

# Modelling the Impact of Palaeogeographical Changes on Weathering and CO<sub>2</sub> during the Cretaceous-Eocene Period

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**Abstract.** The feedback between atmospheric CO<sub>2</sub> concentrations and silicate weathering is one of the key controls on the long term climate of the Earth. The potential silicate weathering flux (as a function of conditions such as temperature, runoff, and lithology), or "weatherability", is strongly affected by continental configuration, and thus the position of continental landmasses can have substantial impacts on CO<sub>2</sub> drawdown rates. Here, we investigate the potential impact of palaeogeographical changes on steady-state CO<sub>2</sub> concentrations during the Cretaceous-Eocene period (145-34 Ma) using a coupled global climate and biogeochemical model, GEOCLIM, with higher resolution climate inputs from the HadCM3L General Circulation Model (GCM).

We find that palaeogeographical changes strongly impact CO<sub>2</sub> concentrations by determining the area of landmasses in humid zones and affecting the transport of moisture, that runoff is a strong control on weatherability, and that changes in weatherability could explain long term trends in CO<sub>2</sub> concentrations. As Pangaea broke up, evaporation from the ocean increased and improved moisture transport to the continental interiors, increasing runoff rates and weathering fluxes, resulting in lower steady-state CO<sub>2</sub> concentrations. Into the Cenozoic however, global weatherability appears to switch regimes. In the Cenozoic, weatherability appears to be determined by increases in tropical land area, allowing for greater weathering in the tropics.

Our modelled CO<sub>2</sub> concentrations show some strong similarities with estimates derived from proxy sources. Crucially, we find that even relatively localised changes in weatherability can have global impacts, highlighting the importance of so-called weathering "hot-spots" for global climate. Our work also highlights the importance of a relatively high-resolution and complexity forcing GCM in order to capture these hot-spots.

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## 1 Introduction

20 One of the key controls on the long-term climate of the Earth is the concentration of atmospheric CO<sub>2</sub>. Throughout Earth's history, concentrations of CO<sub>2</sub> in the atmosphere have changed substantially, for example, from several thousand ppm in the early Phanerozoic to under 200 ppm during the last glacial maximum (e.g. Royer, 2006; Bouttes et al., 2011; Foster et al., 2017). Atmospheric CO<sub>2</sub> concentrations are ultimately determined by the balance of CO<sub>2</sub> emitted from volcanic sources (volcanic degassing), and the drawdown of CO<sub>2</sub> by the chemical weathering of silicate rocks (Bernier et al., 1983). This "simple" balance  
25 is complicated by the complexity of chemical weathering processes, especially in the geological past. It should be noted that the burial of organic matter is also significant for atmospheric CO<sub>2</sub> concentrations (e.g. Bernier, 1990; Hilton and West, 2020). The impact and extent of this process can be difficult to quantify in the geological past and is not assessed as part of this study.

Chemical weathering rates of silicate minerals are sensitive to climate variables such as temperature and precipitation in addition to intrinsic factors (i.e. factors inherent to the rocks themselves, such as lithology) (Strakhov, 1967; Walker et al.,  
30 1981; Bernier et al., 1983; White and Blum, 1995; Oliva et al., 2003; West et al., 2005). While there is evidence from individual sites that weathering is accelerated under warm, humid conditions (e.g. White and Blum, 1995), it is unclear whether such a trend can be applied at the global scale. Evidence from field and modelling studies of weathering profiles suggests a stronger precipitation or runoff-based control on weathering rates than temperature (e.g. Oliva et al., 2003; Maher, 2010; Hayes et al., 2020), while global models (e.g. GEOCARB and derivatives) have often invoked temperature as a stronger control (Bernier  
35 et al., 1983; Bernier and Kothavala, 2001). Other global models, including GEOCLIM, have suggested that runoff may be a more significant control on weathering rates and thus long-term climate (Godd ris et al., 2014). The geographical distribution of lithology may play a role (e.g. Donnadi u et al., 2004), but palaeolithologies are difficult to constrain.

General Circulation Models (GCMs) have been used extensively to model climates on multi-million year timescales (e.g. Lunt et al., 2021). These models have been evaluated using records from proxies such as stable isotope ratios, TEX86, Mg/  
40 Ca ratios, and alkenones (e.g. Hutchinson et al., 2021). Although GCM simulations do not always produce results that agree with proxy data (e.g. Huber and Caballero, 2011; Keating-Bitonti et al., 2011; Lunt et al., 2012; Jagniecki et al., 2015), they have greatly expanded the spatial knowledge of past climates, whereas proxy data have been gathered from a comparatively limited number of sites (e.g. Huber and Caballero, 2011). Furthermore, proxy data often produce conflicting reconstructions, such as differing atmospheric CO<sub>2</sub> levels (e.g. Jagniecki et al., 2015) or significantly different temperature and precipitation  
45 reconstructions (e.g. Keating-Bitonti et al., 2011). GCMs can be used to reconstruct global climates based on different proxy records and compare the results (e.g. Huber and Caballero, 2011; Lunt et al., 2012, 2016). Thus, GCMs have become a powerful tool in reconstructing palaeoclimates.

The large quantities of GCM-derived climate outputs have been incorporated into global geochemical models over the last 10-15 years (e.g. Bernier and Kothavala, 2001; Donnadi u et al., 2004), enabling some of those models to provide spatially  
50 varying estimates of global weathering rates under different climate and palaeogeographic configurations. However, as noted above, these climate reconstructions can vary significantly depending on the model configuration and the boundary conditions used.

## 1.1 Controls on Mesozoic-Cenozoic Climate

Over multi-million year timescales, climates during the Mesozoic and Cenozoic were affected by a gradual increase in solar forcing and, in general, a gradual decrease in CO<sub>2</sub> forcing (Foster et al., 2017, ; Figure 1). A number of theories have been advanced to explain changes in atmospheric CO<sub>2</sub>, and thus climate, over multi-million timescales. The gradual breakup of Pangaea and associated rifting likely resulted in increased volcanic degassing fluxes (Tajika, 1998; Lee and Lackey, 2015; Lee et al., 2015). The late Cretaceous and the Cenozoic saw a period of mountain building associated with the latter stages of the break-up of Pangaea. The Laramide orogeny began approximately 70 Ma (Humphreys et al., 2003) and the development of the Northern Andes and Himalaya occurred during the early Cenozoic (Schellart, 2008). The development of new mountain ranges, particularly Himalaya, has been implicated in the reduction of CO<sub>2</sub> concentrations during the Cenozoic by increasing silicate weathering rates and thus increasing CO<sub>2</sub> drawdown (Raymo and Ruddiman, 1992). Changes in weathering fluxes have been linked to a number of climate shifts in the Earth's past, such as the global cooling which occurred during the Carboniferous (e.g. Godd ris et al., 2014, 2017).

Changes in palaeogeography can have substantial effects (Lunt et al., 2016). For example, the presence of mountain ranges may affect monsoon circulation and thus precipitation, or the presence of large continental interiors isolated from moisture can lead to large arid deserts. As such, the potential silicate weathering flux, or 'weatherability' of the continents is likely to represent a key climate forcing during the late Mesozoic-early Cenozoic and forms the focus of this study.

At steady-state, silicate weathering fluxes are equal to emissions from volcanic degassing (e.g. Walker et al., 1981; Berner et al., 1983; Godd ris et al., 2014). However, a distinction should be drawn between weathering (here, the chemical breakdown of silicate rocks by water and carbonic acid) and "weatherability". Weatherability corresponds to the susceptibility of the land surfaces to be weathered, under a given climate (temperature and runoff) (Kump and Arthur, 1997). Weatherability can be affected by the presence of land plants, or relief, erosion rates (through the exposure of fresh material to weathering), or by changes in the lithology, or by the palaeogeographic configuration (Dessert et al., 2003; Oliva et al., 2003; West et al., 2005; Maher, 2010; West, 2012; Bazilevskaya et al., 2013; Bufe et al., 2024). The role of land plants is difficult to constrain and studies have produced contrasting results (e.g. Porada et al., 2016; Godd ris et al., 2017), so vegetation is often excluded or held constant within simulations of weathering.

Global palaeolithologies are poorly constrained and become even less so further back into the geological record. Similarly, factors such as topography and uplift rates, which affect erosion rates (e.g. Riebe et al., 2004; West et al., 2005; Gabet and Mudd, 2009), are also poorly constrained into geological time. In contrast, broad scale climate parameters like temperature and precipitation patterns are comparatively well constrained through proxy data and GCM studies. Climate states are in turn strongly controlled by the positioning of the continents, which can affect oceanic and atmospheric circulation patterns (e.g. Gyllenhaal et al., 1991; Von der Heydt and Dijkstra, 2006). The positioning of the continents also affects weatherability by determining the land area in intense weathering environments, such as the tropics (e.g. Godd ris et al., 2014).

85 In this study we focus on the impact of the palaeogeographic setting on the weatherability, assuming that all the other factors remain constant over the simulated period (this is certainly not the case, but our objective is to quantify exclusively the role of palaeogeography).

For an idealised case where degassing is constant (Walker et al., 1981), the global weathering flux will equilibrate the volcanic degassing of CO<sub>2</sub> and the global CO<sub>2</sub> consumption by silicate rock weathering will stay constant. But the evolving  
90 continental configuration modulates weatherability, allowing CO<sub>2</sub> to fluctuate (Goddéris et al., 2014).

