The emergence of subaerial crust and onset of weathering 3.7 billion years ago

Desiree Roerdink (✉ Desiree.Roerdink@uib.no)
University of Bergen  https://orcid.org/0000-0002-1182-3606

Yuval Ronen
University of Bergen

Harald Strauss
University of Muenster

Paul Mason
Utrecht University

Keywords:

DOI: https://doi.org/10.21203/rs.3.rs-98804/v1

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Abstract

Reconstructing the emergence and weathering of continental crust in the Archean is crucial for our understanding of early ocean chemistry, biosphere evolution and the onset of plate tectonics. However, considerable disagreement exists between the elemental and isotopic proxies that have been used to trace crustal input into marine sediments and data are scarce prior to 3 billion years ago. Here we show that chemical weathering modified the Sr isotopic composition of seawater as recorded in 3.52-3.20 Ga stratiform barite deposits from three different cratons. Using a combination of Sr, S and O isotope data, barite petrography and a hydrothermal mixing model, we calculate a novel Sr isotope evolution trend for Paleoarchean seawater that is much more radiogenic than the curve previously determined from carbonate rocks. Our findings require the presence and weathering of subaerial and evolved (high Rb/Sr) crust from 3.7 ± 0.1 Ga onwards. This Eoarchean onset of crustal weathering affected the chemistry of the oceans and supplied nutrients to the marine biosphere 500 million years earlier than previously thought.

Background

One of the most accurate and complete records for tracing the weathering of continental crust over geological time are strontium isotopes ($^{87}\text{Sr}/^{86}\text{Sr}$) in marine chemical sediments. Strontium in seawater is derived from two sources with distinct radiogenic isotopic compositions: hydrothermal alteration of oceanic crust with low and mantle-like $^{87}\text{Sr}/^{86}\text{Sr}$, and subaerial weathering of continental crust with higher $^{87}\text{Sr}/^{86}\text{Sr}$ \textsuperscript{1}. Because Sr has a long residence time relative to the ocean mixing rate, $^{87}\text{Sr}/^{86}\text{Sr}$ in seawater is globally homogenized and the balance between oceanic and continental inputs can be recorded in the Sr isotope ratios of authigenic marine minerals. Typically the lowest $^{87}\text{Sr}/^{86}\text{Sr}$ (least radiogenic) are taken as the best estimate of seawater at any time since post-depositional alteration is most likely to increase Sr isotope ratios \textsuperscript{2}. Throughout the Phanerozoic, carbonate shells and limestones show many secular variations in seawater $^{87}\text{Sr}/^{86}\text{Sr}$ that can be linked to changes in seafloor spreading rates, as well as shifting tectonic, geographic and climatic controls on the Sr isotopic composition of riverine runoff \textsuperscript{3}. Further back in time, the seawater Sr isotope record becomes compromised by the scarcity of unaltered sedimentary carbonate rocks. Sparse Archean carbonates with highly unradiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ have therefore been interpreted to represent seawater, suggesting a mantle-dominated ocean chemistry at this time \textsuperscript{4, 5, 6}. This curve and the concomitant rise in seawater $^{87}\text{Sr}/^{86}\text{Sr}$ at the Archean-Proterozoic boundary supports models and proxies that argue for the late emergence of continental crust and onset of weathering around 2.5 Ga \textsuperscript{7, 8, 9, 10, 11, 12, 13}. However, recent work on two barite deposits suggested much higher seawater $^{87}\text{Sr}/^{86}\text{Sr}$ than the mantle-like value assumed from the Archean carbonate record at 3.2 Ga \textsuperscript{14, 15}. This finding questions the assumption that the unradiogenic carbonates truly reflect seawater, and challenges the validity of a mantle-like Sr isotope seawater evolution curve for the Paleoarchean. The earlier onset of weathering inferred from these barite data would be consistent with Ti isotopic evidence for emerged felsic crust at 3.5 Ga \textsuperscript{16}, weathering-induced decoupling of Hf and
Nd isotope systems in 3.4 Ga chert \(^{17}\), and Hf isotope ratios in black shales indicating weathering of evolved crust by 3.0 Ga \(^{18}\).

To correctly reconstruct the onset of crustal weathering and emergence of continental crust, here we investigate the Sr, S and O isotopic compositions of six stratiform marine-hydrothermal barite deposits from three different cratons covering a time window of 320 million years. Using a mixing model to assess hydrothermal influence on the barite \(^{87}\)Sr/\(^{86}\)Sr, we calculate a new Sr isotope evolution trend for Paleoarchean seawater and use this result to constrain the timing at which weathering began to modify early ocean chemistry.

