# THE COMPOSITION AND WEATHERING OF THE CONTINENTS OVER GEOLOGIC TIME

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# Abstract

The composition of continental crust records a history of construction by tectonics and destruction by physical and chemical erosion. Quantitative constraints on how both igneous addition and chemical weathering have modified the continents' bulk composition are essential for understanding the evolution of geodynamics and climate. We have extracted temporal trends in sediments' protolith composition and weathering intensity from the largest available compilation of sedimentary major-element compositions, of ~ 15,000 samples from 4.0 Ga to the present. To do this we used a new analytical method which inverts whole sedimentary compositions for protolith composition and weathering intensity simultaneously. We find that the average Archean upper continental crust was silica rich and had a similar compositional diversity to modern continents. This is consistent with an early-Archean, or earlier, onset of plate tectonics. In the Archean, chemical weathering was ~ 25 % more efficient at sequestering CO<sub>2</sub> than in subsequent time periods. Since 2.0 Ga, over long (> 0.5 Ga) timescales, the crustal weathering intensity has remained largely constant. On shorter timescales over the Phanerozoic, the intensity of weathering is correlated to global climate state, consistent with silicate weathering feedback acting in response to changes in CO<sub>2</sub> outgassing.

Keywords Sedimentary Geochemistry · Weathering · Provenance · Archean Continents · Crustal Evolution · Phanerozoic Climate

## INTRODUCTION

The rocks at Earth's planetary surface are compositionally divided between dense, silica-poor oceanic crust and a buoyant, silica-rich continental crust. It is generally accepted that this dichotomy was generated and is maintained by plate tectonics. When the first felsic continents emerged remains highly debated. The Archean (i.e., 2.5 - 4.0 Ga) continental crust, is often viewed as dominated by basaltic rocks, and having an overall mafic composition similar to the oceanic crust (Taylor and McLennan 1986; Tang et al. 2016; Smit and Mezger 2017; Chen et al. 2019). This view suggests a relatively late onset for plate tectonics during the Neoarchean,  $\sim 2.5$  Ga. However, recent analyses of the sedimentary trace element and titanium isotope record suggest instead evolved, silica-rich Archean continents (Greber et al. 2017; Greber and Dauphas 2019; Keller and Harrison 2020; Ptáček et al. 2020). This view suggests an earlier onset for plate tectonics during the Paleo- or Eoarchean, before 3.5 Ga. Resolving this debate is essential for determining when and how plate tectonics began on Earth.

Not only is the history of the crust necessary for understanding geodynamics, the reaction of the crust with the hydrosphere is integral to the stabilisation of the planet's climate (Broecker and Langmuir 1985). Continental chemical weathering (the alteration of silicate minerals by reaction with water at Earth's surface) transfers atmospheric  $CO_2$  into carbonate minerals deposited on the ocean floor. This reaction is thus the major long-term sink for  $CO_2$  outgassed by the mantle (Walker et al. 1981).

The geochemical composition of sedimentary rocks is our primary record of crustal evolution on Gyr timescales. However, this archive is challenging to interpret. Chemical weathering strips sediments of mobile elements which means their composition is altered relative to the rocks from which they derive. Signals of changing crustal composition are thus obscured by alteration. In addition, most sediments record the signals of the local catchment they come from, not the continental crust as a whole. In this paper, we attempt to provide provide new perspectives into the long-term composition and alteration of the upper continental crust (UCC. To do this we use novel data analytical methods and the extensive geochemical database produced by the Sedimentary Geochemistry & Paleoenvironments Project.

# METHODS AND DATA

Most studies aiming to track changes in crustal composition claim to circumvent the alteration of sedimentary compositions by carefully selecting immobile elemental ratios. Whilst this approach could be successful at resolving protolith changes, by design it cannot provide information on how weathering has changed through time. In this study we extract signals of both the weathering intensity of sediments and protolith composition at once.

To do this we use a new method which explains the majorelement (Si, Al, Fe, Mg, Na, Ca, K) composition of sediments in terms of the composition of their protolith, and the intensity of weathering they have experienced (Lipp et al. 2020). This method works by constructing a model for a centred log-ratio transformed composition (after Aitchison 1986)  $\mathbf{x}'$ , as the sum of a weathering vector,  $\hat{\mathbf{w}}$ , and a protolith vector,  $\hat{\mathbf{p}}$ , relative to the composition of modern UCC:

$$\mathbf{x}' = \mathbf{U}\mathbf{C}\mathbf{C} + \omega\hat{\mathbf{w}} + \psi\hat{\mathbf{p}} + \mathbf{E}.$$
 (1)

The coefficients of these vectors correspond to the weathering intensity experienced by a sediment,  $\omega$ , and its protolith composition,  $\psi$ . Deviations from this model cause the misfit, **E**, to rise. Sediments with protoliths more(/less) felsic than modern UCC have positive(/negative)  $\psi$  values. Weathering of rocks causes  $\omega$  to rise. In this study, we modify this method to correct for the effect of cation exchange (e.g., Sayles and Mangelsdorf 1979), and use a recalibrated  $\hat{\mathbf{w}}$  vector. Both of these modifications improve model fit and reduce the possibility of biases. Other potential biases including diagenesis, are discussed in the Supplementary Information but do not significantly affect our results.

