



A Whiff of Oxygen Before the Great Oxidation Event?

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- through the geological record, and time-series data from a broadly equivalent South African core are available (along with data tables) as supporting material on *Science* Online.
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- 18. All bulk samples were analyzed by the sulfur monoxide method in triplicate, and the reported values are averages of these measurements. Uncertainties are better than 0.3% for both $\delta^{34}\mathrm{S}$ and $\Delta^{33}\mathrm{S}$ values, which is comparable with uncertainties based on multiple standard measurements during each analytical session. For SF₆ analyses, uncertainties are 0.14, 0.008, and 0.20% for $\delta^{34}\mathrm{S}$, $\Delta^{33}\mathrm{S}$, and $\Delta^{36}\mathrm{S}$, respectively.
- 19. Monte Carlo resampling of data suggests that the two intervals carry a unique isotopic mean. In the lower half, we calculated means of $\delta^{34}S = 3.04~(\pm 0.50)$ and $\Delta^{33}S = 1.98~(\pm 0.44)$. In the upper half, including the organic and pyrite-rich horizon, the data indicate means of $\delta^{34}S = -1.41~(\pm 0.34)$ and $\Delta^{33}S = 1.10~(\pm 0.29)$. Considered alone, data from the pyritic interval between 153 and 135 m indicate means of $\delta^{34}S = -0.80~(\pm 0.47)$ and $\Delta^{33}S = -0.54~(\pm 0.17)$.
- 20. Atmospheric photochemistry is presently the only known mechanism that can account for the nonzero Δ³³S data and their relationship to Δ³⁶S values in the Archean record (1–4, 17). The principal source of S in the Archean atmosphere was volcanic (although biogenic sources may have also existed). Gas-phase photochemistry involving

- sulfur dioxide or sulfur monoxide has been shown in closed-cell photochemical experiments (2) to result in NMD sulfate (SO₄²⁻; with negative, and in some cases positive, Δ^{33} S values) and elemental S (S₈, with positive Δ^{33} S values). These reactions are sensitive to the wavelength of available ultraviolet radiation, and this parameter depends on, among other things, atmospheric O2 concentrations. The transfer pathways of S from the atmosphere to Earth's surface also depend on O2 concentration. An atmospheric model (4) constrains an upper limit of $<10^{-5}$ of present atmospheric levels of O_2 for the transfer of nonzero Δ^{33} S (and Δ^{36} S) containing SO₄²⁻ and S_o to Earth's surface: at a higher partial pressure of O2, the two S reservoirs homogenize in the atmosphere, and the photochemical signal is not preserved.
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- 28. A Monte Carlo resampling of the entire Mount McRae data set $(\Delta^{33}S = 1.25 \pm 0.17)$, which mirrors that of the entire Archean (27), points to a negative $\Delta^{33}S$ reservoir of the Archean S cycle largely lost from the geological record. Evidence from banded iron-formations, volcanic massive sulfide deposits, and other sea-floor environments may ultimately provide tests for a sulfate concentration gradient in the Archean ocean and solve the mystery of the missing negative $\Delta^{33}S$ reservoir. Ultimately, the balance of sources and sinks maintained generally low sulfate concentrations that allowed for spatial isotopic heterogeneities.

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Supporting Online Material

www.sciencemag.org/cgi/content/full/317/5846/1900/DC1 SOM Text Figs. S1 to S4 Tables S1 and S2 References

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A Whiff of Oxygen Before the Great Oxidation Event?

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High-resolution chemostratigraphy reveals an episode of enrichment of the redox-sensitive transition metals molybdenum and rhenium in the late Archean Mount McRae Shale in Western Australia. Correlations with organic carbon indicate that these metals were derived from contemporaneous seawater. Rhenium/osmium geochronology demonstrates that the enrichment is a primary sedimentary feature dating to 2501 ± 8 million years ago (Ma). Molybdenum and rhenium were probably supplied to Archean oceans by oxidative weathering of crustal sulfide minerals. These findings point to the presence of small amounts of O_2 in the environment more than 50 million years before the start of the Great Oxidation Event.

any lines of evidence point to a rapid rise in the partial pressure of atmospheric $O_2(P_{O_2})$ from $<10^{-5}$ times the present atmospheric level (PAL) between 2.45 and 2.22 billion years ago (Ga) (1, 2), a transition often referred to as the Great Oxidation Event (GOE).