**Does barite reflect seawater?**

We studied field and drill core samples from barite deposits at Londozi (3.52 Ga, Theespruit Fm), Vergelegen (3.41 Ga, Kromberg Fm), Stentor/Amo (3.26 Ga, Bien Venue Fm) and Barite Valley (3.24 Ga, Middle Mapepe Fm) in the Kaapvaal craton, North Pole (3.49 Ga, Dresser Fm) in the Pilbara craton and Sargur (3.20 Ga, Sargur Group) in the Dharwar craton (Table S1). All barite deposits occur in volcanic-sedimentary successions consistent with shallow to deep marine environments \(^{19,20,21,22,23,24,25}\).

Multiple sulfur isotope data provide evidence for atmospherically-derived sulfate in the barites from a well-mixed global seawater reservoir \(^{26,27,28}\) or local felsic volcanic eruptions \(^{29,30}\). In contrast, field observations suggest a hydrothermal origin for the barium \(^{25}\). In all localities, barite is strongly associated with chert \(^{21}\), silica dykes feed into barite horizons at Barite Valley and North Pole \(^{25,31}\), and barium-rich hydrothermal alteration zones with Ba-feldspar underlie barite at Londozi and Sargur \(^{19,29}\). However, all deposits lack polymetallic sulfide deposits, indicating that hydrothermal fluid temperatures were relatively low and likely below 150°C \(^{32}\). Low-temperature hydrothermal activity is also consistent with the lack of an underlying magmatic system, absence of sintering and the small vertical extent of chert dykes at Barite Valley \(^{25}\).

Two types of barite are observed in the six deposits: bladed barite consisting of course blades up to several centimeters long, and fine-grained granular barite (Fig. S1). Crystal morphology alone does not reveal the origin of the barite. However, a combination of sedimentological evidence and morphology supports a primary origin for bladed barite at Barite Valley, except for some isolated blades cutting through barite sands that appear to have grown diagenetically \(^{23,25}\). A primary origin for bladed barite is consistent with the well-formed, tabular to bladed crystal morphology that is predicted to grow in settings with low to moderate degrees of barite oversaturation \(^{33}\) and is found in modern hydrothermal settings \(^{34}\). In contrast, granular barite is often found in association with heavy minerals and reworked quartz, indicating a detrital origin for this morphological type \(^{23,25}\). Equigranular textures can also form by recrystallization of barite during diagenesis or metamorphism that has affected all deposits \(^{35}\), from lower greenschist facies (300-400°C) at North Pole, Barite Valley and Vergelegen \(^{36,37}\) to upper greenschist facies (400-500°C) at Stentor/Amo \(^{20}\) and amphibolite facies (500-650°C) at Londozi and
Sargur. In order to determine which barite can be used to constrain Paleoarchean seawater $^{87}\text{Sr}/^{86}\text{Sr}$, field data and mineral morphology must therefore be integrated with geochemical proxies.

Within individual deposits, bladed barite samples are Sr isotopically distinct from granular barite. Measured $^{87}\text{Sr}/^{86}\text{Sr}$ values are lowest in bladed barite (Fig. 1a), with weighted averages (± 95% confidence intervals) ranging from 0.700562 ± 0.00015 at Londozi (n = 2), 0.700841 ± 0.00004 at Vergelegen (n = 6), 0.701295 ± 0.00008 at Barite Valley (n = 4) to 0.701333 at Sargur (Table S2). No bladed barite was found in samples from the Stentor/Amo deposit. In contrast, granular barite is characterized by higher average $^{87}\text{Sr}/^{86}\text{Sr}$ values of 0.700757 ± 0.00010 at Londozi (n = 21), 0.701112 ± 0.00001 at Vergelegen (n = 2), 0.701240 ± 0.00013 at Stentor/Amo (n = 5), 0.701478 at Barite Valley and 0.701814 ± 0.00033 at Sargur (n = 3). These higher values cannot be explained by in situ $^{87}\text{Rb}$ decay, because measured Rb concentrations in acid-leached fractions from both types of barite are very low and would require corrections less than our analytical precision (Table S3). In addition, we carefully selected least weathered samples to avoid contamination with high $^{87}\text{Sr}/^{86}\text{Sr}$ phases, as exemplified by the two highly weathered samples from Barite Valley (Fig. 1a).