We apply this method to the compilation of sedimentary geochemical data produced by the Sedimentary Geochemistry & Paleoenvironments (SGP) research consortium (sgp.stanford.

Table 1: Average sediment protolith composition (wt %)through time.

Age, Ga	SiO <sub>2</sub>	$Al_2O_3$	$Fe_2O_3$	MgO	Na <sub>2</sub> O	CaO	$K_2O$
0 - 0.5	66.5	14.6	5.32	2.32	3.98	4.32	2.95
0.5 - 1.0	67.3	14.5	4.99	2.05	4.04	3.98	3.17
1.0 - 1.5	67.1	14.5	5.06	2.11	4.03	4.05	3.11
1.5 - 2.0	67.3	14.5	5.00	2.06	4.04	3.99	3.16
2.0 - 2.5	66.1	14.7	5.48	2.46	3.95	4.48	2.85
2.5 +	64.1	14.9	6.22	3.16	3.82	5.27	2.46

edu). The SGP database compiles geochemical data and geological context information from three sources: 1) direct data entry by SGP team members (mainly Neoproterozoic-Paleozoic shales with global geographic coverage), 2) the USGS National Geochemical Database (consisting of data from USGS projects from the 1960-1990s; mainly Phanerozoic samples of all lithologies from the United States), and 3) the USGS Critical Metals in Black Shales database (a global shale database spanning all of Earth history; samples from ore deposits removed). In total we analyse 17,472 major-element compositions each associated with an age. Full details of data, preprocessing and analysis is found in the Supplementary Information. Scripts and data are found at github.com/AlexLipp/crustal-comp.

## THE ARCHEAN PROTOLITH

First, we investigate changes in the average composition of UCC through time. To overcome local heterogeneities we calculate composite samples using the arithmetic mean of all samples in 500 Myr time intervals (Table S1). Because of a low sample density in the oldest part of Earth's history, we average all samples older than 2.5 Ga to create an Archean sediment composite. We solve Equation 1 for each composite to calculate  $\omega$  and  $\psi$ , the weathering and protolith coefficients.

We can use Equation 1 to reconstruct the full major-element composition of a sediment's protolith from just its  $\psi$  value. By substituting the  $\omega$  value of a sediment in Equation 1 for that of pristine igneous rocks ( $\omega_0 = -0.271$ ; see Supplementary Information) the composition of a sediment's protolith is returned. The calculated compositions of the average sediment protoliths through time are shown in Table 1. These protoliths can be analysed as igneous rocks, with, for example, a Total Alkali Silica plot (Figure 1). We find that the average Archean protolith was silica-rich and Dacitic in composition. However, it was slightly more mafic than younger protoliths. This evolved composition for Archean protoliths is similar, albeit marginally more felsic, to the estimates of Ptáček et al. (2020) who used independent methodologies and data. We find that the composition of the average protolith has remained constant since 2.5 Ga.

As most sediments derive from broad regions, their compositions, in aggregate, can be assumed to be representative of the average crustal composition (Rudnick and Gao 2003). Our estimate for the average protolith of recent, < 0.5 Ga, sedimentary rocks (Figure 1) is within error of the estimate of UCC as averaged by surface sampling, validating this approach. Hence, the composition of the Archean sedimentary protolith would suggest that the average Archean UCC was Dacitic. A uniformitarian



Figure 1: A Dacitic composition for Archean UCC. Total Alkali-Silica plot displaying the protoliths of the average sediment for different time periods (Le Maitre et al. 2005). Ellipses indicate 68.3 % (i.e., 1  $\sigma$ ) confidence range (see Supplementary Information). '×' is the pristine igneous precursor of the modern upper continental crust (Rudnick and Gao 2003). Dashed line is trend described by  $\hat{\mathbf{p}}$ . See Supplementary Information for details of how uncertainties are generated.

interpretation of this felsic Archean UCC is that plate tectonics commenced no later than the early Archean.

Nonetheless, we exercise caution about inferring global conditions from the small inventory of Archean samples. The low sample size of Archean sedimentary rocks makes inferences about global conditions highly susceptible to preservation and sampling biases (Korenaga 2013). For example, the sedimentary record will overepresent tectonically active regions. Furthermore, it is plausible that evolved igneous rocks could be generated in the absence of plate tectonics (Reimink et al. 2014).