In contrast to palaeo degassing rates, palaeogeographies (within the last 150 million years) are relatively well constrained. Thus, there is scope to investigate the impact of changing weatherability on long-term CO<sub>2</sub> concentrations. Early studies using zero-dimensional global models such as GEOCARB used estimations of global mean annual temperature (MAT) and runoff values, but no geographical constraints, to estimate global weathering fluxes and steady-state CO<sub>2</sub> concentrations in geological  
95 time (Berner, 1991). Further studies refined the GEOCARB approach by including palaeogeographical reconstructions, both conceptual (e.g. ‘ringworld’ configurations) and realistic, which demonstrated that continental configurations had significant impact on steady-state CO<sub>2</sub> concentrations (e.g. Barron et al., 1989) and that palaeogeography and climate interact to determine steady-state CO<sub>2</sub> concentrations (Otto-Bliesner, 1995). Further model developments provided spatial patterns in climate forcing data, demonstrating that regional changes in weatherability can have global climate impacts (Donnadieu et al., 2004). More  
100 recently, such models have shown that the positioning of the continents also affects weatherability by determining the land area in intense weathering environments, such as the tropics (e.g. Goddéris et al., 2014).

The impact of changing weatherability as a result of changes in palaeogeography during the Phanerozoic has previously been investigated by Goddéris et al. (2014) using the GEOCLIM model with climate inputs from the FOAM GCM, which found that the formation of supercontinents resulted in a continental configuration favourable for high atmospheric CO<sub>2</sub> concentrations  
105 (10-25 x PAL), as the converged continental configuration resulted in arid continental interiors and reduced weatherability. Conversely, dispersed continental configurations were more favourable to lower atmospheric CO<sub>2</sub> concentrations (1-8 x PAL) by favouring higher continental runoff and weatherability. Furthermore, Goddéris et al. (2014) found that continents or even smaller landmasses crossing warm-humid climate belts can result in significant draw down of atmospheric CO<sub>2</sub>. For example, during the late-Triassic, the gradual northward drift of Pangaea brought larger areas of landmass into more humid zones. In  
110 response, CO<sub>2</sub> concentrations fell sharply over ~20 Myr from 19 x PAL to 3 x PAL (Goddéris et al., 2014).

Goddéris et al. (2014) had a relatively low temporal resolution of 22 simulations over the last ~520 Myr (approximately 1 simulation per 20-30 Myr) to produce a simulated CO<sub>2</sub> record, and from that glean insights on the effects of changing weatherability (through palaeogeographical changes) on long-term CO<sub>2</sub> concentrations and thus global climate. In contrast, this study will use 19 simulations over the Cretaceous-Eocene period (145-36 Ma) in addition to better constrained and higher  
115 resolution (factor of 3.65 increase) palaeogeography and climate inputs to produce our own simulated CO<sub>2</sub> record, assuming a constant degassing rate. The higher temporal resolution will allow for the investigation of the impacts of higher temporal resolution changes (~5 Myr) in palaeogeography on long-term CO<sub>2</sub> and weatherability, while the higher spatial resolution will better constrain regional variability in weatherability relative to Goddéris et al. (2014). The use of climate inputs produced by

the more complex HadCM3L model in this study (relative to the FOAM inputs used by Godd ris et al. (2014)) also represents  
120 an improvement. These model-based improvements should provide a more accurate picture of how global climate responds to  
changes in weatherability, and better constrain the role of chemical weathering feedbacks on global climate through geological  
time.

## 2 Methodology

This study will investigate the impact of varying palaeogeography from the Early Cretaceous to the Late Eocene on weather-  
125 ability, using a coupled global climate and geochemical model, GEOCLIM, with climatic reconstructions derived from the  
HadCM3L model.

### 2.1 GEOCLIM

GEOCLIM is a coupled global climate and biogeochemical model initially developed in the early 2000's and has been used  
to investigate interactions between climate and geochemistry in deep time settings on geological timescales (Donnadieu et al.,  
130 2004). GEOCLIM uses temperature and runoff inputs from climate models at specified CO<sub>2</sub> concentrations (e.g. 280, 560 ppm)  
and models the climate conditions through interpolation between as atmospheric CO<sub>2</sub> concentrations evolve within GEOCLIM  
simulations. Should CO<sub>2</sub> concentrations in GEOCLIM fall outside of the specified range, it will be unable to update climate  
conditions and thus will use climate conditions associated with the nearest available CO<sub>2</sub> concentration. GEOCLIM initially  
used CLIMBER inputs in early studies (Donnadieu et al., 2004), but more recent studies have used FOAM inputs (Lefebvre  
135 et al., 2013; Godd ris et al., 2014, 2017).

The COMBINE model within GEOCLIM handles biogeochemical processes for both land and ocean environments. Oceans  
are divided into 9 "boxes": 2 high latitude oceans (>60 N/S) (each separated into a photic layer and a deep ocean layer), a  
low-mid latitude ocean (60 S - 60  N) divided into photic, thermocline, and deep ocean layers, and an epicontinental sea also  
divided into photic and deep layers. A final 10th box represents the atmosphere. A range of biogeochemical processes are  
140 modelled within the ocean layers. As this study focuses on terrestrial processes we will omit a detailed description of the ocean  
processes, but full descriptions can be found in Godd ris and Joachimski (2004) and Donnadieu et al. (2006).

Chemical weathering in GEOCLIM occurs in terrestrial grid cells, and the total sum value of silicate weathering from each  
terrestrial grid cell, along with a prescribed volcanic degassing value is used to calculate changes in atmospheric CO<sub>2</sub> at each  
timestep. Ocean carbon and alkalinity are kept balanced at steady-state. Chemical weathering calculations for silicate rocks are  
145 based on equations from Oliva et al. (2003) and Dessert et al. (2003).

$$F_{sil}(t) = k_{sil} * (\alpha_j(t) * \rho_j(t) * A e^{-E_a/RT}) \quad (1)$$

Equation 1 calculates the granitic weathering flux in moles of C per year at a timestep ( $t$ ) in a given grid cell,  $F_{sil}(t)$ ,  $k_{sil}$   
is the silicate weathering constant,  $\alpha_j(t)$  is the area of a grid cell ( $10^6$  km<sup>2</sup>),  $\rho_j(t)$  is the runoff value of that grid cell (cm

yr<sup>-1</sup>). The final term (dimensionless) is an Arrhenius equation for granite weathering, representing the granitic dependence on  
150 dissolution based on air temperature(T) and the universal gas constant (R), which uses the activation energy for granite (E<sub>a</sub>)  
defined by Oliva et al. (2003) and the air temperature in a given grid cell, T<sub>j</sub>.

$$F_{bas}(t) = k_{bas} * (\alpha_j(t) * \rho_j(t) * Ae^{-E_a/RT}) \quad (2)$$

A similar calculation exists for basalt (Equation 2) based on values in Dessert et al. (2003), and for carbonate rocks. Both  
calculations were derived from and calibrated on weathering reaction processes within granite and basaltic watersheds, de-  
155 scribed in detail in Oliva et al. (2003) and Dessert et al. (2003), respectively. We omit the carbonate equation here as this study  
focuses on variations in CO<sub>2</sub> drawdown via changes in silicate weathering fluxes. GEOCLIM assumes that each grid cell has  
an equal area of granitic, basaltic, and carbonate rocks. While this is obviously a significant simplification of the real-world ge-  
ology of the time, palaeolithologies are poorly constrained. In the absence of reliable global palaeolithologies from the studied  
time period, we assume an even distribution of lithologies for simplicity and consistency with Godd ris et al. (2014). A newer  
160 version of GEOCLIM, recently developed, includes an erosion dependency on chemical weathering rates based on equations in  
Gabet and Mudd (2009) (Maffre et al., 2021). As a key part of this study relates to comparing our new simulations to previous  
results with the GEOCLIM model (Godd ris et al., 2014) this version of the model was not used in this study for reasons of  
consistency. The most recent version of GEOCLIM, GEOCLIM7, is fully detailed in Maffre et al. (2024).

## 2.2 HadCM3L

165 HadCM3L is part of the HadCM3 “family” of coupled GCMs originally developed by the UK Met Office (Gordon et al.,  
2000; Valdes et al., 2017). The HadCM3 family of models have been in use for over 15 years, with various modifications  
to the configuration of the original HadCM3 model being made in that time. HadCM3L is a version of HadCM3 using a  
“low-resolution” ocean, where ocean and atmosphere share the same 96x73 (3.75° x 2.5°) resolution (Valdes et al., 2017).  
Although described as “low-resolution”, 96x73 is nonetheless 3.65 times higher resolution than FOAM. However, like FOAM,  
170 HadCM3L is used for long term simulations where higher resolution models would be too time intensive to be practical. To that  
end, HadCM3L has been used extensively in pre-queternary climate studies, where long term climate simulations are required  
(e.g. Lunt et al., 2010, 2011, 2012; Valdes et al., 2017). This study uses climate inputs from 19 HadCM3L simulations of the  
Cretaceous-Eocene period at intervals of 3-13 Ma, the same simulations used by Farnsworth et al. (2019). These simulations  
were run with CO<sub>2</sub> concentrations at 560 and 1120 ppm, which are considered to be a reasonable representation of potential  
175 CO<sub>2</sub> concentrations during the Cretaceous-Eocene, which are subject to considerable uncertainty (Lunt et al., 2016). To match  
the range of climate conditions at the 11 concentrations used for the FOAM climate inputs (160-1400ppm) in Godd ris et al.  
(2014), the HadCM3L climate inputs were extrapolated from the two CO<sub>2</sub> concentrations (560-1120ppm) to the same 11  
concentrations (Appendix A). This process was devised and evaluated using a sensitivity study in Hayes (2019) and found to  
be robust, although this is largely irrelevant as the modelled CO<sub>2</sub> concentrations produced remained very close to or within the  
180 560-1120ppm range. The simulations in Farnsworth et al. (2019) (used for this study) differ from those in Lunt et al. (2016)

slightly in that they have a spin up time of 10,422 years, as opposed to 1422 years in Lunt et al. (2016). The extended spin up time was to enable the simulations to more closely approach an equilibrium state and to make use of the fully dynamic mode within the coupled vegetation model. Runoff calculations are affected by the vegetation model, with both canopy interactions and soil infiltration considered within HadCM3L. A full description of the runoff parameterisation can be found in Gregory et al. (1994). Estimating uncertainties in modelled runoff is challenging, but the higher resolution of HadCM3L relative to FOAM is likely to provide a better parameterisation of features significant to runoff, such as relief, even when accounting for uncertainties in reconstructing palaeo-relief.

