

Inhibition of carbonate synthesis in acidic oceans on early Mars

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Several lines of evidence have recently reinforced the hypothesis that an ocean existed on early Mars^{1–7}. Carbonates are accordingly expected to have formed from oceanic sedimentation of carbon dioxide from the ancient martian atmosphere^{7,8}. But spectral imaging of the martian surface has revealed the presence of only a small amount of carbonate, widely distributed in the martian dust⁹. Here we examine the feasibility of carbonate synthesis in ancient martian oceans using aqueous equilibrium calculations. We show that partial pressures of atmospheric carbon dioxide in the range 0.8–4 bar, in the presence of up to 13.5 mM sulphate and 0.8 mM iron in sea water⁸, result in an acidic oceanic environment with a pH of less than 6.2. This precludes the formation of siderite, usually expected to be the first major carbonate mineral to precipitate⁸. We conclude that extensive interaction between an atmosphere dominated by carbon dioxide and a lasting sulphate- and iron-enriched acidic ocean on early Mars is a plausible explanation for the observed absence of carbonates.

A survey of the literature suggests four alternative classical explanations for the detection of only small concentrations of carbonate minerals: (1) no primary carbonate formation in a cold and/or dry environment; (2) the inability of the Thermal Emission Spectrometer (TES; on board the Mars Global Surveyor spacecraft) and the Thermal Emission Imaging System (THEMIS; on board the Mars Odyssey spacecraft) to detect carbonates; (3) secondary chemical alteration of ancient carbonate sediments; or (4) carbonates that are obscured by younger rock materials. The first explanation, which is in disagreement with the geomorphological evidence, supports the interpretation of early martian conditions as having been similar to those prevailing at present¹⁰. The second highlights the possibility that substantial regional carbonate deposits can be 100% exposed and remain undetected at the sensitivity of TES/THEMIS¹¹. The third includes the possibility of water vapour and sulphates combining to form acid rain⁴ and thereby promoting chemical decomposition of superficial carbonate layers, acid-fog weathering¹², and/or photodecomposition¹³. And the fourth includes large carbonate deposits hidden beneath several-centimetres-thick secondary alteration rinds that envelop rock materials on the surface¹⁴, or carbonates masked by the formation of relatively recent volumetrically abundant soils¹⁴; it also includes other resurfacing processes that may have mantled carbonate deposits, such as eolian deposition during the dominant phases of Mars' mostly frozen state¹⁵, sedimentation in a later Hesperian-age ocean that occupied the northern plains⁵, emplacement of basaltic lavas and pyroclastic materials in and around the Tharsis and Elysium magmatic complexes³, and the deposition of 100-m-thick Vastitas Borealis Formation materials, a potentially ice-rich sublimation residue of outflow events¹⁶.

Using aqueous equilibrium calculations, we analyse here the feasibility of carbonate synthesis in ancient martian oceans, taking into account the geochemical cycles that were probably in operation then. To this end, our analysis is based on two key assumptions: (1)

that well over 90% of the carbonate rock formation on Earth occurs in open bodies of liquid water¹⁷; and (2) that the fate of the martian oceans was rapid freezing solid followed by sublimation and cold-trapping of ice at higher latitudes^{1,10}. Thus, the problem of carbonate formation on early Mars basically concerns the chemical conditions operating in open bodies of liquid water. Local sedimentation driven by sequential evaporation in closed basins⁸ is considered here an interesting, but minor, mode of water evolution on early Mars; and where such processes are likely to have occurred, the subsequent increase of iron and sulphate in the solution would have further lowered the pH, thus maintaining ionic undersaturation with respect to carbonates. On the other hand, more recent transitory reduced lakes would not have been able to control the process of carbonate synthesis either, as Mars has been mainly cold and dry throughout the Hesperian and the Amazonian, except during brief and intriguing episodes^{5,15}.