Focusing exclusively on the average sediment protolith neglects other features in our dataset. If we look at the protolith coefficients,  $\psi$ , for individual samples through time (Figure 2a) we see a large diversity in protoliths throughout Earth's history, including prior to 2.5 Ga. If each individual  $\psi$  represents a catchment averaged protolith, then the diversity of rocks at the Earth's surface has remained largely constant over the entire age range of our dataset. We formally compare the  $\psi$  distributions from before 2.5 Ga against those from 0 - 0.5 Ga in Figure 2b. Whilst the median of the two distributions is different, there is considerable overlap between the two age groups. The high diversity of exposed rocks, and their on-average evolved nature, suggests that during the Archean the continental crust was more similar to the modern crust than it was different. Protolith diversity appears to have remained broadly constant since Archean times, which is additional evidence for long-lived plate tectonics.

#### CRUSTAL WEATHERING ON BILLION YEAR TIMESCALES

Second, we explore how the efficiency of  $CO_2$  drawdown by crustal weathering has evolved through time. The weathering of mafic rocks sequesters more CO<sub>2</sub> than felsic rocks due to their higher concentrations of Ca and Mg (e.g., Dessert et al. 2003). The slightly more mafic Archean UCC could result in continental weathering being more efficient at sequestering atmospheric CO<sub>2</sub> than modern UCC. This increased efficiency could potentially bring the weathering CO<sub>2</sub> sink in balance with mantle outgassing despite a smaller exposed continental area (Bindeman et al. 2018; Johnson and Wing 2020; see also Korenaga et al. 2017). To quantify this increased efficiency, we calculate the chemical depletion fraction for any  $\omega - \psi$  pair, assuming that Al<sub>2</sub>O<sub>3</sub> is immobile (Brimhall and Dietrich 1987; see Supplementary Information). The mass of each element mobilised due to weathering per kg of protolith eroded can then be converted into moles of carbonate-bound CO<sub>2</sub> drawn down assuming the following stoichiometry

$$CaSiO_3 + CO_2 \longrightarrow CaCO_3 + SiO_2$$
,

and that Mg exchanges for Ca at mid-ocean ridges (Holland 1984). This calculation assumes all weathering acidity is donated by carbonic acid and is therefore an upper-bound (Torres et al. 2014). Any  $\omega - \psi$  pair can thus be converted into a maximum amount of CO<sub>2</sub> deposited per kg of weathered protolith (Figure S2a).

Weathering the more mafic Archean protolith sequesters ~ 25 % more CO<sub>2</sub> than the 0 - 0.5 Ga protolith for the same weathering intensity (Figure S2b). We note that since the end of



Figure 2: Archean protoliths were more mafic than the present day but equally diverse. (a) Grey points are protolith coefficients,  $\psi$ , for individual samples. Mean  $\psi \pm \sigma$  for each 0.5 Ga time period given by black circles. Means for > 3 Ga greyed out to emphasise low sample coverage. (b) Box-and-whisker comparison of protolith distributions for samples of age > 2.5 and 0 – 0.5 Ga. Box spans the interquartile range (IQR). Whiskers extend 1.5 × IQR from the upper/lower-quartiles. 200 randomly selected samples shown for each age group.

the Archean, this long term  $CO_2$  drawdown efficiency has not changed considerably, due to constant protolith composition and largely constant weathering intensities (Figure S2a). On timescales greater than 0.5 Gyr therefore, any change that may have occurred in Earth's total weathering flux must have been achieved by changing the absolute amount of erosion, not the weathering intensity. A corollary is that, at this timescale, any secular changes in volcanic  $CO_2$  outgassing must have been compensated for by changes in physical erosion and uplift to keep  $\omega$  constant, assuming that other controls on the geologic carbon cycle are constant.

## CRUSTAL WEATHERING OVER THE PHANEROZOIC

Finally, we explore how global weathering intensity has changed during the Phanerozoic. On 10 - 100 Myr timescales, weathering intensity is believed to respond to global climate state as part of a negative feedback (Walker et al. 1981). Individual sediments only record the weathering intensity of their source regions, but collectively they reflect global shifts in silicate weathering intensity. Previous studies of this global weathering thermostat relied mostly on isotopic proxies (e.g., Pogge von Strandmann et al. 2017), but here we explore the secular changes in Phanerozoic weathering intensity using the SGP dataset of ~ 12,000 Phanerozoic sediments.

Figure 3 shows the weathering coefficient,  $\omega$ , for samples less than 600 Ma in age alongside a smoothed trend. The lowest weathering intensity occurs in the Neogene. Other periods of

low intensity are observed in the late Ordovician/Silurian, late Permian and the Jurassic. Peaks in weathering intensity are found in the Carboniferous, Triassic and Cretaceous. Bootstrap resampling of our data and averaging of data within stratigraphic units indicate that sampling biases do not strongly affect our conclusions (Supplementary Information; Figure S4).

