applying our measured rates of gross productivity to an area equivalent to, for example, 5% of the late Archean land area (2.5 $\times$ 10$^7$ km$^2$ (ref. 50), lower end of values reported for the Neoarchean) yields a terrestrial productivity of 17 Tmol yr$^{-1}$ (Fig. 6), or up to 10% of total estimated global production with minimal coverage of land surfaces. Benthic lacustrine mats could contribute to some of this production, but modern lakes and rivers cover only ~2% of the land surface and just a small fraction of that is suitable habitat for mat growth (Fig. 6). The fraction of the available land mass colonized by mats would depend on precipitation patterns and water availability. Modern climate and topography allow development of forests and grasslands on ~55% of today’s land surface, which when translated to the Archean would have allowed terrestrial mats to contribute up to ~190 Tmol yr$^{-1}$ to global productivity. Landmasses in the tropics today are typically overgrown with dense forest and ‘tangled’ vegetation referred to as jungles. We suggest, by analogy, that tropical regions in the Archean Eon were similarly overgrown with dense and ‘tangled’ microbial communities, such as
the Balambano mat, that thus formed Precambrian continental jungles.

Combining cell counts and C-fixation rates allows us to evaluate the possible importance of subaerial microbial mats in driving evolutionary processes during the Archean Eon. At 10% surface coverage, microbial mats likely harboured tenfold fewer microbial cells, globally, than the Archean surface ocean (Supplementary Table 6 (ref. 55)). Based on our measurements in Balambano mat, global rates of cell division, on the other hand, could have been similar in the two ecosystems (Supplementary Fig. 7 and Supplementary Table 6), and this implies similar annual probability for genetic mutations on land and in the ocean. In microbial mats, cells also live in much closer proximity than in the open ocean, which fosters horizontal gene transfer, and thus rates of genetic exchange are expected to be higher in mat communities than in the ocean. Together, similar mutation rates and higher frequencies of horizontal gene transfer in land-based ecosystems would thus have made them prime locations for

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**Table 1 Gross and net oxygen production**

| Single sources and sinks | Gross O₂ production (mmol m⁻² d⁻²) | Method |
|--------------------------|------------------------------------|--------|
| 1D model                 | 23                                 | 1      |
| O₂ flux atmosphere + rock| 8.4                                | 2,3    |
| CH₄ flux atmosphere + OM production | 24            | 4,5    |

**Gross O₂ production** (mmol m⁻² d⁻²) Method

1. 1D transport reaction based on daytime greenhouse profile; 2, flux determination from daytime greenhouse profile; 3, flux determination of in situ profile; 4, in situ mat accretion; 5, headspace analysis in dark mat incubation; 6, 1D transport reaction modelling based on night time greenhouse profiles, see also Supplementary Fig. 4

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**Fig. 5** Profiles of oxygen concentrations as a function of depth in the mat. **a** Diel oxygen profiles measured in the greenhouse every 2 h. **b** Net oxygen production modelled from the 15:12 profile from **a** using a 1D transport-reaction model. **c** Net oxygen production modelled from the 3:40 profile from Fig. 3a using a 1D transport-reaction model. Respiration is restricted to the oxic surface layers. High oxygen concentrations during the day likely stimulate respiration throughout the mat. The areal rates calculated from the rate profiles are presented in the box model (Supplementary Fig. 5) and used for the calculations in Table 1 and Fig. 6.
microbial evolution in the Archean Eon, implicating continental mats as possible drivers of the Archean expansion in biological diversity.

Continental microbial communities have also been implicated in oxidative weathering, global oxygen budgets, and the evolution of atmospheric O2 and CH4 in the Precambrian Eons. To directly test the potential role of continental mat communities in global O2 cycling, we partitioned net mat O2 sinks between loss to the atmosphere and consumption through oxidative weathering of the underlying bedrock. Under the modern oxygenated atmosphere, 6.3 mmol m−2 d−1 or 16% of gross O2 production escapes the mat surface to the atmosphere, 6% is consumed through weathering of the underlying bedrock, and the balance channeled through respiration. Under low-O2 early-to-mid-Archean atmospheres, 24 mmol m−2 d−1 or up to 70% of the O2 production would escape from the mat surface. With minimal continental freeboard, such an O2 flux from continental mats would be entirely consumed through reactions with atmospheric O2 sinks even at 100% land coverage (Fig. 6). Growth of landmass in the late Archean to early Palaeoproterozoic to about 10% of the Earth surface, however, could have allowed O2 effluxes from continental mats to overcome these sinks and O2 to accumulate in the atmosphere, if residual organic carbon was ultimately buried in marine sediments, even with minimal contribution from marine or benthic freshwater photosynthesis. Rates of organic carbon reoxidation would decrease upon export from the mat into surrounding anoxic environments, promoting transport to the oceans, burial in sediments, and net oxygenation of the atmosphere. At the same time, CH4 effluxes from terrestrial microbial mats to the atmosphere could have contributed to atmospheric haze formation in the Archean Eon and climate warming throughout the Archean and Proterozoic Eons. The combined effects of mat O2, organic C, and CH4 efflux should be investigated further, but overall the growth of continental landmass would have redistributed biogeochemical cycles between the oceans and continents with strong potential to alter the atmospheric chemistry between the late Archean and early Proterozoic Eras.