Unlike the marine carbonate record, the lowest $^{87}\text{Sr}/^{86}\text{Sr}$ values in bladed barite cannot be unambiguously interpreted to reflect seawater as ratios may have been lowered by hydrothermal input of unradiogenic Sr. We therefore combine $^{87}\text{Sr}/^{86}\text{Sr}$ data with oxygen and sulfur isotopic compositions to select which barite is most representative of seawater (Fig. 1b-d). Bladed and granular barite is characterized by $\delta^{18}\text{O}$ and $\delta^{34}\text{S}$ values that fall within the range reported for Paleoarchean seawater sulfate from sulfate minerals and carbonate-associated sulfate (Fig. 1b, 1c). Seawater-like $\delta^{18}\text{O}$ and $\delta^{34}\text{S}$ values do not give direct evidence for seawater Sr isotope ratios in hydrothermal barite, since the source of Sr is not directly coupled to that of SO$_4^{2-}$. However, the observation above is consistent with the low-temperature hydrothermal settings inferred from field data, as modification of seawater-like $\delta^{18}\text{O}$ values is expected above 150°C due to rapid oxygen isotope exchange between dissolved sulfate and water. This in turn suggests that measured $^{87}\text{Sr}/^{86}\text{Sr}$ have been relatively little affected by Sr from a non-seawater source, because leaching of Sr from rocks is limited at these low temperatures and low- to intermediate-temperature hydrothermal fluids are dominated by seawater-derived Sr.

Importantly, we observe the highest $\delta^{18}\text{O}$ values for each deposit in bladed barite and in association with the most negative, and therefore most seawater-like, anomalous sulfur isotope signatures (D$_{33}\text{S}$, see Methods for calculation, Table S2). These samples also display a strong positive correlation ($R^2 = 0.95$) between $^{87}\text{Sr}/^{86}\text{Sr}$ and $\Delta^{33}\text{S}$ (Fig. 1d), in contrast to a weaker correlation for granular barite ($R^2 = 0.64$, not shown in Fig. 1d). Previous work has demonstrated that the magnitude of seawater sulfate D$_{33}\text{S}$ decreases throughout the Paleoarchean, as shown in Fig. 1d. The observed correlation between Sr and S isotopes in bladed barite is therefore best explained by co-evolution of D$_{33}\text{S}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ in seawater due to progressive decay of $^{87}\text{Rb}$. In contrast, this correlation may have been blurred in the
granular barite as a result of alteration or contamination, which is consistent with granular textures resulting from recrystallization processes. At water-rock ratios of 1 to 10, metamorphic fluids with 50-1000 ppm Sr and $^{87}\text{Sr}/^{86}\text{Sr} \sim 0.703-0.706$ can shift Sr isotopic compositions from those measured in the bladed barite towards the higher values in granular barite samples (Fig. S2).

Based on the O, S and Sr isotope systematics outlined above, we conclude that $^{87}\text{Sr}/^{86}\text{Sr}$ of bladed barite is as close to Paleoarchean seawater as possible for a hydrothermal deposit. The $^{87}\text{Sr}/^{86}\text{Sr}$ values in the bladed barite samples define a strong regression line (Fig. 2, $R^2 = 0.98$), and are more radiogenic than the Paleoarchean primitive mantle (Fig. 1a) calculated from the Basaltic Achondrite Best Initial (BABI) at 4.56 Ga of $^{87}\text{Sr}/^{86}\text{Sr} = 0.69897$ and bulk Earth $^{87}\text{Rb}/^{86}\text{Sr} = 0.085$. Values also plot above estimates of the depleted mantle based on $^{87}\text{Sr}/^{86}\text{Sr} = 0.703$ for modern mid-ocean ridge basalt and an initial value of $^{87}\text{Sr}/^{86}\text{Sr} = 0.69950$ (DM1) or 0.69897 (DM2) (Fig. 1a). Our findings are consistent with previously reported Sr isotopic compositions of the Barite Valley, Sargur, Vergelegen and North Pole deposits. To go further, we explore next the hydrothermal influence on barite $^{87}\text{Sr}/^{86}\text{Sr}$ values to assess the Sr isotopic composition of the Paleoarchean oceans.