We investigate the relationship between weathering intensity to climate state by comparing our weathering intensity trend to the detrended oxygen isotope composition of marine carbonates, which is considered a proxy for global climate state (Veizer et al. 2000). Generally, when the  $\delta^{18}$ O of marine carbonates is heavy (associated with cooler climates) we observe a lower weathering intensity while the opposite is true for lighter  $\delta^{18}$ O (associated with warmer greenhouse climates). It is argued that the marine carbonate  $\delta^{18}$ O record has been overprinted by diagenetic reactions (e.g., Ryb and Eiler 2018) so we also compare our  $\omega$ record to an independent record of climate state: evidence of glaciated poles. We observe local minima in chemical weathering intensity during ice-house climates (Ordovician-Silurian, Permian, Neogene). We note however that the end-Devonian glaciation coincides with a period of observed high weathering intensity. These observations show a relationship between global weathering intensities and climate state.

The  $\omega$  fluctuations we observe are on timescales too long (10's Myr) to be explained by the silicate weathering feedback acting in response to short-term climatic perturbations. The weathering intensity trend is instead consistent with long-term CO<sub>2</sub> massbalance forced by the solid Earth. An increased flux of volcanic



Figure 3: Weathering intensity of sedimentary rocks across the Phanerozoic. Grey points are individual samples, black line is smoothed trend calculated using 30 Myr bandwidth Gaussian kernel. Grey lines show the bounding envelope of smoothed versions of 1000 bootstrap resamples of SGP dataset (see Supplementary Information). Red line is detrended Oxygen isotope composition of carbonates smoothed using 30 Myr bandwidth Gaussian. Data from  $\delta^{18}$ O compilation of Jaffrés et al. (2007). Periods of glaciation and land-plant expansion given by Macdonald et al. (2019) and McMahon and Davies (2018) respectively. Star emphasises short-lived Gaskiers glaciation (Pu et al. 2016)

 $CO_2$  will result initially in an imbalance in the geologic carbon cycle, as the weathering sink is unchanged. However, as atmospheric  $CO_2$  rises on Myr timescales, the weathering intensity of rocks should rise due to higher pCO2 driving warmer and wetter conditions. The carbon cycle will then remain balanced albeit at an elevated  $CO_2$  level and altered climate state. The opposite response is expected for a reduction in volcanic outgassing. In this way, the weathering carbon sink changes in concert with solid Earth degassing (e.g., Berner and Caldeira 1997; Kump et al. 2000; McKenzie et al. 2016). The observed correlation between weathering intensity and climate state in Figure 3 is consistent with this mass-balance.

The lack of a significant state-change in weathering intensity following the Paleozoic emergence of land-plants further illustrates the importance of carbon mass balance. It is argued that the expansion of land plants, by increased organic acids and root-action, caused a permanent increase in weathering intensity of the land-surface (e.g., Algeo et al. 1995). However, no step-change in  $\omega$  is observed, a result inconsistent with this hypothesis. A step-increase in weathering intensity and thereby carbon drawdown could only maintain balance with volcanic outgassing if there is a concomitant reduction in physical erosion. Given that physical erosion is ultimately controlled by tectonics, this is implausible. Hence, the 10's Myr  $\omega$  fluctuations we observe record weathering intensity changes in response to solid-earth forcing by CO<sub>2</sub> degassing, not in response to internal climate perturbations.

## CONCLUSIONS

A large inventory of sedimentary rock major-element compositions has been deconvolved into a record of crustal composition and weathering intensity. Results indicate an evolved and heterogeneous Archean crust, which suggests an early onset of plate tectonics. Weathering of this Archean crust was ~ 25 % more efficient at sequestering atmospheric CO<sub>2</sub> than modern day UCC. On long, Gyr, timescales the weathering intensity of the crust has remained constant. By contrast, on short, 100 Myr, timescales weathering intensity responds to global climate shifts consistent with a silicate weathering feedback balancing by solid earth CO<sub>2</sub> degassing.

## DATA AND CODE AVAILABILITY

An annotated R-markdown notebook which performs all the calculations described below in R and python can be found at github.com/ AlexLipp/crustal-comp (Van Rossum and Drake 2009; R Core Team 2018). The data used for our reported results and the SQL command used to query the SGP database are also found at the same location.

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# Author Contributions

AGL and OS conceived of the study. AGL performed data analysis and prepared the manuscript. EAS led development of the Sedimentary Geochemistry and Paleoenvironments Project. JJB, DC, PWC, LDM, KD, SQD, JFE, UCF, AJ, BWJ, PK, CBK, MK, AJM, NTM, BO, SEP, NJP, SRR, SDS, PRW and JY all contributed to the Sedimentary Geochemistry and Paleoenvironments Project. JFE guided discussion of the role of diagenesis. All authors contributed to manuscript revision.

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## SUPPLEMENTARY INFORMATION

An annotated R-markdown notebook which performs all the calculations described below in R and python can be found at github.com/AlexLipp/crustal-comp (Van Rossum and Drake 2009; R Core Team 2018). The data used for our reported results and the SQL command used to query the SGP database are also found at the same location.