To raise the mean surface temperature of Mars above the freezing point, a greater atmospheric CO₂ pressure is required. The present CO₂ pressure (6 mbar), in addition to that adsorbed in the regolith (30–40 mbar; ref. 18) and the minimum quantity sequestered as ice or clathrate in the poles, represent nearly 40 mbar of CO₂. If the elimination processes by impact erosion and pick-up ion sputtering removed 95–99% of the Noachian atmosphere¹⁸, then the martian surface CO₂ pressures 4 Gyr ago could have been of the order of 800–4,000 mbar. However, without the additional warming provided by the presence of other strong greenhouse gasses, such as CH₄ and NH₃ (ref. 19), even 4 bar of CO₂ would have been insufficient to yield mean global temperatures above the freezing point. Carbon dioxide ice clouds in the troposphere may have also contributed to warming early Mars, as ice particles reflect the outgoing thermal infrared radiation back to the surface²⁰.

Nevertheless, this approach for high CO₂ pressures in early Mars has been classically disputed by two lines of arguments: the loss of atmospheric volatiles¹⁸, and the lack of carbonates on the martian surface¹⁰. But substantial amounts of atmospheric CO₂ may have been maintained during the Noachian by volcanic outgassing from Tharsis^{2,3,5}. In addition, if the oceans were acidic, the lack of carbonates no longer qualifies as a reliable indicator of CO₂ partial pressures in the Noachian martian atmosphere. Consequently, the possibility of substantial atmospheric CO₂ pressures in early Mars cannot be ruled out. Later, nearly all of the martian atmosphere would have been lost by impact erosion during the heavy bombardment period^{18,21}, and by sputtering acting throughout most of the martian history¹⁸.

Such a carbon dioxide-dominated atmosphere in early Mars would have rendered an ocean mildly acidic by releasing free protons in the sequence $\text{H}_2\text{O} + \text{CO}_2(\text{g}) \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{H}^+ + \text{HCO}_3^-$. The addition of iron contributed even more to oceanic acidification. Also, sufficient iron in solution acted as an excellent buffer, as appreciable amounts of iron can be dissolved in moderately acidic to neutral solutions. Hydrolysis of ferrous iron generates free protons by the equilibrium $\text{Fe}^{2+} + 2\text{H}_2\text{O} \leftrightarrow \text{Fe}(\text{OH})_2 + 2\text{H}^+$, and the process is much more effective if the iron is in the form of ferric iron: $\text{Fe}^{3+} + 3\text{H}_2\text{O} \leftrightarrow \text{Fe}(\text{OH})_3 + 3\text{H}^+$.

The potential source of iron in terrestrial Archaean oceans was iron emitted from hydrothermal vent fluids produced by intense submarine magmatic activity creating hot Fe²⁺-enriched plumes²²; and the subsequent photolytic oxidation of ferrous iron to Fe³⁺ would provide the ocean with its major oxidized species²³. Consequently, iron is thought to have been abundant in the oceans of early Earth, reaching concentrations as high as 50 μM (ref. 24). On early Mars, the iron-rich nature of the mantle and active volcanism³ produced igneous minerals containing ferrous iron. In fact, the martian surface is much richer in iron than the surface of Earth: the reddish colour of the martian surface is due to the optical properties of ferric-iron-bearing minerals present in the oxidized surface layer²⁵. In this sense, up to 800 μM of Fe²⁺ for basin feedwater⁸

on the surface of early Mars has been considered a reasonable estimate for an iron-rich fluid.

Moreover, jarosite minerals, as well as evaporite deposits containing magnesium sulphate salts, a typical sublimation residue, have been detected by the Mars Exploration Rover (MER) Opportunity in the sediments deposited at Meridiani Planum. The amount of salts detected, up to 40% in the outcrop⁷, leads us to expect that sulphate levels in ancient martian oceans were at least at the same order of magnitude as those derived from weathered ultramafic rocks on Earth (0.18 mM; ref. 8), since the salinity of present-day terrestrial oceanic water is 3.5×10^5 p.p.m. (ref. 26). Following the Opportunity results, however, we assume a Fe^{3+} -enriched solution in equilibrium with jarosite²⁷ at least locally and/or temporarily, so potentially displaying even higher sulphate levels, up to 13.5 mM (see Table 1).