Increased fluxes of nutrients and redox-sensitive trace metals to the oceans 100’s of millions of years before the Great Oxidation Event (GOE) are used to infer high atmospheric oxygen levels in the late Archean Eon. Indeed, there is abundant evidence for the emergences of oxygenic photosynthesis by at least the Mesoarchean Era and this comes from both geochemical proxies and molecular phylogenies. Rapid recycling of oxygen produced in terrestrial microbial mats through respiration or coupled to rock weathering and sulphide mineral oxidation, however, would have allowed for delivery of such redox tracers to the oceans even under an anoxic atmosphere. It is thought that aerobic respiration may have evolved before oxygenic photosynthesis could have emerged as the immediate response. A 0.02% coverage of the late Archean landmass by continental photosynthetic microbial mats would have been sufficient to produce the sulphur flux proposed for the Mesoarchean, and at a 1% coverage, a modern sulphur flux can be sustained (Fig. 6). The presence and activity of nitrogen fixing and nitrifying bacteria in the mat also implies that terrestrial mats could have supplied the coastal ocean with oxidized N-species under an anoxic atmosphere, as previously suggested for biological soil crusts. Nitrification likely emerged in response to oxygenic photosynthesis, even though widespread evidence for nitrification at global scales seems only to emerge following the GOE. Based on our measured net production and C:N ratios, the mats could have supplied 1 Tmol yr−1 nitrate to the coastal ocean and such a flux may be recorded in late Archean sedimentary N-isotope signals. Enhancement of fluxes of nitrate and sulphate could also have stimulated coastal marine primary production, fuelling sulphate reduction and supporting the development of late Archean sulphidic oceans.

Our work on Balambano mat qualitatively and quantitatively shows how oxygenic photosynthesis supports taxonomically and metabolically diverse microbial jungles that can drive fluxes of
matter and energy at global scales. Export of nutrients from terrestrial mats to the coastal ocean can be sustained by aerobic metabolisms and weathering reactions that operate within the mat and largely independent of atmospheric oxygen. Terrestrial mat CH4 effluxes can contribute to atmospheric chemistry and climate, possibly playing an outsized role compared to marine communities in the Proterozoic Eon. In this way, terrestrial microbial jungles would have established the first links between biogeochemical cycles on land and in the ocean during the Archean Eon and these links could have continued throughout the Proterozoic Eon until plants colonized the continents and usurped the microbial continental biosphere in the mid-Ordovician some 470 Ma.

Methods

Sampling site. Samples were collected from an outcrop of ultramafic bedrock exposed during the construction of the Bambelano hydroelectric dam (Supplementary Fig. 1). The dam was constructed in 1997 and the samples collected in 2006 and again in 2009, 9–12 years after bedrock exposure. The outcrop measures several tens of square-meters and is periodically wet by direct precipitation as well as overland flow (Fig. 1a). The outcrop is mostly covered by the microbial mat, but bedrock is exposed in places, and is also partly covered by grasses. The maximum observed thickness of the mat was 6 cm, but typical observations were ~3 cm (Fig. 1b).

Mat description. The microbial mat forms distinct layers that vary on a millimeter-centimeter scale (Fig. 1b). The mat is mechanically robust allowing removal of intact square-decimeter sections. The dominant colour of the mat is ochre, which is characteristic of iron oxohydroxides. A green layer is visible at about 2–5 mm depth, coinciding with a peak in oxygen concentrations (Fig. 5 and Supplementary Fig. 10). Light microscopy revealed opaque minerals and rock fragments trapped in a translucent matrix that also hosted microbial filaments (Fig. 1c). Fluorescence microscopy of individual filaments revealed the presence of heterocystous cyanobacteria (Fig. 1d), enabling the mat to fix nitrogen from the atmosphere.