# Data

The Sedimentary Geochemistry and Paleoenvironments Project (SGP) is a research consortium that has produced the largest available compilation of sedimentary geochemical data from across Earth's history. This compilation is being analysed for a range of different research questions, in addition to this study. The project combines pre-existing large datasets, such as the United States Geological Survey, Critical Metals in Black Shales compilation, with new data gathered from temporal gaps identified in the record. Each sample in the dataset is accompanied with contextual data relating to their stratigraphic and geographic position, including an interpreted absolute age. More information can be found at the project's homepage at sgp.stanford.edu/about. The current phase of SGP focuses primarily on the Phanerozoic and so for the Precambrian aspects of this study it was supplemented with the compilation from Lipp et al. (2020) and further literature data (Fedo et al. 1996; Nesbitt et al. 2009; Devaraju et al. 2010). A histogram of how these samples are distributed in time is given in Figure S1.

From this database, we query for samples which contain measurements of all the seven elements required for our method (Si, Al, Mg, Fe, Na, Ca, K), by any analytical method except for Handheld X-Ray Fluorescence due to lower accuracy. We also query for a range of additional metadata. These elemental compositions are converted into wt% oxides, with total iron given as  $Fe_2O_3$ . The results of our method are the same whether or not the composition is normalised to 100 % prior to analysis.

## Data Screening

To ensure the reliability of our input data, we screen certain samples from the data extracted from the SGP database. First, we remove any manually identified duplicates which are present due to overlap between the literature compiliation of Lipp et al. (2020) and any literature data within the SGP dataset. Second, we exclude any samples for which the lithology is not listed as siliciclastic. Finally, from these siliciclastic sediments we exclude samples which are likely to have been affected by carbonate contamination. How these carbonate contaminated samples are identified is described in further detail below.

## **Composite Samples**

Composite sediment samples are generated by mixing (i.e, taking the arithmetic mean) of all samples within 500 Ma intervals. Due to low sample density in the Archean, all samples older than 2.5 Ga were mixed to generate an Archean composite. If composite samples are created in 500 Ma bins in the Archean, there are some minor changes to the results. The 3.5 - 4.0 Ga interval produces an Andesitic protolith, and the 3.0 - 3.5 Ga has a protolith similar to that from the time period 0 - 0.5 Ga.



2000

1500

a

Supplementary Figure 1: **Temporal sampling density of dataset** (a) Histogram of ages for all samples included in dataset with binwidth equal to 25 Myr. Coloured boxes indicate geological periods (see Figure 3). (b) Same as panel a but for the number of individual stratigraphic units sampled, as defined by the SGP dataset.

# Supplementary Table 1: Composite sediment compositions through time.

Age, Ga	SiO <sub>2</sub>	$Al_2O_3$	$Fe_2O_3^T$	MgO	Na <sub>2</sub> O	CaO	$K_2O$
0 - 0.5	71.7	13.7	5.12	2.35	1.14	2.99	3.06
0.5 - 1.0	70.0	15.6	5.42	2.18	1.26	1.79	3.77
1.0 - 1.5	68.4	16.4	5.97	2.34	1.32	1.53	4.08
1.5 - 2.0	66.6	17.1	6.00	2.51	1.09	2,13	4.58
2.0 - 2.5	66.0	17.6	7.46	2.41	1.86	1.25	3.46
2.5 +	65.1	17.0	7.70	3.29	2.08	1.84	3.00

However, given the low sampling density (the 3.5 - 4.0 Ga interval contains only 30 samples all from the Isua Greenstone Belt), this temporal variance is likely strongly affected by local variability. There is no reasonable binning procedure which results in a protolith of any time-interval more mafic than an Andesite.

# Data Analytical Methods

The method we use was developed and detailed in full in Lipp et al. (2020). This method deconvolves the major-element composition of a sediment into the contribution due to changes in protolith and changes caused by chemical weathering. This method has a number of benefits relative to previous compositional analytical approaches such as: insensitivity to the issues of the 'closure effect', the ability to reconstruct the full composition of protoliths, and a quantitative measure of misfit.

This approach works by constructing a 2D vector addition model to explain the major-element compositions of sediments. This model is shown in Equation 1 in the main manuscript and repeated here:

$$\mathbf{x}' = \mathbf{UCC} + \omega \hat{\mathbf{w}} + \psi \hat{\mathbf{p}} + \mathbf{E}$$

This model is only applied to a composition, **x**, after they have undergone a centred log-ratio, *clr*, transformation to resolve the 'closure' effect inherent to compositional data (Aitchison 1986). Hence,  $clr(\mathbf{x}) = \mathbf{x}'$ . The unit vectors corresponding to weathering and protolith,  $\hat{\mathbf{w}}$  and  $\hat{\mathbf{p}}$  respectively, were calibrated on independent data.  $\hat{\mathbf{w}}$  was calibrated using a soil profile, and  $\hat{\mathbf{p}}$  was calibrated using a suite of cogenetic igneous rocks. The model calibrated in this way was successful in extracting information of weathering intensity and protolith from sedimentary compositions and explained the majority of the observed variance. However, residual analysis indicated two potential points at which the model could be improved.