**Paleoarchean seawater evolution trend**

We use a hydrothermal mixing model to calculate a plausible seawater Sr evolution trend from the $^{87}\text{Sr}/^{86}\text{Sr}$ values measured in bladed barite samples, based on the low-temperature hydrothermal setting inferred from field and oxygen isotope data. In our model (see Methods), we calculate mixing ratios of seawater (20°C, 0.6M NaCl salinity) and hydrothermal fluid (150°C) that lead to oversaturation with respect to barite. We use a local seawater sulfate concentration of 8 µM based on observed 12-20‰ differences between barite and pyrite $^{34}\text{S}$, and assume the same $\text{SO}_4^{2-}$ concentration for the hydrothermal fluid as anhydrite precipitation and thermochemical sulfate reduction were likely negligible at 150°C. Our calculations for a range of Ba concentrations show that the highest degree of oversaturation occurs for a mixture consisting of 10-40% hydrothermal fluid (Table S5). Calculated saturation indices are low (<0.2), which is consistent with our interpretation above that bladed barite formed as primary crystals.

We use this result and a two-component mixing model to calculate $^{87}\text{Sr}/^{86}\text{Sr}$ values of seawater-hydrothermal fluid mixtures from which barite precipitated. By varying the input value for seawater $^{87}\text{Sr}/^{86}\text{Sr}$, we assess which seawater composition is feasible with the highest and lowest $^{87}\text{Sr}/^{86}\text{Sr}$ values measured in bladed barite from Londozi, Vergelegen and Barite Valley. To constrain the Sr isotopic composition of the hydrothermal fluids, we assume that Sr is predominantly derived from seawater (80%) with a small contribution from leached crust (20%), as observed in low-temperature (150°C) hydrothermal experiments and modern low-intermediate temperature hydrothermal fluids, and consistent with seawater-dominated REE patterns in Paleoarchean alteration zones. For each deposit, we explore $^{87}\text{Sr}/^{86}\text{Sr}$ variations during hydrothermal leaching of mafic and felsic rocks, represented by Sr isotope evolution curves for the depleted mantle and continental crust, respectively.
From our constraints on the chemical conditions required for barite precipitation and comparison of theoretical seawater-hydrothermal mixtures with those recorded in the bladed barite, we obtain a seawater Sr isotope evolution trend with a slope corresponding to $^{87}$Rb/$^{86}$Sr values of 0.194-0.198 (Fig. 2). Sr isotopic compositions of single bladed barite samples from North Pole and Sargur fall well within the predicted trend, as well as granular barite from the Stentor deposit. Calculated $^{87}$Rb/$^{86}$Sr values are substantially higher than the estimated value for the early Archean depleted mantle (0.07 ± 0.007) $^{50}$ and contrast strongly with the mantle-dominated curve inferred from carbonate $^{87}$Sr/$^{86}$Sr $^{4,5,6}$ (Fig. 3). Our results empirically constrain the seawater Sr isotope evolution trend significantly further back in time compared to the curve predicted from the extrapolation of 3.2 Ga barite $^{87}$Sr/$^{86}$Sr $^{15,56}$ (Fig. 3).

### Onset of subaerial crustal weathering

The radiogenic Sr isotope values for 3.52-3.20 Ga Paleoarchean seawater imply detectable weathering of an emerged and felsic crustal source at least 300 million years further back in time than what has previously been reported at 3.2 Ga $^{15}$. We further advance this by constraining the start of subaerial weathering from the intersection of our seawater evolution trend and mantle curves, which represents the time at which the input of crustal Sr started to modify the $^{87}$Sr/$^{86}$Sr of seawater away from mantle-dominated values. Our calculated trend for Paleoarchean seawater $^{87}$Sr/$^{86}$Sr indicates an onset at approximately 3.7 ± 0.1 Ga based on the intersection with the primitive mantle curve, 3.6 ± 0.1 Ga from depleted mantle curve DM1 and 3.8 ± 0.1 Ga from depleted mantle curve DM2 (Fig. 2). Improved constraints on the Paleoarchean Sr isotope mantle curve are required to further reduce the uncertainty on this estimate. However, the late Eoarchean onset of crustal weathering observed from chemical signatures is consistent with the siliciclastic rock record which shows that physical weathering products appear for the first time in greenstone belts around 3.4 Ga $^{57}$ and possibly as early as 3.7 Ga $^{58}$ (Fig. 3), as well as examples of exposed land surfaces by 3.5 Ga $^{59}$ and evidence for the existence of felsic crust at this time $^{60}$. Our findings indicate that weathering substantially modified the Sr isotope budget of Paleoarchean seawater, in contrast to its oxygen isotope composition that was recently shown to be unaffected by this $^{12}$.