For the Noachian oceans, aqueous thermodynamic calculations considering a solution enriched in iron hydroxides and sulphate result in a pH between 5.3 and 6.2 for the siderite- Fe^{2+} equilibrium; and when the ferrous iron was photolytically oxidized to ferric, the final pH was between 1.9 and 2.1 for the siderite- Fe^{3+} equilibrium (Fig. 1; see Methods). The subsequent evolution of the acidic environment over time results in Fe^{3+} -sulphatic species dominating the chemistry of the oceans (Fig. 2). Thus, the TES analyses, which indicate an absence of carbonate minerals at the surface of Mars, can be comprehensively explained by the acidity of the ancient martian oceans. These results address the recently posed contradiction between TES-detected martian mineralogy, and planetary geomorphology and MER-unveiled geochemistry, the former reporting small amounts of disseminated carbonates and the latter evidencing the geomorphologic modification and chemical alteration of surface materials through significant amounts of running water during long periods of early Mars history. Also, this new scenario is qualitatively different from all those argued before (and listed above), as they account for masking or altering the carbonates actually synthesized in a reducing environment. On the contrary, in our model, carbonate rock layers simply never formed, but instead we envisage the possibility of a warmer and wetter Mars, where extensive interaction between significant amounts of moderate to extreme acidic liquid water and a CO_2 atmosphere might have occurred (Fig. 3).

The viability of the model proposed here for ancient Mars is supported by the latest observational evidence provided by MER Opportunity at Meridiani Planum. Also on the modern Earth, the headwaters of the Tinto river system (in the southwest of Spain), an extreme acidic environment controlled by iron biogeochemistry, produces ferric-iron-enriched sediments dominated by sulphate and oxihydroxide parageneses, resulting in goethite, haematite and jarosite, analogous to the minerals found in Meridiani²⁸. In the living Tinto river system, the values of pH, redox potential, and

Table 1 Assumed average composition for surface waters on early Mars

Constituent	Concentration (p.p.m.)	Molality ($\times 10^{-3}$ mol kg ⁻¹)	Activity
$\text{SiO}_2(\text{aq})$	10.8–60.1	0.18*–1.0	$10^{-3.75}$ – 10^{-3}
$\text{Fe}^{2+}/\text{Fe}^{3+}$	3–44.7	0.05†–0.8	$10^{-4.3}$ – $10^{-3.1}$
SO_4^{2-}	17.3 (624–1,263)	0.18 (6.5–13.5)‡	$10^{-3.745}$ ($10^{-2.19}$ – $10^{-1.88}$)
Mg^{2+}	24.3	1.0	10^{-3}
Cl^-	23.0	0.65	10^{-7}
Ca^{2+}	20.0	0.5	$10^{-3.19}$
Na^{2+}	18.4	0.8	$10^{-3.1}$
K^+	2.7	0.07	$10^{-4.16}$

Concentration and molality data after ref. 8, except as shown by footnote symbols as follows.

*From ref. 30.

†From ref. 24.

‡Obtained assuming a solution at 273 K in equilibrium with jarosite for $a_{\text{Fe}^{3+}} = 10^{-4.3}$ and $a_{\text{Fe}^{2+}} = 10^{-3.1}$, respectively, as governed by the equation $\log a_{\text{SO}_4^{2-}} = 3\log a_{\text{H}^+} - 0.5\log a_{\text{K}^+} - 1.5\log a_{\text{Fe}^{3+}} - 2.91$.

HCO_3^- is not a fixed parameter here, as it has been swapped for $\text{CO}_2(\text{g})$, the y-axis variable in Fig. 1a.