The palaeogeographic reconstructions used in this study were developed by Getech for Lunt et al. (2016). The reconstructions were produced using methods developed by Markwick and Valdes (2004), and include data from well-constrained geological databases such as The Paleogeographic Atlas Project. These palaeogeographies were originally constructed at  $0.5^\circ \times 0.5^\circ$  resolution - from these high resolution palaeogeographies, topographies, bathymetries, and land-sea masks were generated at  $3.75^\circ \times 2.5^\circ$  resolution for the HadCM3L model (Lunt et al., 2016). Additionally, HadCM3L is coupled to the TRIFFID model (Top-down Representation of Interactive Foliage and Flora Including Dynamics), which is a dynamic global vegetation model (Cox et al., 2002). The two modules are coupled via the MOSES 2.1 land surface scheme (Cox et al., 1999). The TRIFFID model calculates the fraction of five plant functional types (broadleaf trees, needleleaf trees,  $C_3$  grasses,  $C_4$  grasses, and shrubs) in each grid cell. Although these are modern plant types, studies have suggested that models such as TRIFFID can nonetheless perform acceptably in providing vegetation feedback signals back to at least 250 million years (Donnadieu et al., 2009; Zhou et al., 2012). Estimating uncertainties introduced by the vegetation model is challenging, however Donnadieu et al. (2009) indicates that vegetation models adequately reproduce the spatial distributions of major biogeographical areas, such as desert, temperate, and polar regions.

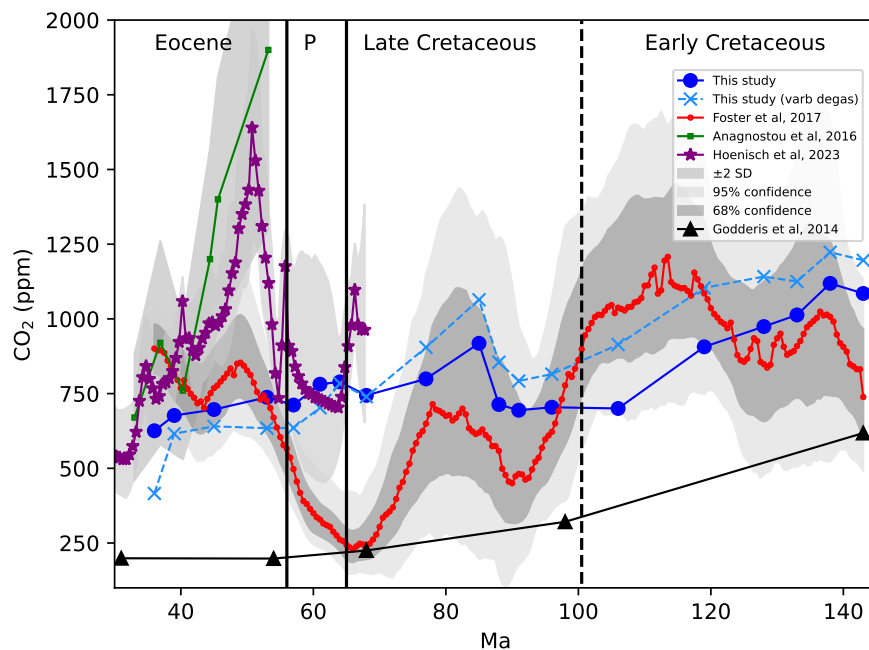
### 2.3 GEOCLIM Configuration

The degassing rate for the paleogeography simulations was set to a constant value of  $1 \times 10^{13}$  mol C yr<sup>-1</sup>, similar to previous GEOCLIM studies (Lefebvre et al., 2013). A value of  $1 \times 10^{13}$  mol C yr<sup>-1</sup> is within the range of estimates (~1.2 – 2 times modern) for degassing rates during the Cretaceous-Eocene period (Van Der Meer et al., 2014). These simulations were also run using variable degassing rates ranging from a low value of  $7.82 \times 10^{12}$  mol C yr<sup>-1</sup> in the latest-Eocene to a high of  $1.23 \times 10^{13}$  mol C yr<sup>-1</sup> in the early-Cretaceous (Appendix Figure A1). This range is based on values calculated by Van Der Meer et al. (2014) using reconstructed subduction zone arc lengths.

We conducted two sets of experiments. We run one simulation per time slice, fixing the initial atmospheric CO<sub>2</sub> to 2.85 times the pre-industrial value of 280 ppm (~ 800 ppm). This CO<sub>2</sub> concentration is intermediate between the 560 and 1120ppm reconstructions. The first set of experiments allows us to isolate the effect of the palaeogeography on the weatherability of the continental surfaces, all other factors being fixed. The second set of experiments was conducted by allowing GEOLCIM to run until atmospheric CO<sub>2</sub> reaches steady-state to provide an estimate of atmospheric CO<sub>2</sub> under the aforementioned palaeo-

geophysical and climate conditions. Atmospheric CO<sub>2</sub> will adapt in response to weathering until global silicate weathering fluxes compensate for solid Earth degassing.

## 215 3 Results

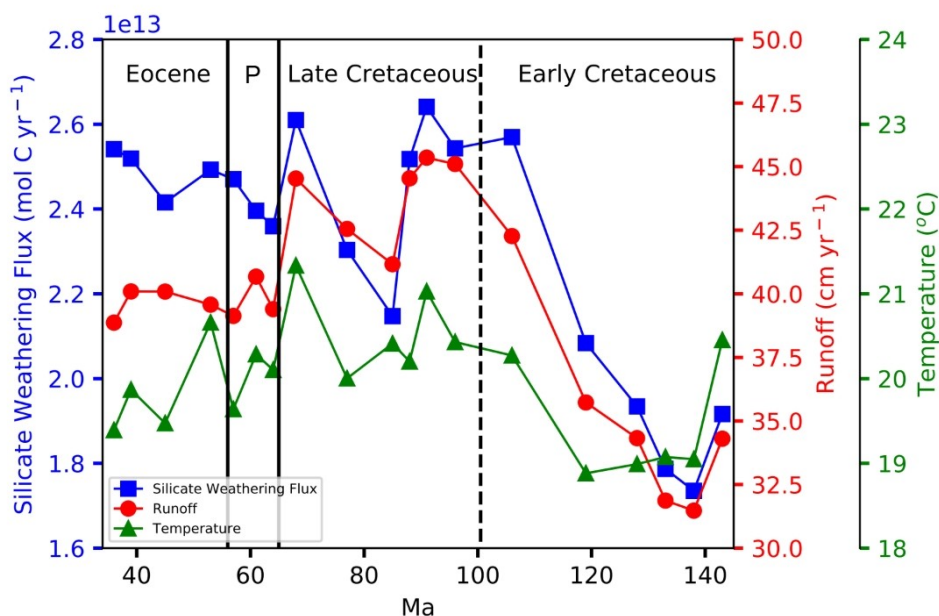


**Figure 1.** Steady-state CO<sub>2</sub> as modelled by GEOCLIM plotted against three CO<sub>2</sub> proxy records from the literature and simulated record from Godd ris et al. (2014). The GEOCLIM model output displays some agreement with the record from Foster et al. (2017), particularly during the early-Cretaceous where CO<sub>2</sub> concentrations are comparably high, the late-Cretaceous, and during the late-Eocene. The CO<sub>2</sub> record from Consortium\*† et al. (2023) (referred to here as Hoenisch et al (2023)) is generally higher than the GEOCLIM model output, but shows a similar pattern in the late Cretaceous and early Palaeocene. The record from Anagnostou et al. (2016) displays considerably higher CO<sub>2</sub> concentrations during the early-Eocene than those in the GEOCLIM and Foster records, but shows better agreement with both records towards the mid and late Eocene. A second GEOCLIM model output (dashed red) displays the results of running GEOCLIM simulations using variable degassing rates calculated in Van Der Meer et al. (2014). Note: Palaeocene is abbreviated to P for brevity.

Our simulations for the 19 time slices produced a simulated CO<sub>2</sub> record from 145-36 Ma. Throughout the results and discussion, patterns within the record will be compared with changes in global weathering fluxes and climate variables to investigate if, and how, changes in weatherability have affected the CO<sub>2</sub> record. To begin, we discuss the role of degassing and silicate weathering fluxes in driving these variations.