Oxygen measurements. In situ profiles of oxygen concentrations were determined using a gold amalgam voltammetric microelectrode according to Brendel and Luther (Supplementary Fig. 10). The profile was measured in 2006 directly on the outcrop hosting the mat. These measurements are thus distinct from the measurements used to construct the profiles shown in Fig. 5, which were conducted in a greenhouse with a Clark type microelectrode. We used 25-μm-diameter Clark-type oxygen sensors (Unisense, DK) and m-profile software (MPI-Bremen, D) for diel oxygen concentration profile measurements. Subsamples of the mats were transferred in a greenhouse in 2009 to conduct measurements of photosynthesis and respiration. These samples were kept in flumes in a greenhouse and oxygen profiles measured under subaerial conditions and within 1 month of collection. Profiles were measured every ~3 h over a 24-h cycle (Fig. 5a). Profiles measured at 3 a.m. and 3 p.m. were used to calculate respiration and net oxygenic photosynthesis, respectively, using a 1D reaction-transport model (Fig. 5b, c).

Flux determinations from profiles. Oxygen fluxes to both the atmosphere and the underlying rock were calculated based on oxygen concentration gradients using Fick’s law of diffusion:

$$J = -\frac{\partial C}{\partial x}$$

(1)

where $J$ is the diffusive flux (mol m$^{-1}$ s$^{-1}$), $\partial C/\partial x$ the concentration gradient (mol m$^{-1}$), and $D$ is the diffusion coefficient (m$^2$ s$^{-1}$). Therefore, the diffusion coefficient can be used for all calculations. The flux to the rock was determined from an in situ profile measured with a voltammetric electrode (Supplementary Fig. 10), and the atmospheric flux from a profile measured by a Clark-type electrode in the greenhouse at a higher resolution (Fig. 5).

Oxygen fluxes from the mat to the atmosphere would be different under an anoxic atmosphere, like in the Archean Eon, than in the modern-day atmosphere with 20% oxygen. To estimate fluxes from a mat like Bambelano to an anoxic atmosphere, we modeled oxygen production rates in the mat based on oxygen profiles (Supplementary Figs. 8 and 9). Oxygen concentrations were modelled as a function of rates of photosynthesis that are themselves influenced by light availability and rates of respiration, which in turn were affected by oxygen availability and described by Michaelis–Menten-type equations.

Oxygen production ($O_{2\text{Prod}}$) is modelled according to:

$$O_{2\text{Prod}} = \frac{\text{PAR} \times V_{\text{max Pro}}}{(K_{\text{m Pro}} + \text{PAR})}.$$  

(2)

Photosynthetically active radiation (PAR) was set to 1500 μmol m$^{-2}$ s$^{-1}$ at the surface decreasing by 1% every depth interval of 0.071 mm. $V_{\text{max Pro}}$ was set to 2.1 nM s$^{-1}$; $K_{\text{m Pro}}$ is the ‘affinity’ for light, set to 1000 μmol m$^{-2}$ s$^{-1}$.

Respiration ($O_{2\text{Resp}}$) was modelled according to:

$$O_{2\text{Resp}} = \frac{O_{2} \times \text{Resp} \times \text{Min Pro}}{V_{\text{max Resp}} + O_{2} + \text{Resp}}.$$  

(3)

$O_{2\text{Resp}}$, is the concentration at time $n - 1$ (previous time step) and depth $x$, $V_{\text{max Resp}}$ is set to 1.75 nM s$^{-1}$, and $K_{\text{m Resp}}$ was set to 230 nM of $O_{2}.$

The model was parameterized to reproduce the measured $O_{2}$ profiles (Fig. 5b, c) and steady-state was approached within the first 1200 iterations of 1 s each (Supplementary Fig. 9). Next the oxygen concentration in the water was changed to 0 μM and the model run again with 1200 iterations of 1 s each (Supplementary Figs. 8 and 9) to reach a new steady-state.

The sensitivity of the model was tested by changing the variable parameters by ±10% (change of 0 and 5% for the decrease of PAR with depth) and evaluating the influence of these changes on the $O_{2}$ slope across the mat–water interface at the end of the model run. This 10% change resulted in a change in the slope of ±3% for the high $O_{2}$ scenario and less than ±1% for the no $O_{2}$ scenario (Supplementary Table 7). Changes of the variable parameters had a much stronger effect on the oxygen concentrations deeper in the mat than they did on the slope at the mat–water interface.