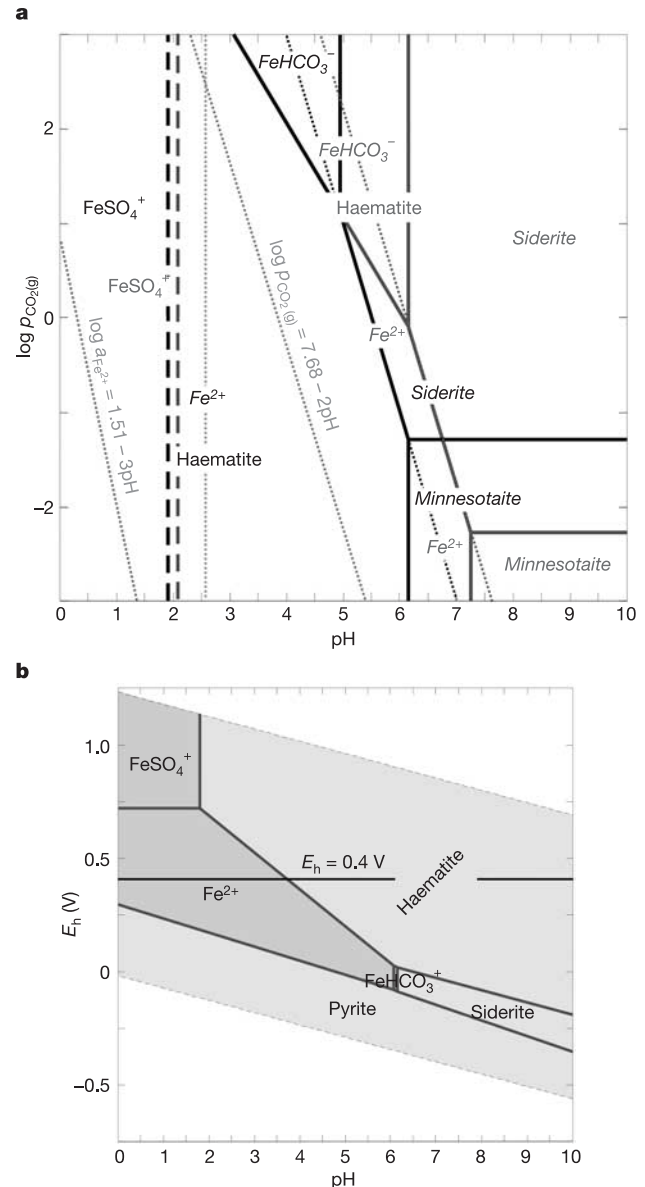


Figure 1 Stability boundaries of siderite and minnesotaite (under reducing conditions) and acidic solutions and haematite (under oxidizing conditions) on early Mars. **a**, Stability boundaries obtained as a function of CO_2 pressure and pH, calculated at 273 K for four different solutions: black lines represent a solution containing Fe and $\text{SiO}_2(\text{aq})$ at their upper limits (800 μM and 1 mM, respectively); grey lines represent the lower limits assumed here (50 μM and 180 μM); solid lines represent reducing conditions (Fe^{2+}) and standard $[\text{SO}_4^{2-}] = 0.18$ mM; and dashed lines represent oxidizing conditions (Fe^{3+}) and upper and lower limits for $[\text{SO}_4^{2-}] = 6.5$ (black) and 13.5 (grey) mM, driving to the formation of haematite and acidic solutions from which jarosite is expected to form at $\text{pH} \approx 2$ (boundary of jarosite stability). Dotted lines represent, from left to right: (1) the pH buffer when Fe^{3+} controls the solution, under acidic and oxidizing conditions, assuming goethite as the ultimate chemical product ($\text{Fe}^{3+} + \text{H}_2\text{O} \leftrightarrow \text{FeOOH}(\text{goethite}) + 3\text{H}^+$); (2) the equilibrium pH of jarosite ($(\text{K}_2\text{Fe}_6(\text{SO}_4)_4(\text{OH})_{12}) + 6\text{H}^+ \leftrightarrow 2\text{SO}_4^{2-} + 6\text{H}_2\text{O} + \text{K}^+ + 3\text{Fe}^{3+}$) when minimum values for Fe^{3+} (50 mM) and sulphates (180 mM) are considered; (3) the changes in pH exclusively controlled by the p_{CO_2} ; and (4 and 5) the relationship between CO_2 pressure and pH for each modelled solution ($\text{pH} = 5.6 - 0.5\log p_{\text{CO}_2}$ and $\text{pH} = 6.1 - 0.5\log p_{\text{CO}_2}$, respectively). All constituents included in the solutions for displaying this graphic are quantified in Table 1. **b**, E_{h} -pH diagram assuming the lower values of Table 1, at $p_{\text{CO}_2} = 1$ atm and $T = 273$ K. The stability fields of jarosite (acidic oxidizing solutions with FeSO_4^+), haematite and siderite are represented. Dark grey indicates phase solutions, while light grey indicates minerals. Jarosite formation is favoured by $E_{\text{h}} > 0.4$ V and $\text{pH} < 2$.