220 The GEOCLIM simulations of the 19 time slices resulted in a range of CO<sub>2</sub> concentrations, with the lowest value of ~650 ppm at the end of the Eocene and the highest value of ~1100 ppm during the early Cretaceous (Figure 1). Atmospheric CO<sub>2</sub> exceeds 1000 ppm during the early Cretaceous but declines significantly to ~700 ppm by the mid-Cretaceous. There is a brief increase in CO<sub>2</sub> around 85 Ma to ~900 ppm, followed by a gradual decrease into the Cenozoic and through to the end of the Eocene. The simulations run with variable degassing rates produced a similar pattern, albeit with higher atmospheric CO<sub>2</sub> concentrations during the Early Cretaceous and lower concentrations in the Late Eocene relative to the simulations using a constant degassing rate.

225 concentrations during the Early Cretaceous and lower concentrations in the Late Eocene relative to the simulations using a constant degassing rate.



**Figure 2.** Modelled silicate weathering fluxes (blue squares) plotted against global mean annual runoff (red circles) and temperature (green triangles). Trends in silicate weathering fluxes are similar to those of global runoff, but show less similarity to temperature. Note that Palaeocene has been abbreviated to P.

For the simulations performed with a fixed initial atmospheric CO<sub>2</sub> level at 2.85 times the pre-industrial value, the calculated global silicate weathering flux is inversely related to the atmospheric CO<sub>2</sub> concentrations produced by the simulations where atmospheric CO<sub>2</sub> was allowed to reach steady state (i.e. high initial silicate weathering fluxes result in lower steady state CO<sub>2</sub> concentrations) (Figure 2). The early Cretaceous is marked by low silicate weathering fluxes, although fluxes increase significantly towards the mid Cretaceous and peak at approximately 91 Myr. A brief, but sharp decrease occurs until 85 Myr, followed by a similarly rapid recovery at the end Cretaceous to a similar level of weathering seen at 91 Myr. Another brief decrease occurs into the Palaeocene, followed by a gradual rise through the Eocene. Initial global mean temperature varies

230

little throughout the Cretaceous-Eocene period, although a period of warming occurs in the mid-late Cretaceous, followed  
235 by gradual cooling into the Cenozoic. Farnsworth et al. (2019) noted in their simulations that temperature varied little with  
palaeogeography. Initial mean global runoff increases significantly into the mid-Cretaceous, followed by a brief decrease  
around 85 Ma, and then an increase again towards the end-Cretaceous. Runoff drops sharply at the start of the Palaeocene and  
remains mostly stable through the Eocene.

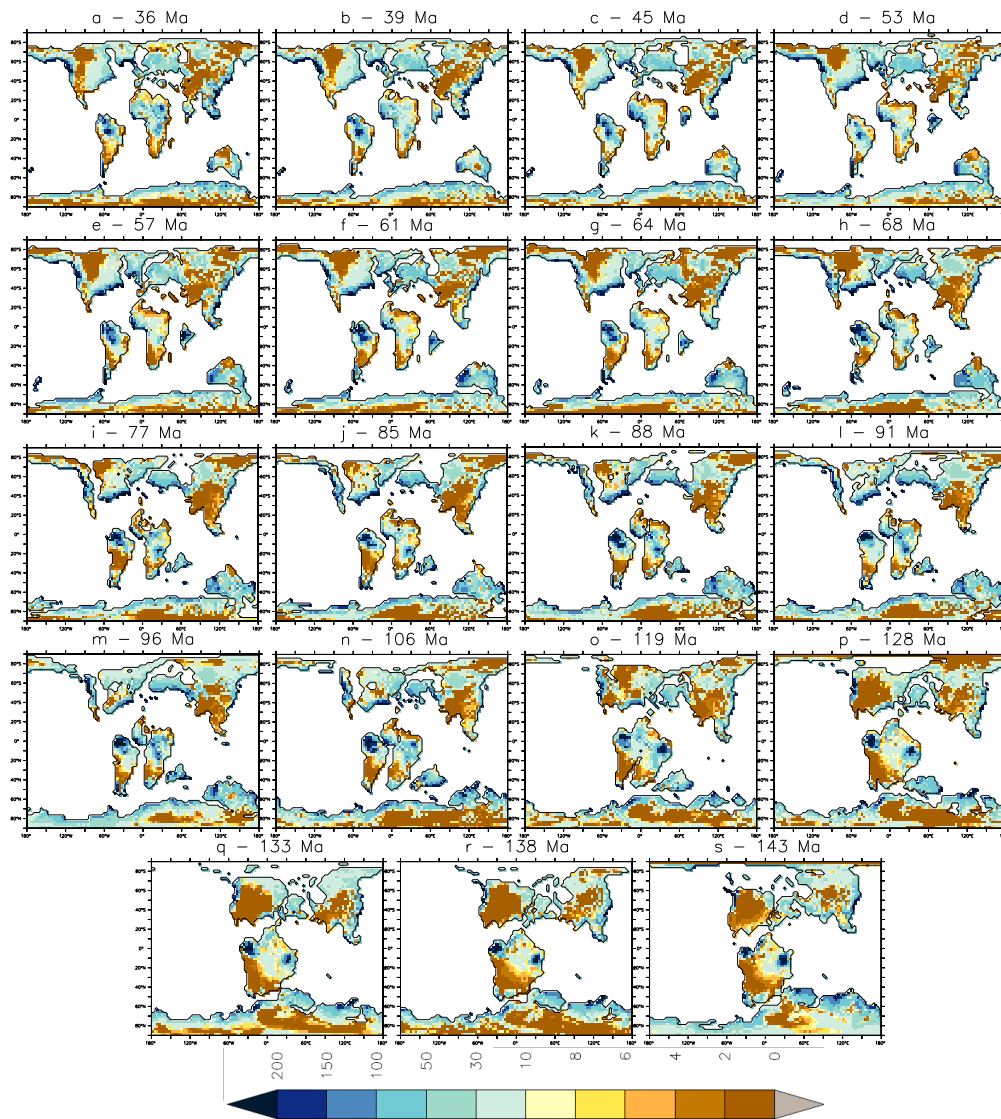
In the fixed CO<sub>2</sub> simulations, the distribution of runoff changes significantly during the Cretaceous-Eocene period (Figure  
240 3). During the early Cretaceous, large areas of North America, South America, Asia, and Antarctica have low or zero runoff,  
likely due to low moisture transport from the oceans given the converged continental configuration at this time. There are  
some areas of high runoff, such as the northern coast of Gondwana, along with parts of the western Amazon and east Africa.  
Into the mid-late Cretaceous, North America and Antarctica become much more humid while Asia becomes more arid. The  
Amazon remains persistently humid, while the Atlantic coast of North America has high runoff, possibly due to increases in  
245 evaporation (and or tropical cyclones) associated with the expansion of the Atlantic. The west coast of North America has  
especially high runoff, with zonal means indicating that runoff here is greater than at the equator, although runoff falls towards  
the late Cretaceous. Into the Cenozoic, areas such as India and western Australia have a significant increase in runoff. India  
crosses the equator at this time, so the increased runoff may be related to interactions with the ITCZ. Towards the end of  
the Eocene, areas such as North America and Asia become more arid, while runoff in the southern mid-latitudes increases  
250 slightly. Much of the world however becomes more arid, reflecting the global drop in runoff at the end of the Eocene, possibly  
linked to the general cooling trend and falling evaporation rates from the oceans from around 49 Ma into the Eocene-Oligocene  
boundary.

## 4 Discussion

Silicate weathering fluxes at 2.85 times the pre-industrial CO<sub>2</sub> show a strong relationship with mean global runoff rates and a  
255 strong inverse relationship with atmospheric CO<sub>2</sub> concentrations (Figures 1-2 and Appendix Figures A2-A3,  $r = 0.88$  and  $-0.96$ ,  
respectively), indicating that changes in global runoff driving changes in silicate weathering fluxes are the primary control on  
steady-state CO<sub>2</sub> concentrations over the modelled period. However, there are some inconsistencies in these trends. Notably,  
global silicate weathering fluxes increase towards the late Eocene despite a fall in global runoff rates. To understand these  
inconsistencies in the global trend, we investigate the relationship between runoff rates and silicate weathering fluxes at the  
260 regional level.