Methane flux determination. A subsample of the mat was incubated in the dark for 1.5 months and the methane production was measured in the headspace with a gas chromatograph (GC) equipped with an flame ionization detector (FID) (SRI, USA). An areal flux was determined assuming a 6-cm-thick mat. Half the flux determined by the incubation was used for the flux determinations to account for inhibition of methanogenesis under the oxic conditions that develop during the incubation. As they have a higher spatial resolution, OM production was calculated from the model balance of organic carbon (C$_{org}$) content, and isotopic composition, as well as the N isotopic composition of the mat was measured through combustion-based elemental analysis of freeze-dried sub-samples on an elemental analyzer coupled to an isotope ratio mass spectrometer.
Inorganic solid-phase composition. The mineralogy and solid-phase chemistry of the microbial mat was determined on freeze-dried sections as described in ref. 74. In brief, the mineralogy was analyzed using X-ray diffraction (XRD; Rigaku MiniFlex) according to Morono et al.83. The mat was aseptically sectioned into three layers: top, middle, and bottom, and the X-ray diffraction pattern and the X-ray fluorescence (XRF) on a lithium tetraborate fused bead using a Philips PW2440 4 kV instrument (Pana-lytical Inc., Natick, MA, USA) (Supplementary Table 3). Accuracy for major and trace elements was within 1% and 5%, respectively, based on the analyses of standard reference materials, and relative precision was within 0.5% determined by repeated analyses of the same fused bead.

Molecular microbiology. Samples for molecular microbial analyses were subsampled in the field and frozen at −20 °C, shipped to Denmark on dry ice, and stored frozen until extraction. The microbial mat was aseptically sectioned into two layers: top (0–2 mm), middle (2–10 mm), and bottom (10–25 mm) and weighed. Microbial community DNA was extracted from 0.25 mg of each microbial mat using the Power soil® DNA Isolation Kit (MoBio Laboratories, Inc., Solana Beach, CA) according to the manufacturer’s protocol. The purity and quantity of DNA extracted were assessed and determined using both a NanoDrop 1000 spectrophotometer (Thermo Scientific) and the Picogreen® fluorosceine assay.

16S rRNA gene amplicon sequencing and bioinformatic analysis. 16S rRNA gene amplicon sequences were generated by the Joint Genome Institute (Walnut Creek, CA) through a community sequencing program project (CSP-4140). Libraries for Illumina MiSeq sequencing (a 2 × 300 bp read) was produced by amplifying region V4-V5 of the 16S rRNA gene using primers 515F (5′-GTGCCAGCMGCGCGTAA-3′) and 806R (5′-GGACTACHVGGGTWTCTAAT-3′) corresponding to the V4-V5 region of the 16S rRNA gene previously described75,76. Sequences were processed using Mothur77 (https://www.mothur.org/wiki/MiSeq SOP). Briefly, sequences were removed from the analysis if they contained ambiguous characters, had homopolymers longer than 8 bp, or did not align to a reference alignment of the correct sequencing region. Unique sequences were identified and their frequency in each sample determined and then a pre-clustering algorithm78 was used to further de-noise sequences within each sample. Unique sequences were aligned against a SILVA reference alignment (available at http://www.mothur.org/wiki/Silva_reference_alignment). Sequences were chimera checked using UCHIME49 and were then clustered into OTUs with 97% sequence identity using OptiClust81. OTUs were classified using a SILVA reference taxonomy database (release 132, available at http://www.mothur.org/wiki/Silva_reference_files (Supplementary Fig. 8, Supplementary Table 4). Diversity metrics were calculated in Mothur.

qPCR and DNA content. Total 16S rRNA gene copy numbers were determined by qPCR using a bacterial-specific (27F, 5′-AGAGTTTGTATCCTGCGCTCA-3′) forward primer coupled to a universal reverse primer (DW519R, 5′-GNT TTACCCGCGCTGCTG-3′) according to Zaikova et al.82 (Supplementary Table 5). Standards for total bacteria quantification were derived from 16S rRNA gene clone libraries using a 16S rRNA from the bacteria SUP05 according to Zaikova et al.82. A tenfold dilution series for each standard ranging from 3 × 105 to 3 × 101 copies μl−1 for bacteria was used in real-time analysis. For the purposes of this study, the limit of detection was set at the Ct values of the no template controls and was determined to be ~5 × 103 copies μl−1 for all sample replicates. The quantity of DNA extracted was also used to estimate cell abundance assuming 4 fg of DNA per cell.