The GOE could have been an immediate consequence of the evolution of oxygenic photosynthesis (3). Alternatively, O_2 biogenesis may be ancient (4). If so, the GOE was a consequence of an abiotic shift in the balance of oxidants and reductants at Earth's surface (5–8). This debate can

be addressed by looking for evidence of localized or short-lived concentrations of $\rm O_2$ before 2.45 Ga.

The abundances of some transition elements in sedimentary rocks are sensitive to the availability of $O_2(9)$. In particular, in the modern oxygenated environment, molybdenum (Mo) exists in rivers and oceans primarily as the unreactive molybdate ion (MoO_4^{2-}) . Oxidative weathering of Mo-bearing sulfide minerals in crustal rocks leads to the accumulation of Mo in the oceans, where it is the most abundant transition element (at a concentration of \sim 105 nM) (10, 11). The abundance of Mo in the oceans is reflected in

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pyritic marine sediments deposited under oxygendeficient conditions, where Mo is removed from solution in association with organic carbon (12, 13), probably after reacting with H_2S to form oxythiomolybdates ($MoO_{4-x}S_x^{2-}$) (14). In such sediments deposited today and through much of the Phanerozoic, Mo contents are typically >100 ppm versus ~1 ppm in average crust (10, 12, 13, 15, 16).

By comparison, on an anoxic Earth, Mo would be largely retained in unoxidized crustal sulfide minerals during weathering. Therefore, Mo concentrations in the oceans would be low, and organic-rich sediments would show little authigenic Mo enrichment as compared to modern equivalents. Similar logic applies to sulfur (S). In fact, studies of Mo and S concentrations and stable isotopes in black shales reveal systematic shifts in ocean budgets from the Archean through the Phanerozoic that are broadly consistent with the GOE and with another rise in P_{O_2} later in the Proterozoic (2, 17–19) (table S1). Rhenium (Re) and uranium (U) are also promising indicators because their aqueous geochemistry is similar to that of Mo.

Here we report Mo, Re, U, and S measurements, as well as other geochemical data obtained at high stratigraphic resolution in the Mount McRae Shale, deposited \sim 2.5 Ga in the Hamersley Basin, Western Australia (20, 21). Approximately 100 samples were analyzed from a freshly recovered continuous drill core obtained for this study (22) (Fig. 1, fig. S1, and table S2). These samples were also analyzed for S isotope variations as part of a companion study (23).

The core intersected two intervals containing pervasive pyritic carbonaceous shale, which we refer to as S1 (from 125.5 to 153.3 m) and S2 (from 173.0 to 189.65 m). Shales in both intervals contain several weight % (wt %) S and typically >3% total organic carbon (TOC), which is consistent with anoxic (and potentially sulfidic) bottom waters and the presence of H₂S in pore waters during these depositional intervals.

The most prominent feature of the data is the excursion in Mo content within S1 (Fig. 1). Mo concentrations below this layer are typically <5 parts per million (ppm), which is near the crustal value and is typical of Archean carbonaceous shales. Concentrations increase gradually up the section from the base of S1 to a peak value of ~40 ppm at 143 m and then decrease to <10 ppm by ~125 m. These variations and the Mo peak at ~143 m are more pronounced when plotted as aluminum (Al)–normalized enrichment factors (24). Viewed this way, Mo content

increases up the section by ~ 50 times before falling sharply over an interval of ~ 2 m. The Mo enrichment correlates with enrichments in TOC and Re and broadly coincides with variations in carbonate and S contents. However, U contents vary little through the section.

The coherent behavior of Mo, Re, and U makes it possible to use Re/Os geochronometry to verify that metal abundances were unaffected by remobilization. Postdepositional addition or loss of Re (or Os) would result in significant isochron scatter. We find that samples taken from 128 to 149 m define an isochron with mean square weighted deviation = 1.1 (22) (Fig. 2 and table S3) and an age of 2501.1 \pm 8.2 million years, which is consistent with previous ages for the Mount McRae shale (20, 21). The element enrichments are therefore primary sedimentary features (as is the lack of associated U variation) deposited at the boundary of the Archean and Proterozoic eons, at least 50 million years before the beginning of the GOE.