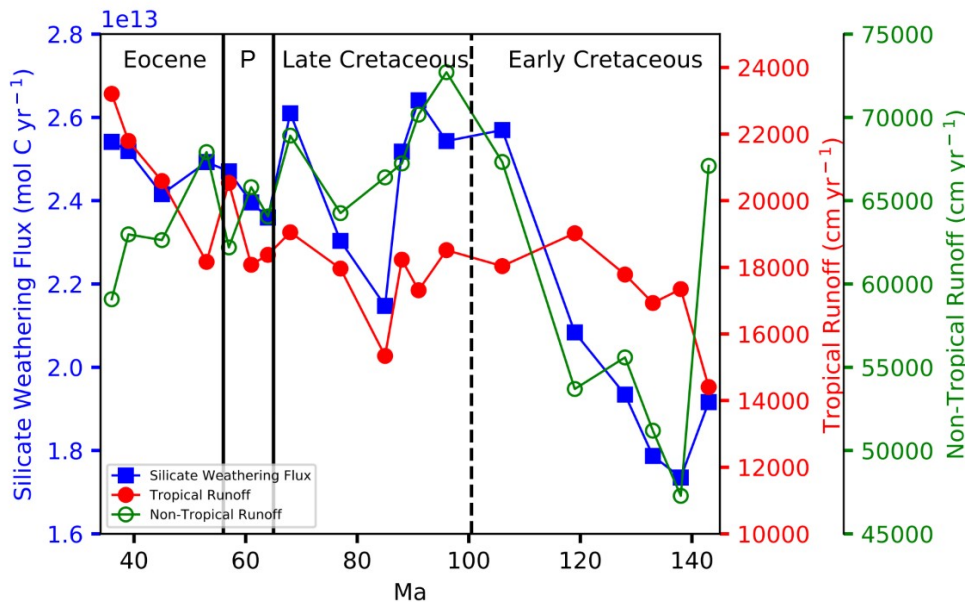
### 4.1 Regional Trends

During the Eocene, within the GEOCLIM simulations both mean and total global runoff from the fixed CO<sub>2</sub> runs falls, yet  
silicate weathering fluxes rise, an inverse of the pattern seen during the Cretaceous where runoff rates and weathering fluxes  
show identical patterns (Figure 2). During the early-Cretaceous, silicate weathering fluxes are strongly controlled by non-  
265 tropical runoff ( $>30^{\circ}$ N/S), showing an almost identical pattern until the mid-Cretaceous (Figure 4). This pattern weakens



**Figure 3.** Regional mean annual runoff ( $\text{cm yr}^{-1}$ ) maps from the earliest-Cretaceous (s) to the latest-Eocene (A). An arid climate prevails during the early-Cretaceous (s-o), particularly in the continental interior of North America. Runoff increases significantly from the mid-Cretaceous, with very high runoff rates present in Amazonia (m-k). High runoff totals are present in India during the latest-Cretaceous and the Palaeocene as it crosses the equator (h-e). The world becomes slightly more arid through the Eocene (d-a).

somewhat around 86 Ma, but strengthens by the end of the Cretaceous. Into the Cenozoic, the pattern weakens again and silicate weathering fluxes become anti-phased to non-tropical runoff. In contrast, silicate weathering fluxes in the Eocene appear to be more strongly controlled by runoff rates in the tropics (Figure 4). Similarly, tropical runoff changes in the Cretaceous does not appear to have a significant impact on silicate weathering fluxes, with the exception of a brief period during the late-Cretaceous



**Figure 4.** Modelled silicate weathering fluxes (blue squares) plotted against total tropical runoff (red circles) and total non-tropical runoff (green open circles). The tropics are defined here as all areas within 30°N/S. Changes in silicate weathering fluxes are driven broadly by non-tropical runoff during the early-Cretaceous and by tropical runoff during the Cenozoic. Note that Palaeocene has been abbreviated to P.

270 (86 Ma). There is a noticeable weakening of the influence of non-tropical runoff on weathering fluxes from the Cretaceous to the Cenozoic, although due to the small number of simulations in this period ( $n = 6$ ), confidence in this relationship is low.

The shift from non-tropical to tropical runoff controlled weathering appears to represent a regime change in the long-term climate pattern seen during the modelled period. Theoretically, a continental configuration with a greater land area in the low-latitudes, where precipitation is generally highest, would favour lower global CO<sub>2</sub> concentrations by increasing silicate weathering rates (Gibbs and Kump, 1994; Otto-Bliesner, 1995; Godd ris et al., 2014). However, changes in continental positioning and orogeny will alter ocean-atmospheric circulation and evaporation patterns, which would then alter climatic variables, especially runoff (Barron et al., 1989; Lunt et al., 2012). Furthermore, runoff patterns will also be affected by whether continents are in a dispersed or converged (i.e. supercontinental) configuration.

280 The shift in weathering controls seen in the GEOCLIM simulations is the result of a change in the effects of palaeogeography on weathering fluxes. During the Cretaceous, the models suggest that increased weathering occurred as a result of the climate becoming more humid due to the break-up of Pangaea. The break-up of Pangaea promoted higher weathering fluxes through evaporation by increased ocean area and more favourable moisture transport to the continental interiors. In contrast, during the Cenozoic both total land areas and tropical land areas increased. During the same period, modelled total global runoff

falls, but tropical runoff increases, resulting in both a greater weatherable area and a more intense weathering environment  
285 in the tropics. As such, the shift in weathering controls in the Cenozoic from non-tropical to tropical runoff is the result of  
the continents moving into a configuration which promotes higher weatherability, while during the Cretaceous changes in  
weathering fluxes were largely caused by climate changes induced by the break-up of Pangaea which increased runoff in the  
previously arid continental interiors.

## 4.2 Comparison to Proxy Data

290 While the GEOCLIM simulations in this study represent an improvement over previous studies due to their higher spatial-  
temporal resolution and better constrained palaeogeography, they are nonetheless an obvious simplification relative to both  
full GCM studies (which provide more realistic modelling of climate change processes, and feedbacks in particular, rather  
than the linear interpolation used here) and real-world settings. Because the primary aim of this study is to assess the impact  
of palaeogeographic changes on potential global “weatherability”, simplifications such as a uniform palaeolithologies were  
295 used in the absence of well-constrained data for such fields. Still, it is naturally of interest to compare the modelled CO<sub>2</sub>  
concentrations produced in this study with those derived from previous modelling studies and proxy data. Such a comparison  
may provide an indication of the potential impact of changing weatherability on long-term CO<sub>2</sub> concentrations through the  
Cretaceous-Eocene period.

Figure 1 presents the CO<sub>2</sub> output produced in this study plotted (henceforth referred to as the GEOCLIM model output)  
300 against three CO<sub>2</sub> records derived from proxy data (Anagnostou et al., 2016; Foster et al., 2017; Consortium\*† et al., 2023)  
and the 145-34 Ma portion of the record from Godd ris et al. (2014). The Anagnostou and Hoenisch records only cover  
the Cenozoic and latest Cretaceous (in the case of the Hoenisch record). Through the Cretaceous-Eocene period, there are  
three periods in which the trends in the GEOCLIM model output agrees with the proxy records: the early-Cretaceous, the  
late Cretaceous, and the mid-Eocene. Both the GEOCLIM model output and the record from Foster et al. (2017) indicates  
305 relatively high (>800 ppm) CO<sub>2</sub> concentrations in the earliest-Cretaceous, followed by a general decreasing trend until 125  
Ma. The high CO<sub>2</sub> concentrations in the early-Cretaceous are likely the result of the converged continental configuration  
which is unfavourable for high weathering fluxes. Both records indicate a rise in CO<sub>2</sub> just after the earliest-Cretaceous (138  
Ma, although only one GEOCLIM simulation is available between 140 and 135 Ma) . At this time, the continental interiors  
become more arid, as reflected by global runoff falling from  $\sim 82 \times 10^3 \text{ cm yr}^{-1}$  to  $\sim 64 \times 10^3 \text{ cm yr}^{-1}$  over a period of 5 Myr  
310 (Figures 2 and 3). The fall in global runoff leads to a reduction of silicate weathering fluxes, resulting in a CO<sub>2</sub> increase.

A rise in CO<sub>2</sub> during the late-Cretaceous ( $\sim 85$  Ma) occurs in both the GEOCLIM model output and the record from Foster  
et al. (2017). During the same period, a drop in global weathering fluxes (especially in the tropics) and a fall in total tropical  
runoff occurs (Figure 4). In both records, the rise in CO<sub>2</sub> is relatively brief (10-20 Myr) and of approximately the same  
magnitude ( $\sim 200$  ppm). There is notable disagreement between the Foster and Hoenisch records at the end of the Cretaceous,  
315 with Foster et al. (2017) showing a significant drop in CO<sub>2</sub> concentrations while Consortium\*† et al. (2023) has a more modest

decrease down to around 700ppm. There is significant uncertainty in the Hoenisch proxy record at this time, and the GEOCLIM model output is similar to the mean value suggested by the Hoenisch record.

The proxy records in the early Eocene show different CO<sub>2</sub> concentrations, ranging from around 600ppm in the Foster record to 1800ppm in the Anagnostou record. The GEOCLIM model output shows relatively steady CO<sub>2</sub> concentrations at this time, while the Anagnostou and Hoenisch records show much higher CO<sub>2</sub> concentrations and a sharper decrease. A sharp increase in CO<sub>2</sub> concentrations is shown in the Hoesnich record but this occurs between simulations in the GEOCLIM model output and thus our climate simulations may not be capturing this high CO<sub>2</sub> period. All records and the GEOCLIM model output indicate a drop in CO<sub>2</sub> of varying magnitudes towards the mid-Eocene, with the GEOCLIM model output showing the smallest drop in CO<sub>2</sub> (< 100 ppm) while the records from Anagnostou et al. (2016) and Consortium\*† et al. (2023) indicate a much larger drop of around 600 ppm and 400ppm respectively. A fall in CO<sub>2</sub> from the early to mid-Eocene is often attributed to increased silicate weathering (e.g. Raymo and Ruddiman, 1992). Indeed, our modelled Eocene weathering fluxes are generally high and with a general rising trend towards the end of the Eocene, which coincides with falling CO<sub>2</sub> concentrations (Figures 1 and 2). Zonal weathering plots indicate a wider area of high weathering fluxes in the tropics (focused on Eastern Asia and India) at this time relative to other time periods (e.g. mid-Cretaceous) in this study (30° N/S)(Appendix Figure A4).