Cell counts. Cell abundances were determined using epifluorescence microscopy. The mat was aseptically sectioned into three layers: top, middle, and bottom, weighed and then preserved with 2.5% (final concentration) of glutaraldehyde. The fixed samples were washed twice with phosphate-buffered saline and then dissociated form the mat matrix according to Morono et al.83. In brief, mat sections were immersed in a solution of 2.5% NaCl, detergent mix (100 mM EDTA, 100 mM sodium orthophosphate, 1% (v/v) Tween 80), and methanol, then vigorously shaken for 120 min at 500 rpm. After shaking, the mat sections were sonicated and milled with 1 mm glass beads, and methanol was added to a final concentration of 10% (v/v) and 15% (v/v) to bring the volume to 30 ml. After centrifugation at 4500 × g for 30 min after which the supernatant, including the high-density layer(s), was carefully removed and transferred to a separate vial. Cells were enumerated following the V4-V5 region of the 16S rRNA gene amplicon sequencing and bioinformatic analysis procedure. Total bacterial 16S rRNA gene copy numbers were determined by qPCR using a bacterial-specific primer (F 5′-GNT TCAG-3′ forward primer coupled to a universal reverse primer (DW519R, 5′-GNT TTACCCGCGCTGCTG-3′) according to Zaikova et al.82 (Supplementary Table 5). The mat consisted of a large number of small bacterial cells and a relatively small number of larger, filamentous microbes. The DAPI counts only represent the small bacterial cells, as the larger filamentous cells were not separated from the matrix. The general agreement between DAPI counts and q-PCR, as well as quantification of DNA yields, gives us confidence that visual enumeration captured the vast majority of the cells. Our cell counts lie at the lower end of previously reported values for mats85, and this is in line with the minimum values we report.

Calculation of cellular carbon and cell divisions in subaerial mats. To calculate cellular biomass, we considered cell abundances of 109–1010 cells cm−2 and cellular carbon contents of 1–150 g C cm−2 (ref. 86–88). This translates to between 10−7 and 1.5 × 10−6 g C cm−3 of wet mat. Based on a porosity of 0.97 and a particle bulk density of 2.65 g cm−3, the total organic carbon content of 1–12% dry mass translates to carbon concentrations in the wet mat of between 0.7 × 10−4 and 9.5 × 10−3 g C cm−3. This means that cellular biomass accounts for between 0.001 and 20% of the total organic carbon. Cell counts (Supplementary Table 6) and gross photosynthesis rates (Fig. 5) were used to calculate a cell turnover time in the mats based on 1–150 fg of C per cell85,86. Cell counts of the top section were used, as the locus of photosynthesis is within this depth interval (Fig. 5b). This resulted in a turnover time of 0.2 d or 5 divisions per cell per day. Based on the range in carbon concentrations per cell, this results in 6.9 × 1027 to 1.0 × 1025 divisions per day for 0.1% continental coverage by microbial mats 6.9 × 1027 to 1.0 × 1025 divisions per day for 100% continental coverage (Supplementary Fig. 10, Supplementary Table 7). An area ocean surface area was calculated based on a land area of 5% leaving an ocean area of 95% of the earth surface or 485 × 106 km2. Using cell densities and turnover times for the surface ocean according to Whitan et al.55 and a tenfold lower primary productivity than the modern results in 4.8 × 1027 cells and 2–8 × 1026 divisions per day (Supplementary Fig. 10, Supplementary Table 7).

Eukaryotes. Microeukaryotes, including diatoms and algae, were visible through microscopy, but were not directly analyzed through, for example, amplicon sequencing of the 18S rRNA gene or ITS regions. 16S rRNA gene sequences recovered from mitochondria and chloroplasts, however, were rare in our amplicon sequence data and this is at least qualitatively consistent with our limited observations of Eukaryotes through microscopy. For example, chloroplast reads represented <0.2% of all reads and <2% of all photosynthetic reads (Supplementary Table 8). Photosynthesis and respiration by microeukaryotes would have been integrated into our rate measurements, but their likely very low abundance, compared to bacteria, implies that they play a limited role. We do recognize, however, that abundance is not a good proxy for activity, but given the extreme dominance of bacteria here, we argue that they also play the most important role.

Data availability. Sequence data that support the findings of this study have been deposited into the SRA database under SRA accession: SRP155785. Additional data that support the findings of this study are available from the corresponding author upon request.

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Author contributions
S.A.C., N.F. and D.A.F. conceived the study; S.A.C., A.H.O., D.A.F., C.H. and S.N. performed the field work; N.F., S.A.C., R.L.S. and A.H.O. conducted the analyses; N.F., R.L.S. and S.A.C. wrote the manuscript with input from all co-authors including L.C.M., D.E.C., K.K. and S.V.L.

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