First,  $\hat{\mathbf{w}}$  was noted to be slightly miscalibrated, causing a systematic increase in misfit with increasing weathering intensity. This miscalibration was likely caused by calibrating  $\hat{\mathbf{w}}$  on a single profile, which will incorporate localised noise. For this study, to derive a better calibrated vector we take a 'consensus'  $\hat{\mathbf{w}}$  of two different profiles. Specifically it is the arithmetic mean of the first principal components of the soil profile reported in White et al. (2001) and the 'Toorongo' soil profile that was used individually to calibrate the original  $\hat{\mathbf{w}}$ . This better calibrated  $\hat{\mathbf{w}}$  ameliorates the issue of rising misfit with greater weathering intensity. All the vectors utilised in Equation 1 are given in Table S2.

Secondly, a relationship between clr(Ca) and clr(Na) residuals was noted. This relationship was interpreted to be related to cation exchange of Ca and Na which is increasingly recognised as playing a significant impact on geochemical cycles (Sayles and Mangelsdorf 1979; Cerling et al. 1989; Lupker et al. 2016). Subsequently it was observed that cation exchange could cause, minor, spurious changes in  $\omega$  and  $\psi$  if it was not explicitly taken into account. As a result, in this study we include a cation exchange correction into our model. Hence, our model is now:

$$\mathbf{x}' = \mathbf{U}\mathbf{C}\mathbf{C} + \omega\hat{\mathbf{w}} + \psi\hat{\mathbf{p}} + f(\chi) + \mathbf{E}$$

This correction factor simply shifts compositions which have been offset from model plane due to cation exchange, back onto the 2D plane indicated in Equation 1. In this formulation  $f(\chi)$ varies the proportion of total Ca and Na which taken up by Ca,  $\chi$ , according to the stoichiometry:

$$Na_{2,clay} + Ca^{2+} \implies 2Na^+ + Ca_{clay}$$

 $\chi$  therefore ranges between 0 and 1.  $f(\chi)$  is non-linear so this equation has no analytical solutions. To find  $\omega$ ,  $\psi$  and  $\chi$  we therefore numerically minimise  $|\mathbf{E}|$  using a gradient descent algorithm implemented in python.

When increasing the components of any model there is always a tradeoff between over- and under-fitting data. We choose to explicitly consider cation exchange as a process as not doing so introduced some minor biases into our results. Nonetheless, this increases the risk that other processes not explicitly included in the model act to alias the results. We found that the major results of this study were invariant to including cation exchange or not.

#### Calculating Protolith Compositions

Consider a *clr* transformed major-element composition  $\mathbf{x}'$ . We solve Equation 1 to calculate its  $\omega$  and  $\psi$  values. These coefficients can be interpreted in terms of translating a protolith composition parallel to the weathering vector a distance equal to ' $\omega$ '. This translation is performed relative to a protolith composition equal to  $\mathbf{UCC} + \psi \hat{\mathbf{p}}$ . Hence, to calculate the protolith composition we simply translate back along the weathering vector to the original  $\omega$  value. As a result, we need to calculate an  $\omega$  value which corresponds to pristine igneous rocks, i.e.,  $\omega_0$ . Previously this was done by calculating the mean  $\omega$  value of a large suite of igneous rocks taken from the NAVDAT (www.navdat.org) database (see Lipp et al. 2020). Performing this calculation on the same compilation of igneous rocks for our updated  $\hat{\mathbf{w}}$  vector gives an  $\omega_0 = -0.271$ .

## Quantifying Uncertainties

Projecting all igneous variability onto a single 1D vector,  $\hat{\mathbf{p}}$ , is obviously a simplification, albeit a useful one. When we calculate protolith compositions as described above, any variability excluded from this 1D vector is neglected. A useful measure of uncertainity in the protolith calculations therefore is how much natural variability there is of real igneous rocks relative to this  $\hat{\mathbf{p}}$  trend. To calculate this variability we solve Equation 1 for the NAVDAT compilation of igneous rocks described above. The variability of igneous rocks around the trend, is hence the misfit matrix  $\mathbf{E}$  for the NAVDAT dataset. To generate the uncertainty distributions shown in Figure 1 we simply add this derived  $\mathbf{E}_{NAVDAT}$  matrix to the calculated protolith compositions. To turn this empirical distribution on the TAS plot into a confidence ellipse we fit a 2D t-student distribution and demarcate the standard error of the mean ellipse, i.e., the 68.3 % interval.