#### 330 4.2.1 Impact of Variable Degassing Rate

Based on calculations from Van Der Meer et al. (2014) (used here as an example of differing degassing rates), degassing rates peak in the early-Cretaceous and fall gradually through the late-Cretaceous and Cenozoic periods. The variable degassing simulations produced a record similar to the GEOCLIM simulations using a fixed degassing rate of  $1 \times 10^{13}$  mol C yr<sup>-1</sup>, which is similar to the range of degassing rates in Van Der Meer et al. (2014). In general, the variable degassing simulations produced slightly higher steady-state CO<sub>2</sub> concentrations during the Cretaceous, but slightly lower steady-state concentrations during the Cenozoic (Figure 1). Steady-state CO<sub>2</sub> concentrations fall sharply in the latest-Eocene in response to a decrease in degassing rates that continues into the Neogene. Both the constant degassing and variable degassing simulations are broadly similar however, suggesting the impact of variable degassing is relatively minor. The similarity of both the constant and variable degassing CO<sub>2</sub> timeseries implies that on timescales of tens of millions of years, and in the context of this time period and the weathering model used, changes in weathering have a greater control on CO<sub>2</sub> than changes in degassing. Given that more recent studies of degassing rates have indicated that degassing during the late Mesozoic and early Cenozoic was relatively stable (Müller et al., 2022), this further suggests that degassing had a relatively limited impact on atmospheric CO<sub>2</sub> concentrations relative to changes in weatherability.

### 4.3 Comparison to Previous GEOCLIM Palaeogeography Study

#### 345 4.3.1 Comparison of Results With Previous Work

The model outputs presented in this study demonstrate that the long-term climate (under constant degassing rates) is strongly controlled by silicate weathering fluxes, which in turn are predominately controlled by runoff rates. Palaeogeographical changes

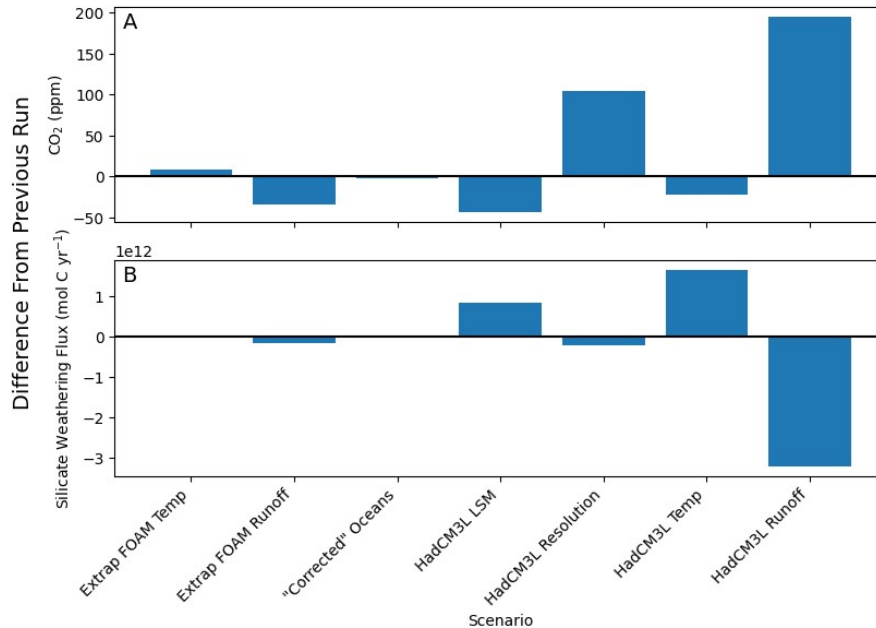
are likely to have had significant impacts on both the intensity and distribution of runoff. These findings are in line with those of Godd ris et al. (2014), which found a strong direct palaeogeographical influence on runoff, and thus weathering rates and long-term atmospheric CO<sub>2</sub>. Both Godd ris et al. (2014) and this study found a general decreasing trend in atmospheric CO<sub>2</sub> from the Cretaceous to the early Cenozoic, although this study has a higher temporal resolution during that period (19 simulations vs 5). Godd ris et al. (2014) noted a number of potential climate impacts from palaeogeographical changes, based on their GEOCLIM study with FOAM inputs.

Godd ris et al. (2014) found that a converged, or supercontinental, arrangement inhibited weathering fluxes leading to high CO<sub>2</sub> concentrations. In contrast, a dispersed continental configuration favours higher weathering fluxes. While the time period in this study does not cover the formation of Pangaea, during the earliest Cretaceous the continents were still in a converged configuration (Figure 3) and coincided with the highest CO<sub>2</sub> concentrations and lowest weatherability in the modelled period. Godd ris et al. (2014) showed significant rises and falls in CO<sub>2</sub> (up to 20x PAL) within ~10 Myr occurring during the formation and subsequent break-up of Pangaea associated with changes in weathering fluxes. Similarly, this study indicates rapid changes in CO<sub>2</sub> (200-300 ppm within 5 Myr) as Gondwana and Laurasia break apart, indicating the potential for geologically rapid CO<sub>2</sub> changes as a result of palaeogeographical changes.

Godd ris et al. (2014) also noted the potential climate impact of high weathering rates on small continental landmasses in tropical areas. During the Rhaetian period of the Triassic, south China crossed the tropics and contributed 17% of the global CO<sub>2</sub> drawdown despite a small land area relative to Pangaea (Godd ris et al., 2014). This disproportionate CO<sub>2</sub> drawdown may be the result of a generally arid climatic context during the Triassic, as a similar result is not seen in this study (Figure 3), despite the passage of India across the tropics during the Cretaceous-Eocene. Although runoff rates (and thus weathering rates) on the Indian subcontinent increase significantly as it crosses the tropics, its passage does not drive a large decrease in atmospheric CO<sub>2</sub> (Figures 2 and 3). The rise in weathering rates occurs just after a global drop in runoff rates and weathering fluxes. Thus, while the passage of India through the tropics raises global weathering fluxes significantly, the effect of this rise is counteracted by the fall in global weathering fluxes as a result of increasing aridity. As such, the influence of small continental landmasses on long-term CO<sub>2</sub> concentrations appears to be variable and largely dependent on the prevailing global climate state, with CO<sub>2</sub> concentrations more sensitive to change under a more arid global climate. In this study, a stronger influence on long-term weathering rates occurs as the result of a continental configuration that favours either increased evaporation from the oceans and moisture transport to continental interiors or a general increase in tropical land areas.

#### 4.3.2 Evaluation of the Role of GCM

A notable advance made by this study on the work of Godd ris et al. (2014) is the higher resolution GCM and palaeogeography inputs, significantly improving our ability to assess the impact of regional changes in climate and palaeogeography on global weathering fluxes. A comparison between a FOAM and a HadCM3L simulation of the early Eocene (52Ma) reveals significant differences between the two inputs used and produced substantial differences in steady-state CO<sub>2</sub> (299 ppm and 505 ppm, respectively). Furthermore, a step-by-step transition of variables from all FOAM to all HadCM3L inputs was also undertaken,



**Figure 5.** Impact of incremental changes between FOAM and HadCM3L climate inputs in GEOCLIM (A) Changes in modelled steady-state CO<sub>2</sub> in ppm (after 1 Myr) for input changes relative to the previous run and (B) changes in global silicate weathering fluxes at CO<sub>2</sub> fixed to 2.85 times the pre-industrial level (mol C yr<sup>-1</sup>) relative to the previous run. The variable changed in each run is shown as the x-axis. Changes in weathering fluxes are essentially an inverse of the pattern of changes in steady-state CO<sub>2</sub>.

to investigate the independent impact of each of the differences between the two GCMs. Initially, the FOAM climate variables were extrapolated using the same method used to generate climate data at 11 CO<sub>2</sub> concentrations for the HadCM3L data (Appendix Equations A.1 and A.2). This process allowed us to test whether our interpolation resulted in any significant deviation from the original dataset. Then a minor change to the ocean layout was made, followed by the introduction of a low-resolution (48x40) version of the HadCM3L LSM. Then the model resolution was increased to match the HadCM3L resolution (96x73). The HadCM3L temperature data was then included, replacing the FOAM temperature inputs, followed finally by replacing the FOAM runoff inputs with those from the HadCM3L simulation.