Comparison of our Sr isotope data and revised seawater trend with other records of weathering (e.g. $^{6,11,61}$) highlights the scarcity of data in the Paleoarchean and the importance of the barite record for this period in Earth’s history. Paleoarchean carbonate from the Pilbara and Kaapvaal craton $^{1,5}$ displays significantly more radiogenic $^{87}$Sr/$^{86}$Sr than barite (Fig. 3), reflecting the higher preservation potential of insoluble barite during diagenesis and metamorphism $^{35}$. The combined updated barite (blue lines in Fig. 3) and carbonate $^{56}$ (green line in Fig. 3) $^{87}$Sr/$^{86}$Sr record for the Archean shows little secular variation from 3.7 to 2.5 Ga, which differs from the recently predicted sharp rise in seawater $^{87}$Sr/$^{86}$Sr at 3.2 Ga $^{56}$. The absence of strong secular variations in the Archean $^{87}$Sr/$^{86}$Sr record may be partially explained by the low temporal resolution of the data compared to the Phanerozoic, although a linear Sr isotope trend in the Paleoarchean is consistent with small $^{87}$Sr/$^{86}$Sr variations in newly formed crust due to little time for
radiogenic ingrowth of $^{87}$Sr. It also tentatively suggests no measurable long-term shifts in tectonic, geographic and climatic controls on seawater $^{87}$Sr/$^{86}$Sr throughout the Archean, in contrast to the changes observed in carbonate $^{87}$Sr/$^{86}$Sr from the Proterozoic and Mesozoic into the Cenozoic $^{62}$.

If the extrapolation of our seawater Sr isotope trend is correct, it implies that the late Eoarchean geodynamic regime generated granitic magmas and sufficient continental freeboard to support weathering of emerged felsic crust from 3.7 Ga. The globally significant changes in seawater $^{87}$Sr/$^{86}$Sr defined by our samples from three different cratons suggest that subaerial weathering was a relatively widespread phenomenon, likely requiring a larger relative area of emerged crust than the 4% previously reported for the Neoarchean (see Supplementary Information) $^{63}$. Following the same reasoning as above, the absence of radiogenic Sr in seawater before 3.7 Ga suggests that prior to this time the extent of felsic landmass and subaerial weathering was very limited. This in turn would have hampered the colonization of land, inhibited the supply of nutrients to the oceans $^{64}$ and restricted the extent of epicontinental seas in the early Eoarchean, possibly limiting the evolution of photosynthetically fueled ecosystems. The lack of undisputed evidence for microbial activity in supracrustal rocks older than 3.5 Ga may reflect such an Eoarchean planet with fewer favorable environments for life to flourish than in the Paleoarchean, when crustal emergence and weathering facilitated life in shallow marine settings.

**Methods**

**Analytical**

Barite powders were drilled from the same spots as used for multiple sulfur isotope analysis $^{27}$ using a diamond microdrill. Circa 100 mg of pure barite was reacted with concentrated HF to form insoluble SrF$_2$, which was centrifuged together with undissolved BaSO$_4$ to remove the supernatant. Strontium was subsequently leached from the residual solids with 2N HNO$_3$ under sub-boiling conditions for 2 days. The dissolved fraction was purified on an Eichrom Sr Spec ion exchange resin and analyzed for Sr isotopes using a Finnigan MAT 262 Thermal Ionization Mass Spectrometer at the University of Bergen. Values are normalized relative to $^{87}$Sr/$^{86}$Sr = 0.710240 for standard SRM987, with an error (2σ) of 0.000009. Aliquots of the supernatant after the HF dissolution step and the dissolved fraction after the 2N HNO$_3$ dissolution step were analyzed for Rb and Sr concentrations using a ThermoScientific Element XR inductively-coupled plasma mass spectrometer at the University of Bergen.