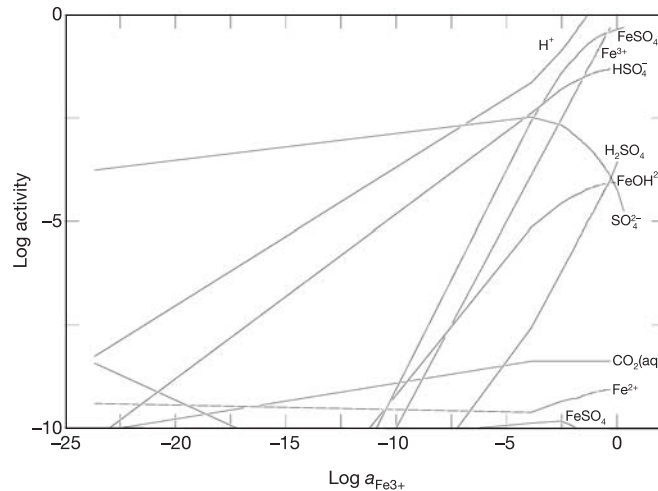


Figure 2 Evolution of the acidic martian environment over time. Evolution of solution and gas compositions, starting from an initial pH = 7 and $E_h = 0.4$ V, at increasing concentration of Fe^{3+} and H_2SO_4 . Fe^{3+} concentration ranges from 1.4×10^{-5} M to 0.5 M, a current record in modern acidic waters at the Tinto river²⁸. Although the ferric ion can acidify different solutions when added at high concentrations, sulphuric solutions generated by H_2S and SO_2 photo-oxidation may indeed generate acidic conditions at first, acting as an acidic matrix for later reactions that produce jarositic associations after

evaporation. Thus, the system has been coupled to an increased concentration of H_2SO_4 from the initial equilibrium to 10^{-4} M, a rough estimation for the acidic waters of the Tinto river²⁸. Although a high diversity of chemical species emerged, following the MER Opportunity results only S and Fe species are displayed, that is, those that would dominate the hypothetical chemistry of the early martian oceans. In this case, Fe^{3+} –sulphatic species would favour the ferric sulphate precipitation.