The step-wise transition between FOAM and HadCM3L revealed that the majority of the differences in steady-state CO<sub>2</sub> concentrations can be explained by differences in model resolution and runoff (Figure 5), as these produced the greatest changes in steady-state CO<sub>2</sub> values. HadCM3L produces a globally more arid world than the FOAM simulation and shows far greater regional variability. These results are significant as previous studies (both from field and model data) have suggested that a significant proportion of global weathering fluxes may be the result of so-called “hot-spots”, i.e. small land areas with high weathering fluxes such as volcanic islands and mountain ranges (e.g. Kent and Muttoni, 2013). While FOAM inputs show areas of higher runoff associated with mountain ranges, the higher resolution of the HadCM3L inputs relative to FOAM inputs may

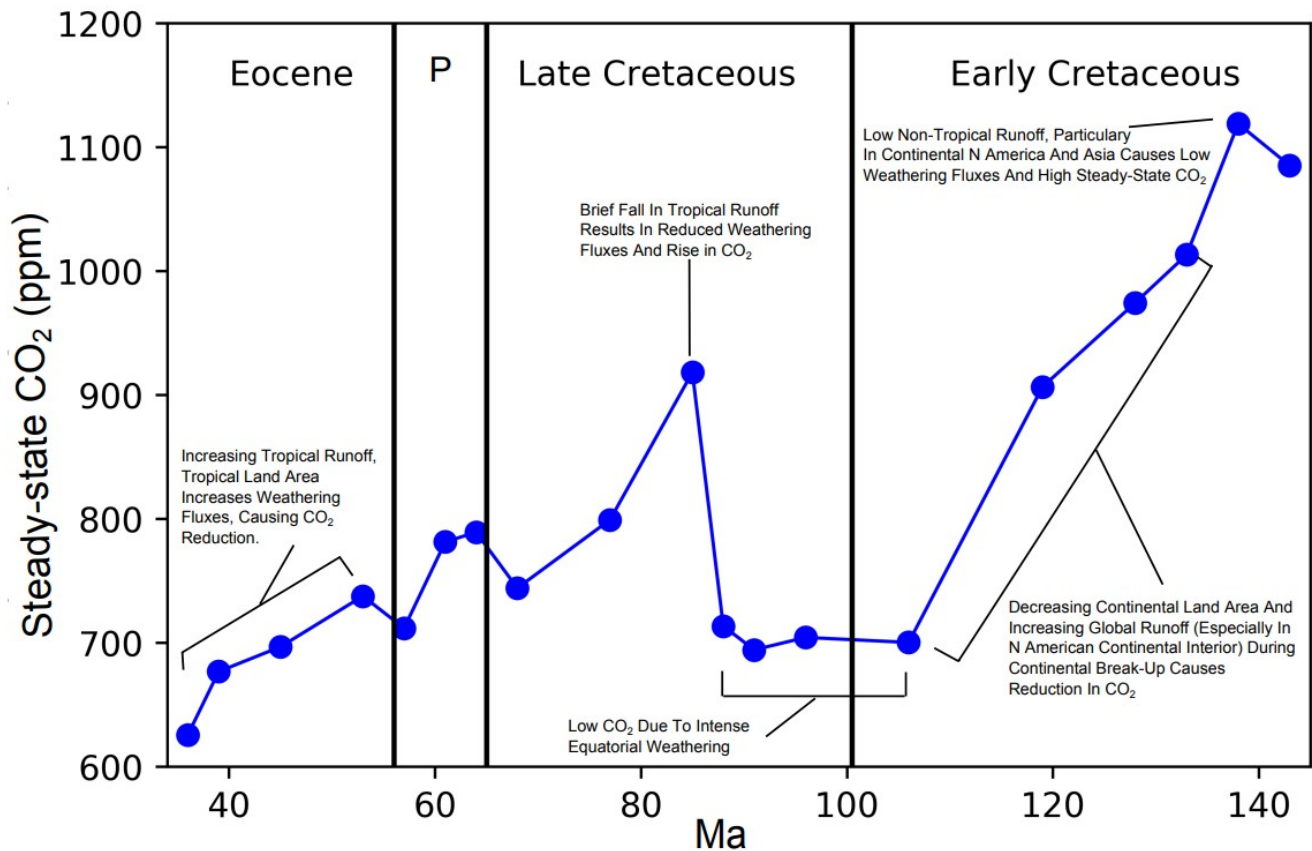
395 be sufficient to more accurately resolve weathering hot spots. Furthermore, given that weathering fluxes are highly spatially  
variable, the HadCM3L simulation likely better constrains such features and thus provides a better estimate of global weather-  
ing fluxes. This study indicates a number of areas of high weathering fluxes, particularly outside of the tropics, associated with  
areas of high relief in the GCM simulations, particularly the Laramides and Appalachians (north America) and the southern  
Andes (Figure 3). In the original HadCM3L simulations these areas have high runoff rates, likely associated with orographic  
400 intensification of rainfall. Thus, the GEOCLIM simulations in this study may provide a better spatial estimate of weathering  
fluxes than those using FOAM inputs. These regions may be partly responsible for the sharp rise in non-tropical runoff in the  
early-Cretaceous (Figure 4), and further support the role of active mountain ranges in influencing CO<sub>2</sub> drawdown (Raymo and  
Ruddiman, 1992; Riebe et al., 2004; West et al., 2005; Godd ris et al., 2017).

#### 4.4 Implications for Palaeoclimates

405 The GEOCLIM modelled CO<sub>2</sub> concentrations presented in this study reflects the changes in global “weatherability” through  
the Cretaceous-Eocene period, indicating a general trend towards increased global weatherability during that time, punctuated  
by a decrease at ~80 Ma. Although comparison with CO<sub>2</sub> records from proxy data suggest that not all changes in long-term  
CO<sub>2</sub> can be ascribed to changes in global weatherability, the three records indicate that changes in global weatherability  
have sufficiently powerful impacts on geologically short timescales (~10 Myr) to be expressed in long term CO<sub>2</sub> records (i.e.  
410 hundreds of ppmv), further supported by the three periods of agreement between the records. These changes in weatherability  
are primarily associated with the break-up of Pangaea during the early-Cretaceous, and changes in tropical land areas and  
runoff rates in the mid-Cretaceous and the Eocene, respectively. Based on these findings, the implications for palaeoclimates  
are twofold.

Firstly, this study reinforces the conclusions of previous studies that continental configuration is a strong control on long-  
415 term CO<sub>2</sub> concentrations by influencing runoff (Otto-Bliesner, 1995; Gibbs et al., 1999; Donnadieu et al., 2004; Godd ris et al.,  
2014). Supercontinental configurations typically reduce global weatherability and favour high CO<sub>2</sub> concentrations by limiting  
global runoff, while dispersed configurations typically increase global weatherability and favour lower CO<sub>2</sub> concentrations as  
the dispersed configuration favours moisture transport to continental interiors.

Secondly, this study demonstrates the potential for relatively localised changes in weatherability to have global impacts. Such  
420 local changes may result in a temporary decoupling of globally averaged runoff rates from globally averaged weathering rates.  
During the Eocene, non-tropical runoff totals fall substantially but tropical runoff increases significantly, likely associated with  
an increase in tropical land areas at the same time. Similarly, during the mid-Cretaceous, tropical runoff totals remain stable  
but non-tropical rainfall increases. The increase in non-tropical rainfall is sufficient to increase silicate weathering fluxes and  
lower atmospheric CO<sub>2</sub> concentrations by 200 ppm over 13 Myr.



**Figure 6.** GEOCLIM steady-state CO<sub>2</sub> record with annotations showing major controls on steady-state CO<sub>2</sub> concentrations. Significant changes in steady-state CO<sub>2</sub> are typically due to changes in runoff and continental land areas. Note that Palaeocene has been abbreviated to P.

## 425 5 Conclusions

This study explores the potential impact of changing “weatherability” during the Cretaceous-Eocene period on long-term CO<sub>2</sub> concentrations, as well as the impact of using high-resolution GCM data on modelled weathering rates and steady-state CO<sub>2</sub> concentrations. The palaeogeography analysis showed that “weatherability” changed significantly through the Cretaceous-Eocene period as Pangaea broke up and the continents entered a more dispersed configuration. Changes in weatherability as a result of palaeogeographical change are due to two processes. The first process is increased evaporation and more favourable moisture transport as continents became more dispersed, which occurred during the early-Cretaceous. The second process is due to continents moving into more humid zones, which occurred during the late-Eocene.

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Steady-state CO<sub>2</sub> concentrations were initially high in the early-Cretaceous due to low total global runoff inhibiting weathering fluxes. As Pangaea broke up, evaporation from the ocean increased and improved moisture transport to the continental interiors, increasing runoff rates and weathering fluxes, resulting in lower steady-state CO<sub>2</sub> concentrations. Into the Cenozoic however, global weatherability appears to switch regimes. In the Cenozoic, weatherability appears to be determined by increases in tropical land area, allowing for greater weathering in the tropics. Furthermore, global runoff fell in the late Eocene but silicate weathering fluxes continued to increase. The increase in silicate weathering is due to an increase in total tropical runoff as land areas in the tropics increased, which is sufficient to allow global weathering fluxes to increase despite a fall in total global runoff.

We also investigated and quantified the impact of using different GCM datasets in GEOCLIM. The high-resolution HadCM3L datasets produced substantially different steady state CO<sub>2</sub> concentrations relative to FOAM datasets, primarily due to changes in runoff reconstructions. The use of high-resolution GCM data was instrumental in revealing these potential impacts, and highlights the benefits of such data in palaeo weathering studies.

While the aim of this study was to investigate the role of palaeogeography on weatherability, the CO<sub>2</sub> concentrations produced by GEOCLIM was also compared with proxy derived records from the literature to determine what aspects, if any, of long-term CO<sub>2</sub> change could have been driven by changes in weatherability. Three periods were identified when the GEOCLIM model output agreed with the proxy CO<sub>2</sub> records and thus suggest that changes in weatherability were perhaps influencing global climate: the early-Cretaceous, the late-Cretaceous, and the mid-Eocene. High CO<sub>2</sub> concentrations in the early Cretaceous are likely due to the arid nature of the continental interior. A brief (10-20 Myr) rise in CO<sub>2</sub> during the late-Cretaceous appears to be the result of falling tropical runoff, causing global weathering fluxes to fall. Finally, a falling trend in CO<sub>2</sub> during the mid-Eocene appears to be linked to increased global weatherability as tropical land areas increase.

The periods of agreement with the three CO<sub>2</sub> records indicate that weatherability changes were sufficient in magnitude to affect the long-term climate of the Cretaceous-Eocene period. Furthermore, this study shows the potential for regional changes (e.g. localised changes in tropical runoff and land area) to have global impacts, and that such regional changes may be more significant than global averages for determining long-term climate.