The Mo excursion cannot be explained by variable carbonate dilution, as documented by extreme enrichment factors (Fig. 1). Instead, the correlations of Mo with TOC are strong evidence of authigenic enrichment (Fig. 3). Such trends are common in sediments from modern anoxic

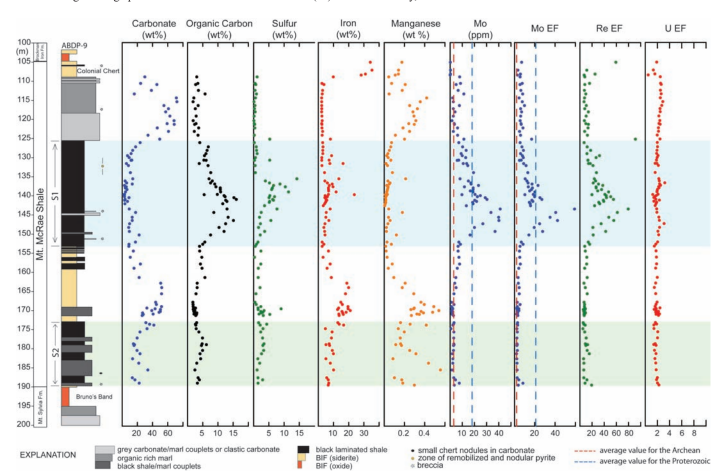


Fig. 1. Stratigraphy and geochemistry of the Mount McRae Shale, including percent of carbonate, TOC, S, Fe, Mn, Mo, Re, and U and EFs (24) for Mo, Re, and U (23). The intervals S1 and S2 span 125.5 to 153.3 m and 173.0 to

189.7 m, respectively. For comparison, dashed lines denote mean Mo concentrations and EFs in Archean and Proterozoic pyritic black shales, as indicated in the legend at bottom (18, 22) (tables S1 and S2).

basins where the concentration of H2S exceeds ~10 µM. In such sediments, Mo/TOC scales with Mo concentrations in deep waters (12). We recognize two trends within S1, corresponding to the zone of increasing Mo enrichment (~143 to 153 m) and the overlying zone in which Mo falls but remains elevated above average crustal values (~125 to 143 m). Mo/TOC slopes in these zones are $\sim 3.4 \pm 0.5~(\pm 1\sigma)$ ppm Mo/wt % TOC and \sim 1.8 \pm 0.2 (\pm 1 σ) ppm Mo/wt % TOC, respectively (25). By comparison, Mo/TOC is 4.5 to 25 ppm Mo/wt % TOC in pyritic sediments from modern anoxic basins (12) and averages ~26 ppm Mo/wt % TOC in Phanerozoic pyritic black shales (18) (table S1). Hence, small but substantial concentrations of dissolved Mo were present during S1 deposition. Similar reasoning can be applied to Re (fig. S2A).

These observations are not easily explained by hydrothermal inputs to the oceans. Enhanced hydrothermal input should result, first and foremost, in enrichments of iron (Fe) and manganese (Mn), yet the S1 unit is depleted in these elements relative to S2. In any event, hightemperature mid-ocean ridge-type systems should

Fig. 2. Re/Os isochron from the Mount McRae Shale, based on data from two subintervals within S1 (128.71 to 129.85 m and 145.22 to 148.32 m). MSWD, mean square weighted deviation. The Re/Os age, 2501.1 ± 8.2 Ma (initial 187 Os/ 128 Os = 0.04 ± 0.06), falls between prior ages of 2479 ± 3 Ma for the overlying Dales Gorge Member of the Brockman Iron Formation

(20) and 2561 \pm 8 Ma for

the underlying Bee Gorge

Member of the Wittenoom

Formation (21). Details are

discussed in (22).

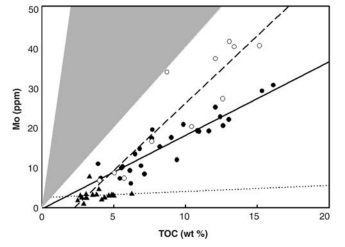
Fig. 3. Relationship between Mo and TOC in organic carbon-rich pyritic intervals in the Mount McRae Shale. Circles are from interval S1 (125.5 to 153.3 m). The metal-enriched zone of S1 below 143 m (open circles) is differentiated from the upper zone (solid circles). Triangles are from interval S2 (173.0 to 189.7 m). For comparison, the shaded region indicates the range of Mo/TOC slopes (forced through the origin) observed in modern sulfiderich anoxic basins.

be sinks, not sources, for Mo and Re because of the low solubilities of Mo and Re sulfides. A small amount of Mo enters seawater today as result of low-temperature hydrothermal seafloor weathering (13), but this Mo is probably derived from modern Mo-rich seafloor sediments.

Instead, these observations can be straightforwardly interpreted as evidence of oxidative weathering during S1 deposition. We hypothesize that O₂ in the shallow oceans and possibly in the atmosphere enhanced the rate of dissolution of submarine and subaerial sulfide minerals, such as molybdenite (MoS₂), that are important for the budgets of Mo and Re in igneous and metamorphic crustal rocks. Mo and Re released in this way would ultimately have produced authigenic enrichments in ocean sediments.