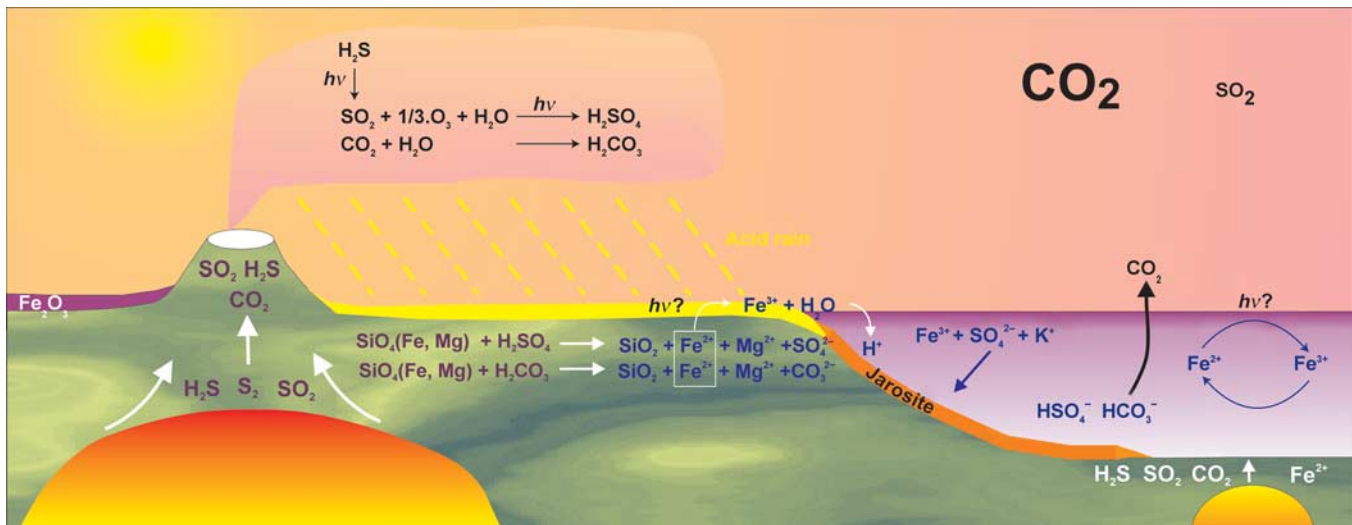


Figure 3 Schematic representation of atmosphere–land–ocean interactions generating acidic environments in early Mars. The acidic attack of ultramafic crust is driven by H_2SO_4 (generated by volcanic SO_2 photolysis) and H_2CO_3 , and consequently weathering of basalt releases Fe and Mg to the sea water. As ferrous iron oxidizes to ferric, Fe^{3+} drives the water more and more acidic, as also does SO_2 forming dilute sulphuric acid. From these solutions, jarosite as well as Mg sulphates are expected to precipitate, as highlighted by

MER Opportunity at Meridiani. Also, the acidic weathering process would generate a sulphate layer over the basaltic crust, potentially able to evolve to haematite by dehydration of the ferric sulphates and related iron-bearing minerals. Under such conditions, HCO_3^- , the main source of carbonates, is at very low concentration, and therefore all C is expected to be in the form of $\text{CO}_2(\text{aq})$ and $\text{CO}_2(\text{g})$, with higher residence time in the atmosphere and therefore maintaining long-term milder surface temperatures.

Fe^{3+} and sulphate concentration remain constant in different years under variable rain regimes. Life is highly diverse in the Tinto system, allowing us to suggest comparable ancient acidic aquatic habitats hosting a putative early biosphere on Mars. In fact, if biological inhabitation of early Mars is considered plausible, moderate acidic oceans represent the closest terrestrial analogue for a biogenic environment, similar to that where life originated on the Hadean–Early Archaean Earth^{23,26}. In the Tinto river system, the remobilization of iron and its subsequent precipitation as ferric

minerals preserves biological remains sometimes in remarkable detail, suggesting an analogous record in the rock exposures derived from the acidic martian early environments. □

Methods

We use the values listed in Table 1 to estimate limits on the pH and composition of the putative Noachian ocean, which includes considering the addition of supplementary cations such as Mg^{2+} , Ca^{2+} , K^+ or Na^+ , and silica (a by-product of the acid dissolution of ferromagnesian silicates, typical of martian igneous rocks in such acidic oceans), as they are effective solutes that raise the pH (ref. 8).