#### Quantifying CO<sub>2</sub> drawdown

If a sediment's protolith composition is known, the relative loss of a specific element due to weathering can be calculated using the chemical depletion fraction, assuming an immobile element (e.g., Jiang and Lee 2019). In this study we calculate the sediment protolith composition,  $x_0$ , from an observed composition,

Supplementary Table 2: Centred log-ratio transformed vectors used to solve Equation 1

	SiO <sub>2</sub>	$Al_2O_3$	$Fe_2O_3^T$	MgO	Na <sub>2</sub> O	CaO	$K_2O$
UCC	2.33	0.869	-0.142	-0.957	-0.681	-0.587	-0.836
ŵ	0.242	0.369	0.235	0.133	-0.487	-0.678	0.186
p	0.234	0.098	-0.231	-0.601	0.248	-0.336	0.589



Supplementary Figure 2: CO<sub>2</sub> drawdown capacity of sedi**ments through time.** (a)  $\omega - \psi$  plot overlain with contours of CO<sub>2</sub> drawdown capacity, calculated using the methods detailed in the main text. This is the amount of CO<sub>2</sub> that could be transferred from the atmosphere to the lithosphere via weathering, assuming all acidity is donated by carbonic acid. Value is given as kg of CO<sub>2</sub> removed from atmosphere per tonne rock eroded. Sediments with more mafic protoliths and greater weathering intensities result in higher potential CO<sub>2</sub> drawdown. Coloured points are the composite sediments from the indicated time periods. Dashed line corresponds to  $\psi$  value of period 0 - 0.5 Ga; Dotted line period 2.5 + Ga. (b) Ratio of Archean to present (0 -0.5 Ga) CO<sub>2</sub> drawdown capacities for different  $\omega$  values. Line is generated by dividing the CO<sub>2</sub> drawdown capacity along the dotted line by the values along the dashed line in Panel a. For the same weathering intensity, the Archean sedimentary protolith sequestered ~ 25 % more  $CO_2$ .

 $\mathbf{x}_1$ , using the method described above and use aluminium as an immobile element. Hence

$$f_i = \frac{\Delta M_i}{M_{i,0}} = \frac{x_{i,1}}{x_{i,0}} \cdot \frac{x_{Al_2O_3,0}}{x_{Al_2O_3,1}} - 1.$$

In this formulation, **f** is the kg of each component lost due to weathering for each kg of initial protolith,  $M_{i,0}$  is the initial mass of the *i*<sup>th</sup> component of a composition and  $\Delta M_i$  is the change in mass of that same component due to weathering. Using this formula and Equation 1 it is possible to calculate the relative loss of each element due to weathering for a given  $\omega - \psi$  pair. Converting from the mass of CaO and MgO lost to moles it is therefore possible to describe the  $\omega - \psi$  plane in terms of maximum kg of CO<sub>2</sub> sequestered per kg protolith eroded (Figure S2a).

## Potential biases and limitations

### **Diagenetic Alteration**

There is increasing evidence that diagenetic reactions between terrestrial sediments and pore-fluids are significant parts of the geochemical cycle of many elements (Sun et al. 2016). As the pH and geochemistry of porefluids evolves with depth, some primary minerals are dissolved, and reprecipitated as authigenic phases, changing the mineralogical composition of sediments as they are lithified. Three lines of evidence suggest our results have not been affected by this process. First, whilst these reactions affect mineralogy, so long as the reactions operate under closed-system conditions, use of bulk major-element data in aggregate is expected to be largely unaffected by this process. Given that the majority of siliciclastic sediments utilised in the SGP database are fine-grained and therefore low porosity, on this basis, closed-system diagenesis is expected (Bjørlykke and Jahren 2012). Assuming dominantly closed-system diagenesis, our results should therefore be insensitive to diagenetic reactions. There is further evidence to support this assertion. Second, we can test the influence of diagenesis on the major element record of marine fine grained sediments by comparing recent sedimentary rocks (0 - 0.5 Ga) to the major-element composition of the modern UCC as determined by surface sampling. This comparison shows that fine grained sediments produce a good match to the independently determined UCC composition. This suggests that diagenetic reactions have not biased the compositions of these recent sedimentary rocks. Finally, all of our principal findings are derived from aggregates of sediment geochemistry, either by generating composite samples (Table S1), or by calculated a smoothed trend of noisy data (Figure 3). Hence, any diagenetic transfer of material within the sedimentary column will likely be undone to some degree during our data analytical process.

We emphasise here that our model is unable to indicate where or when a particular compositional process may have acted. For example, open system chemical weathering which acts in the subsurface (e.g. anoxic marine silicate weathering; Wallmann et al. 2008) could have the same compositional effect as chemical weathering which acts in a sediment source region. By extension, recycling of ancient sedimentary rocks could also impart an inherited weathering signal into any subsequent sediments. The lack of a secular increase in weathering intensity over Earth's history suggests that this inheritance effect is not significant however.