Oxygen isotopes were measured in a selection of barite powders from Londozi, Vergelegen and Stentor/Amo using a ThermoFinnigan TC/EA high-temperature pyrolysis unit coupled to a ThermoQuest Delta Plus XL mass spectrometer at the University of Münster $^{40}$. Results are reported relative to Vienna Standard Mean Ocean Water (V-SMOW) with an average analytical precision of 0.19‰ on δ$^{18}$O. In addition, multiple sulfur isotopes were measured in samples from Sargur after converting barite into sulfide by boiling with a Thode reduction solution (concentrated HCl, HI and H$_3$PO$_4$). Isotope analyses were performed at the University of Maryland on a dual-inlet ThermoFinnigan MAT 253 following
cryogenic and gas chromatographic purification, with an average external precision of 0.1‰ on δ^{34}S and 0.02‰ on δ^{33}S (reported relative to V-CDT) \(^{27}\). Values of δ^{33}S are calculated as \(\delta^{33}S = \delta^{34}S - 1000[(1+\delta^{34}S/1000)^{0.515} - 1]\).

**Model**

Barite solubility products at 0.6M NaCl (modern seawater salinity) were approximated by linear extrapolation of experimental results at 0.2M NaCl and 1M NaCl and 25-250°C \(^{53}\). The temperature of the hydrothermal fluid was estimated at 150°C based on element enrichment patterns and oxygen isotopes in similar silica alteration zones \(^{32}\), and seawater temperature was estimated at 20°C based on isotopic evidence for modern-like temperatures \(^{51}\). Sulfate concentrations were constrained from the model of Crowe et al. \(^{54}\) for sulfate-dependent microbial isotope effects and measured differences in δ^{34}S of 12-20‰ between barite and associated biogenic pyrite minerals \(^{55}\). The saturation index SI was calculated as \(\log Q/K_{sp}\) with \(Q = [Ba^{2+}][SO_4^{2-}]\) for different mixtures of hydrothermal fluid and seawater, assuming mixing temperatures without conductive cooling and Ba concentrations in the hydrothermal fluid and seawater at 75%, 90% and 99% of the solubility concentrations at 150°C and 20°C, respectively (Table S5).

Based on the obtained mixing ratio of hydrothermal fluid (HF) and seawater (SW) during barite precipitation, barite (BA) Sr isotopic compositions were predicted using the following equation modified from Jamieson et al. \(^{34}\):

\[
\frac{^{87}Sr}{^{86}Sr_{EA}} = \frac{^{87}Sr/^{86}Sr_{SW} - \left(\frac{^{87}Sr}{^{86}Sr_{SW}} \times \%HF\right) + \left(\frac{[Sr]_{HF}}{[Sr]_{SW}} \times \frac{^{87}Sr}{^{86}Sr_{HF}} \times \%HF\right)}{1 + \left(\%HF \times \frac{[Sr]_{HF}}{[Sr]_{SW}} - \%HF\right)}
\]

with \%HF as the fraction of hydrothermal fluid in the mixture during barite precipitation (0.1 to 0.4) and \([Sr]_{HF}/[Sr]_{SW}\) as the ratio of Sr concentrations in the hydrothermal fluid versus seawater. Results were calculated for \([Sr]_{HF}/[Sr]_{SW}\) ratios of 0.5, 1, 2 and 3 based on enrichment factors observed in modern seafloor hydrothermal fluids \(^{35}\). Sr isotopic compositions of hydrothermal fluids were calculated as:

\[
^{87}Sr/^{86}Sr_{HF} = 0.8 \times ^{87}Sr/^{86}Sr_{SW} + 0.2 \times ^{87}Sr/^{86}Sr_{L}
\]

with \(^{87}Sr/^{86}Sr_{L}\) as the isotopic composition of Sr leached from host rocks and fractions of seawater and rock-derived Sr based on experimental results at 150°C \(^{45}\). Barite Sr isotopic compositions were
calculated for two end-member scenarios: (1) leaching of mafic rocks, with $^{87}\text{Sr}/^{86}\text{Sr}_L$ obtained from linearly extrapolating BABI (0.69897)$^{47}$ to the average modern mid-ocean ridge basalt value of 0.703 (DM2) at the age of each deposit, and (2) leaching of felsic rocks, with $^{87}\text{Sr}/^{86}\text{Sr}_L$ constrained by linearly extrapolating BABI to the average modern crust value of 0.7119$^9$. Calculations were iterated for different input values of $^{87}\text{Sr}/^{86}\text{Sr}_\text{SW}$ to obtain the highest and lowest possible seawater Sr isotope composition that can explain the range in $^{87}\text{Sr}/^{86}\text{Sr}$ measured in bladed barite samples. No calculations were performed for Stentor/Amo and Sargur due to insufficient bladed barite samples.