Sedimentation models in depositional environments establish that siderite (FeCO_3) is

always the first major carbonate to precipitate from ferrous solutions, because it is the most insoluble of the major common carbonates; whereas minnesotaite ($\text{Fe}_3\text{Si}_4\text{O}_{10}(\text{OH})_2$), a phyllosilicate compositionally close to greenalite, is the simplest of the possible ferrous silicates, as well as a stable phase in terrestrial sediments. According to the reactivity of siderite and minnesotaite given by their corresponding dissociation constants²⁹, the equations governing the limits of the stability fields for both minerals with respect to oceanic pH are $\text{pH} = 3.97 - 0.5 \log p_{\text{CO}_2} - 0.5 \log a_{\text{Fe}^{2+}}$ and $\text{pH} = 5.1 - 0.5 \log a_{\text{Fe}^{2+}}$, respectively, for amorphous precipitates.

The combination of pH, redox potential and concentration of CO_2 determines the stability of carbonates as possible phases, and their precipitation by inorganic processes depends on the saturation state of the solution with respect to these minerals. Oceanic pH specifically related to an equilibrium between aqueous iron and siderite can be determined from the relation $\log K = \log a_{\text{Fe}^{2+}} + \log p_{\text{CO}_2} - 2 \log a_{\text{H}^+}$, as calculated in Fig. 1a. Using values at 273 K and a mean concentration for Fe^{2+} in sea water of between 50 and 800 μM , a pH between 5.3 and 6.2 for the siderite- Fe^{2+} equilibrium is obtained for an ocean that occurs in assumed atmospheric CO_2 pressures of between 0.8 and 4 bar, contrasting to the pH value in modern oceans on Earth, about 8.1; when iron finally photooxidized to Fe^{3+} , pH varies between 1.9 and 2.1. Assuming silica concentrations close to the saturation point of oceanic waters on Earth (180 μM , ref. 30) or even higher for water on early Mars (1 mM, ref. 8), and assuming that no Earth-like biological recycling occurred on Mars during the Noachian period, these relations define the boundaries for siderite, minnesotaite, haematite and jarosite precipitation shown in Fig. 1. System evolution over time, coupled to an increased concentration of H_2SO_4 with Fe^{3+} , which would favour the precipitation of ferric sulphate, is shown in Fig. 2. For the conservative constraints assumed here (see Table 1), neither precipitation nor the metastability of siderite are noted. Increasing the assumed values will result in even more acidic conditions, thus making carbonate formation even less likely. Following the established model for the Earth, additional powerful contributors to the acidity of the Noachian oceans may have been volcanic sulphur-rich gases causing acid rain, as indicated in Fig. 3.

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A high-intensity highly coherent soft X-ray femtosecond laser seeded by a high harmonic beam

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Synchrotrons have for decades provided invaluable sources of soft X-rays, the application of which has led to significant progress in many areas of science and technology. But future applications of soft X-rays—in structural biology, for example—anticipate the need for pulses with much shorter duration (femtoseconds) and much higher energy (millijoules) than those delivered by synchrotrons. Soft X-ray free-electron lasers¹ should fulfil these requirements but will be limited in number; the pressure on beamtime is therefore likely to be considerable. Laser-driven soft X-ray sources offer a comparatively inexpensive and widely available alternative, but have encountered practical bottlenecks in the quest for high intensities. Here we establish and characterize a soft X-ray laser chain that shows how these bottlenecks can in principle be overcome. By combining the high optical quality available from high-harmonic laser sources (as a seed beam) with a highly energetic soft X-ray laser plasma amplifier, we produce a tabletop soft X-ray femtosecond laser operating at 10 Hz and exhibiting full saturation, high energy, high coherence and full polarization. This technique should be readily applicable on all existing laser-driven soft X-ray facilities.

A laser chain consists of several stages: a seed oscillator, delivering a pulse of perfect optical properties, followed by amplifiers, set to increase beam energy. We studied the extension of this scheme to the soft X-ray range. The main difficulties while designing an amplification chain are first to produce a seed with high optical quality, then to maintain the quality over the amplification stages and finally to reduce the spurious self-emission of the amplifiers to a