Similarly, we have only explicitly considered cation exchange as acting on cations absorbed to clay particles. However, any reaction, diagenetic or otherwise, which is a net charge-balanced exchange of Ca for Na would produce the same compositional trend. This ambiguity is a limitation of any approach using purely elemental data.

# Carbonate addition bias

Our model only explicitly considers the siliciclastic portion of a sediment. However, many sediments contain some portion of authigenic or biogenic carbonates, most commonly as calcite. As a result, it is important to understand the impact that calcite addition has on the interpretation of the  $\omega$  and  $\psi$  coefficients. To investigate this effect we performed a synthetic experiment by increasing the amount of CaO in a sediment composition, and recalculated the  $\omega$  and  $\psi$  coefficients. The results of this experiment are shown in Figure S3. Addition of calcite introduces a spurious reduction in weathering intensity. A spuriously more felsic protolith is also introduced, although the magnitude of this effect is much more minor. It is important to minimise the effect of these biases by screening samples from our dataset which are clearly affected by carbonate addition. Lacking mineralogical data it is generally challenging to identify carbonate contaminated samples. One approach is simply to remove samples which have CaO values above a certain cut-off value. However, given that cation exchange is another process which can increase the CaO this approach may be removing sediments which do not contain significant carbonate but simply have absorbed Ca. As a result, we identify the maximum amount of calcium which could be expected to be found in a sediment, if all of the sodium exchanges for calcium, for a standard range of protoliths. This corresponds to a sediment which derives from a basalt but has not undergone significant weathering. Hence, any sediment which has more calcium than this cut off value, if all the sodium is also exchanged, must also contain calcite and is thus excluded from consideration. This procedure is detailed more explicitly in the accompanying code (github.com/AlexLipp/crustal-comp).

Despite this filter, it is still likely that small amounts of calcite are present in many sediments from our dataset. As a result, our interpreted weathering intensities are likely an underestimate. To make sure that this bias is not the cause of the temporal trends in weathering intensity we see, we repeated our analysis with a more stringent carbonate filter (excluding all samples which had a positive CaO residual when cation exchange is not considered), but we observed similar trends. This suggests that whilst our data does likely incorporate the effect of carbonate addition, it does not affect the conclusions we have drawn.



Supplementary Figure 3: **Exploring biases due to calcite addition** (a) Changes in  $\omega$  coefficient resulting when increasing amounts of calcite are added to a composition with initial ( $\omega$ ,  $\psi$ ) values of (2,-1). This suggests that the presence of any carbonate results in a spurious reduction in weathering intensity. (b) Changes in  $\psi$  coefficient resulting from same synthetic experiment as panel a. Calcite addition therefore introduces a spurious felsic bias. Note however that this bias is much smaller in magnitude than the bias introduced for  $\omega$ .

# Sampling Bias

The samples in our dataset are not homogenously distributed in time (Figure S1). As a result, for periods of low sample density, the average weathering trend we calculate will be more uncertain. To quantify this changing uncertainty interval we utilised a bootstrap resampling technique. Here, we resampled our dataset 1000 times with replacement. For each of these resampled datasets we generated a smoothed trend with the same 30 Myr Gaussian kernel. Then for each point in time we simply calculate the minimum and maximum possible values of the smoothed trend from these 1000 possible trends. These envelopes are shown as grey lines in Figure 3. This bounding envelope is wider in periods of lower sampling density reflecting the greater uncertainty of the average trend (Figure S1a).

An additional issue that can arise is if some particular stratigraphic units are 'oversampled' due to various reasons including accessibility of samples and particular economic interest. To investigate the robustness of our findings in the Phanerozoic against this issue we generate an analogous figure to Figure 3 but we instead take the mean  $\omega$  value for individual stratigraphic units. We subsequently smooth these unit-averaged data points in the same way as before (i.e., applying a Gaussian filter of 30 Myr bandwidth and calculating the bootstrap uncertainties). The results of this analysis are displayed in Figure S4. The location of the the peaks and troughs in this trend are largely similar to the original trend in Figure 3. The trough in weathering intensity during the Ordovician/Silurian in Figure 3 is not however visible in this unit-averaged trend. It is notable that due to the smaller number of data points the bootstrap uncertainties are much higher than if samples are treated individually. This is particularly true for regions where only a small number of stratigraphic units have been sampled (Figure S1b).



Supplementary Figure 4: Weathering trend after averaging of  $\omega$  values within stratigraphic units. Grey points indicate average  $\omega$  value of individual stratigraphic units. Black curve generated by smoothing data using 30 Myr Gaussian kernel. Grey lines indicate bounding envelope of 1000 bootstrap resamples of data. The general shape of this trend is similar to that displayed in Figure 3 although without a reduction in weathering intensity during the Ordovician/Silurian. Note the much greater uncertainty envelope than if all samples are treated individually.