Declarations

Acknowledgements

We thank Axel Hoffman for samples from the Sargur barite, Siv Hjorth Dundas for ICP-MS analyses, and Joost Hoek, James Farquhar and the University of Maryland stable isotope laboratory for multiple sulfur isotope analyses. Cedric Hamelin and Helen King are thanked for discussions, and Andreas Beinlich, Douwe van Hinsbergen, Hari Tsikos and Mark van Zuilen are thanked for critically reading through an earlier version of the manuscript and providing constructive comments. Aaron Satkoski and two anonymous reviewers are thanked for providing valuable comments that have improved the manuscript significantly. This research was funded by the Research Council of Norway through the Centre for Geobiology and fieldwork in South Africa was supported by the Stichting Dr Schurmannfonds.

Author contributions

D.R. conceived the study, D.R. and P.M. provided and prepared samples, Y.R. and H.S. measured data, D.R. and PM. interpreted data and wrote the manuscript.

Data availability

Strontium isotope data obtained in this study is available in Supplementary Table S2 and the Pangaea data repository under doi.pangaea.de/10.1594/PANGAEA.913541. We have used published strontium isotope data for Barite Valley, Sargur and North Pole barite samples from $^{14,15,50}$, sulfur isotope data for Londozi, North Pole, Vergelegen, Stentor/Amo and Barite Valley from $^{27}$ and oxygen isotope data for Barite Valley from $^{40}$. Ages of the barite deposits are from $^{19,21,23,27,29,31}$. Sr isotope data for Archean carbonate samples were obtained from $^{56,65}$.

Competing interests

The authors declare no competing interests.

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**Figures**
Figure 1

Sr, S and O isotopic compositions of Paleoarchean stratiform barite. (a) Measured $^{87}$Sr/$^{86}$Sr in bladed, granular and weathered barite versus age, with the primitive and depleted mantle curves shown for comparison (see text for calculations). Deposit names are indicated, with BV = Barite Valley. (b) Measured $^{87}$Sr/$^{86}$Sr in bladed and granular barite versus $\delta^{18}$O (this study, 40), with the published range of seawater sulfate $\delta^{18}$O 15, 26. (c) Measured $^{87}$Sr/$^{86}$Sr in bladed and granular barite versus $\delta^{34}$S (this study, 27), with the published range of seawater sulfate $\delta^{34}$S 26, 41, 42, 66. (d) Measured $^{87}$Sr/$^{86}$Sr in bladed and granular barite versus $\Delta^{33}$S (this study, 27), showing the linear trend (with 95% confidence intervals) defined by the bladed barite samples reflecting temporal evolution of seawater. Error bars are indicated when errors are larger than the symbols.
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Figure 2

Measured and modelled Paleoarchean seawater curves. Sr isotope evolution trend for Paleoarchean seawater calculated from the hydrothermal mixing model and bladed barite $\text{Sr}^{87}/\text{Sr}^{86}$ (dark blue lines). Also shown is the regression line for the bladed barite (light blue line with 95% confidence intervals). Primitive and depleted mantle curves are the same as those shown in Fig. 1a (see text for calculations). The onset of large scale subaerial crustal weathering is defined by the intersection of the seawater and mantle curves at $3.7 \pm 0.1$ Ga.
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![Figure 3](image)

 Compilation of Archean barite and carbonate Sr isotope data. Barite $^{87}\text{Sr}/^{86}\text{Sr}$ from this study (blue diamonds and yellow circles), published barite $^{87}\text{Sr}/^{86}\text{Sr}$ (green triangles) 14, 15, 50 and published carbonate $^{87}\text{Sr}/^{86}\text{Sr}$ (grey triangles) 9, 56, 65 for the Paleoarchean, Mesoarchean and Neoarchean. Our seawater Sr isotope evolution trend is shown (dark blue lines, cf. Fig. 2) in comparison with the curve predicted by Satkoski et al.56 based on 3.2 Ga barite and 2.94-2.8 Ga carbonate (green line), and the carbonate-based mantle-dominated seawater curve by Shields and Veizer (grey line) 6. The first appearance of siliciclastic rocks in the rock record is indicated for each craton on the lower axis, with DW = Dharwar craton, KV = Kaapvaal craton, PB = Pilbara craton and IS = Isua Greenstone Belt (see Table S6 for details and references).
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