Sulfide minerals weather rapidly in the presence of O_2 , so P_{O_2} need not have been high. For example, even if P_{O_2} is only $\sim 10^{-5}$ PAL, a pyrite crystal of $100~\mu m^3$ volume will dissolve completely in $<\sim 20,000$ years (26,27). This is a short time compared to the likely duration of S1 (28). Consistent with such low P_{O_2} , Mo/TOC values in

Age = 2501.1 \pm 8.2 Ma 80 MSWD = 1.1 (n = 9)45.22 m 60 146.08 m 870s/1880s 148.32 m 47.10 m 40 129 85 m 20 128.84 m 400 800 1200 1600 2000 187Re/188Os



S1 do not exceed those of sediments accumulating in the modern Black Sea, which implies that the concentration of Mo in contemporaneous seawater was of similar magnitude as that in the deep waters of the Black Sea, or <1% that of fully oxygenated modern oceans.

The same process could have contributed to the excursion in S content and $\delta^{34} S$ in S1 (23). The long-term $\delta^{34} S$ record of sedimentary sulfides exhibits a negative shift between 2.4 and 2.3 Ga that is thought to indicate an increase in ocean sulfate concentrations. This increase is ascribed to an increased rate of oxidative weathering of pyrites in crustal rocks during and after the GOE (2). The negative shift in sedimentary $\delta^{34} S$ beginning at ~153 m in the Mount McRae Shale may record the effects of less extreme oxygenation at 2.5 Ga.

Our hypothesis of mild oxygenation is supported by the absence of U enrichment coincident with Mo and Re enrichments (Fig. 1) and the lack of correlation between U and TOC in S1 (fig. S2B), observations indicating that dissolved U concentrations were very low. U in the crust is primarily hosted by feldspars, zircon, apatite, and sphene, but not sulfides. Therefore the rate of release of U from rocks is only weakly affected by oxygenation, unlike that of Mo and Re; experimental studies suggest that the rate of pyrite oxidation exceeds that of feldspar minerals when $P_{\rm O_2} > 10^{-6} \text{ PAL } (29, 30). \text{ U may also be less}$ mobile than Mo and Re when O_2 is low (31). Hence, enhancements of Mo and Re influx without U enhancement are expected in the presence of small amounts of O2.

Our interpretation is also consistent with the extremely nonradiogenic initial \$^{187}Os/^{188}Os\$ in the Mount McRae Shale (Fig. 2). Such low values, also seen in shales \$\sim\!200\$ million years younger (32), indicate that the ocean Os budget was dominated by hydrothermal sources rather than by radiogenic Os derived from the weathering of high-Re/Os crustal rocks. As with U, oxidative weathering of sulfide minerals in igneous or metamorphic rocks might have had little effect on the balance between hydrothermal and crustal sources of Os to the oceans, because the Os content of crustal sulfide minerals, particularly molybdenite, can be low.

The low levels of O_2 that can account for our data are similar to the upper limit of 10^{-5} PAL for typical Archean P_{O_2} derived from the observation of nonzero Δ^{33} S in Archean sediments (33, 34), possibly explaining the juxtaposition of Mo and Re enrichments with the small nonzero Δ^{33} S signals seen throughout S1 (23). Alternatively, P_{O_2} above this threshold could have been present ephemerally within geographically restricted areas such as biologically productive regions of the oceans.

In contrast to the observations in S1, Mo concentrations and enrichment factors are very low below \sim 153 m, including in the organic carbon–rich and pyritic S2, where Mo is essentially invariant with TOC [Mo/TOC = 0.15 \pm

 $0.35~(\pm 1\sigma)$ ppm Mo/wt % TOC] (Fig. 3). Weak correlations appear between Mo and Al in S2, suggesting that at these depths the Mo budget was influenced by detrital components; such correlations are absent from S1. These observations point to much less authigenic Mo enrichment in S2 than in S1.

The difference in Mo enrichment suggests that the Mo inventory in overlying waters was much larger during S1 deposition than during S2 deposition, as would follow from an increase in environmental oxygenation up the section beginning at ~153 m. This interpretation is complicated by the fact that the Mo concentration differences between the units were also affected by differences in local depositional conditions that increased the efficiency with which Mo was transferred from water to sediments during S1 time (35). However, a difference in the dissolved Mo inventory and in ocean oxygenation provides a compelling explanation for the sharp difference in Mo/TOC between the units (36).

Other data from the core also point to greater surface ocean oxygenation above ~153 m, including changes in $\delta^{34}S-\Delta^{33}S$ systematics that may record the onset of an oxidative sulfur cycle (23). A redox shift can also explain differences in Fe and Mn concentrations above and below ~160 m (Fig. 1). Below this depth, most of the Fe is present as siderite (FeCO₃), and both elements are much lower in S1 than in S2. Fe and Mn would have been easily mobilized during anoxic weathering, enriched in anoxic Spoor Archean oceans and hence available for incorporation into sediments. Oxygenation of surface environments would have reduced the availability of both elements. At the same time, Re concentrations are slightly elevated above crustal average values throughout the core, and there is a positive correlation of Re with TOC in S2 as well as S1 (fig. S2B). Re can be more mobile than Mo during oxidative sulfide weathering (37), so this persistent Re enrichment suggests that some small degree of oxidative weathering occurred throughout.

The decrease in Mo content and Mo/TOC above 143 m may record a drop in the dissolved Mo inventory after its initial rise, even though the surface environment apparently remained persistently, if mildly, oxygenated (23). Re and S also decrease. Diagenetic complications notwithstanding (23), it is tempting to speculate that these decreases mirror a drop in atmosphere or ocean redox potential (38), as a result of biological or nonbiological feedbacks (39). However, declining trace metal abundances could simply reflect the exhaustion of exposed crustal sulfide sources, or areal expansion of sulfidic basins in the oceans in response to rising sulfate reduction, drawing down seawater Mo, Re, and S inventories.

The onset of oxidative weathering at 2.5 Ga was probably widespread. A recent examination of contemporaneous sediments from the

Ghaap Group in South Africa found that authigenic Mo and Re increased between 2.64 and 2.5 Ga (40), although that study did not have the stratigraphic resolution to capture the scale of variations reported here. Changes in S isotope systematics like those in the Mount McRae Shale also appear in time-correlative units from South Africa (23). Theoretical models show that a shift toward more oxidizing conditions can occur before the rise of an oxygenated atmosphere (5). Hence, the whiff of oxygen in the Mount McRae Shale may presage the global and irreversible transition to an oxygenated world.

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- 28. The duration of S1 is difficult to estimate but is <16 million years based on the uncertainty in the Re/Os isochron. Assuming a typical average shale accumulation rate of ~2.5 m/million years, the duration is ~11 million years. These estimates are consistent with accumulation rates determined from prior geochronology of sampled units (20, 21).
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- 36. We interpret this stratigraphic shift as indicating a change in environmental oxygenation with time. It is alternatively possible that the shift records sediments accumulating at different water depths in a redox-stratified water column. In either case, the Mo and Re enrichments are evidence of oxidative weathering ~2.5 Ga.
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- 38. Consistent with this notion, the rise in Mn content in carbonates above ~125 m without the associated Fe observed deeper in the core indicates segregation of these metals in waters that were only mildly oxygenated. Fe²⁺ and Mn²⁺ are immobile when oxidized to Fe³⁺ and Mn⁴⁺, respectively, because of the formation of insoluble oxyhydroxides. However, Fe²⁺ is oxidized at a lower Eh than Mn²⁺. Segregation is also facilitated by the slower oxidation and hydrolysis kinetics of Mn²⁺ versus Fe²⁺.
- 39. For example, biologically, rising O₂ could inhibit N₂ fixation or limit the availability of bioessential Fe, thereby reducing productivity and the rate of O₂ production. Nonbiologically oxidative erosion of a methane greenhouse would lead to lower surface temperatures, hence higher solubility of O₂ in surface oceans, lowering P_{O2}.
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- 41. The authors thank A. H. Knoll for inspiration and encouragement; B. Runnegar and R. Grymes for launching the Astrobiology Drilling Program; B. Blumberg for supporting the Mission to Early Earth Focus Group of the NASA Astrobiology Institute; and J. Farquhar, R. Raiswell, D. Johnston, A. Bekker, T. Algeo, J. Kasting, and J. Hannah for helpful discussions. Two anonymous reviewers helped improve the manuscript. J. S. R. Dunlop, L. C. Bonser, M. van Kranendonk, A. Hickman, and the Geological Survey of Western Australia assisted with core recovery. Funding was provided by the NASA Astrobiology Institute and the NSF Geobiology and Low Temperature Geochemistry program.

Supporting Online Material

www.sciencemag.org/cgi/content/full/317/5846/1903/DC1 Materials and Methods Figs. S1 and S2 Tables S1 to S3

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