*Author contributions.* N.H: Conceptualization, Methodology, Formal Analysis, Investigation, Writing – Original Draft

D.L: Conceptualization, Methodology, Resources, Writing – Review and Editing, Supervision

Y.G: Methodology, Resources, Writing – Review and Editing

R.P: Conceptualization, Writing – Review and Editing, Project Administration, Funding Acquisition

H.B: Writing – Review and Editing, Supervision, Project Administration

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The views expressed in the paper are those of the authors and do not necessarily reflect the position of the Environment Agency.

*Competing interests.* Some authors are members of the editorial board of *Climates of the Past*.

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## Appendix A

- 610 Because GEOCLIM interpolates climate variables based on climate data at specified CO<sub>2</sub> concentrations, it was necessary to ensure that both the FOAM and HadCM3L input data had consistent CO<sub>2</sub> concentrations. FOAM was previously run at 11 CO<sub>2</sub> concentrations between 160 and 1400 ppm, while HadCM3L was previously run only at 560 and 1120 ppm. For consistency,

a methodology was developed to produce “synthetic” model output from HadCM3L at the same CO<sub>2</sub> concentrations as for FOAM. This was deemed particularly important because GEOCLIM model runs with Ypresian FOAM inputs have previously  
 615 produced CO<sub>2</sub> concentrations below 560 ppm (Lefebvre et al., 2013). In GEOCLIM, if modelled CO<sub>2</sub> concentrations exceed the available CO<sub>2</sub> range, the climate variables will cease to update.

To address these issues, the HadCM3L climate inputs were extrapolated to the same 11 CO<sub>2</sub> concentrations to cover the same range as the FOAM data. Two separate extrapolation methods were used to calculate the temperature and runoff to better reflect their sensitivity to CO<sub>2</sub> concentration changes in observed data. The HadCM3L temperature ( $T_x$ ) for a CO<sub>2</sub> concentration of  
 620  $CO_{2x}$ , was approximated as:

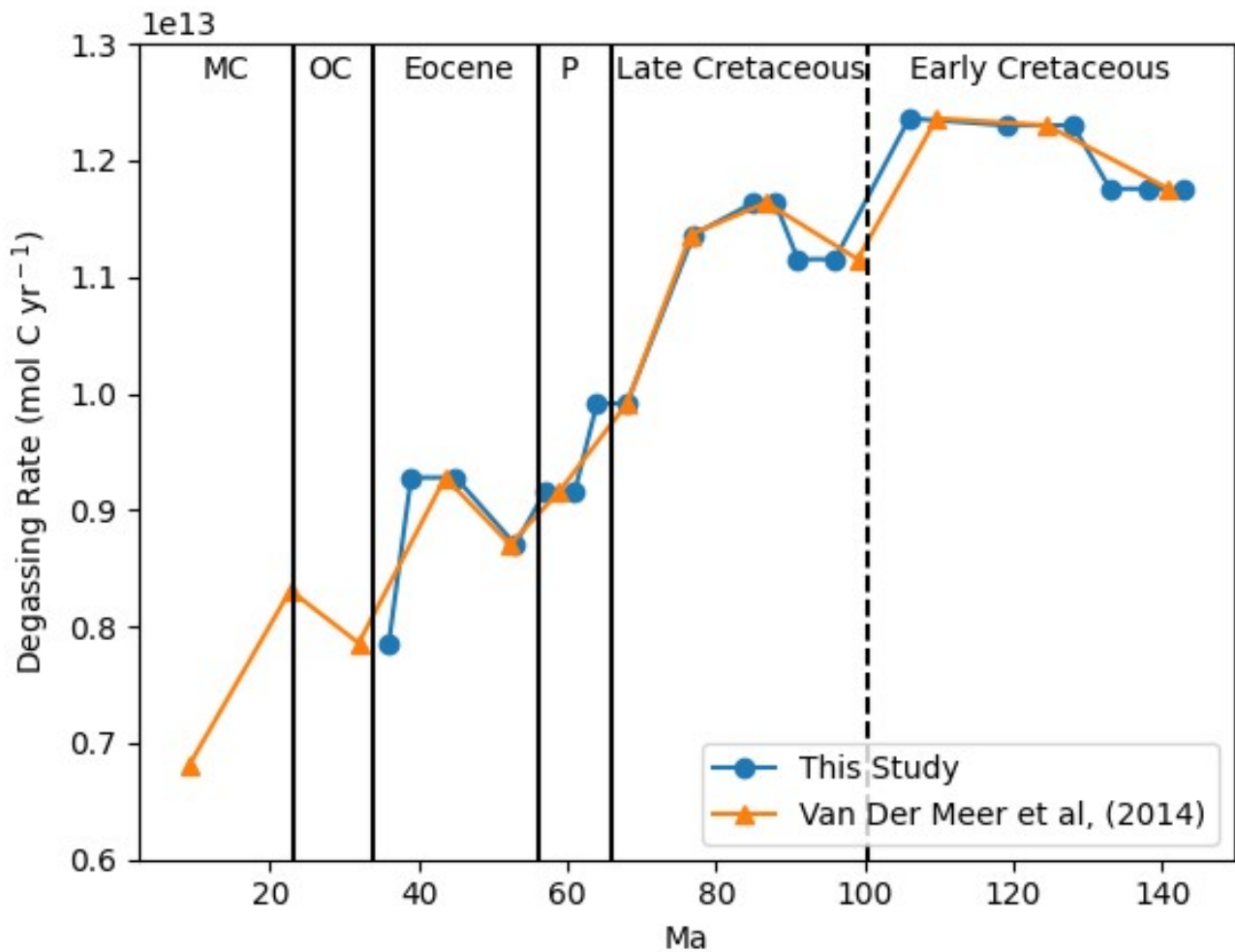
$$T_x = T_{560} + (T_{diff} * \frac{\ln \frac{CO_{2x}}{280}}{\ln 2} - 1) \quad (A1)$$

Temperature values were extrapolated based on absolute differences between the 560 and 1120 ppm temperature outputs (Equation A.1).  $T_x$  is the temperature value at any given CO<sub>2</sub> concentration, where  $T_{560}$  is the temperature at 560 ppm,  $T_{diff}$   
 625 is the difference in temperature between 1120 and 560 ppm, and  $CO_{2x}$  is a specific concentration of atmospheric CO<sub>2</sub> (ppm).

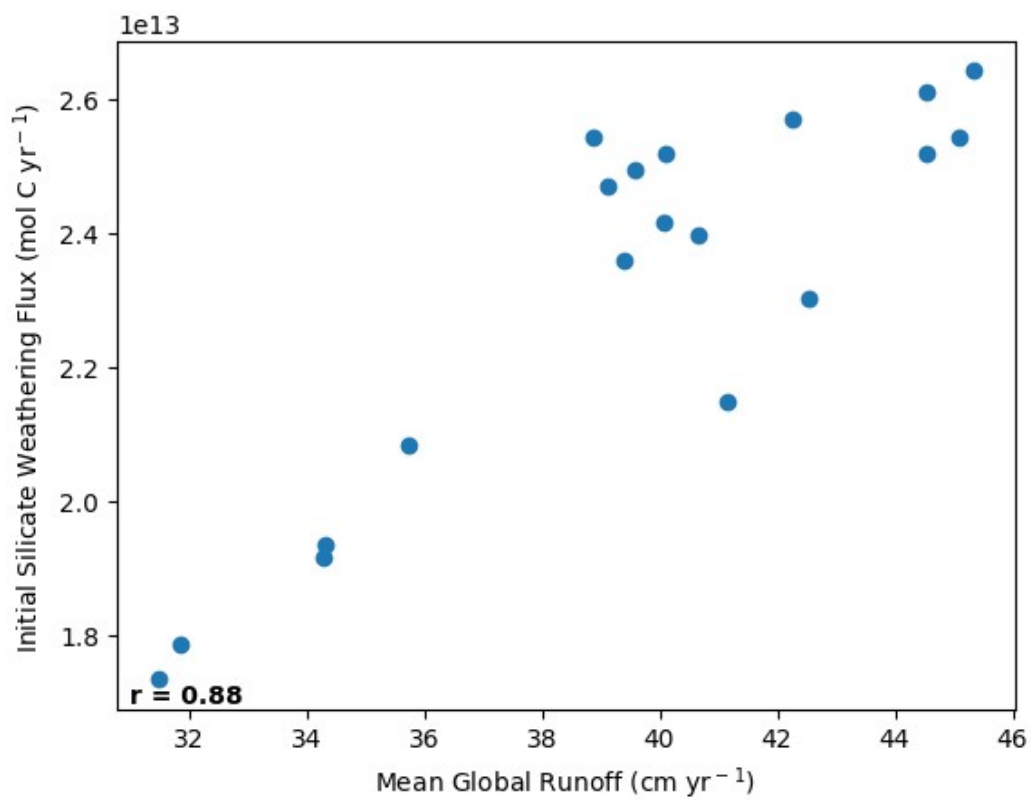
Runoff was extrapolated in a similar fashion to temperature, except that a constant ratio rather than a constant absolute difference was assumed per CO<sub>2</sub> doubling, because this relationship is found in the model results and absolute value changes would result in large areas of negative runoff and excessive aridity at lower CO<sub>2</sub> concentrations which were deemed to be unrealistic:

$$630 \quad R_x = R_{560} * \exp\left(\left(\frac{CO_{2x} - 560}{560}\right) * \ln(R_{diff})\right) \quad (A2)$$

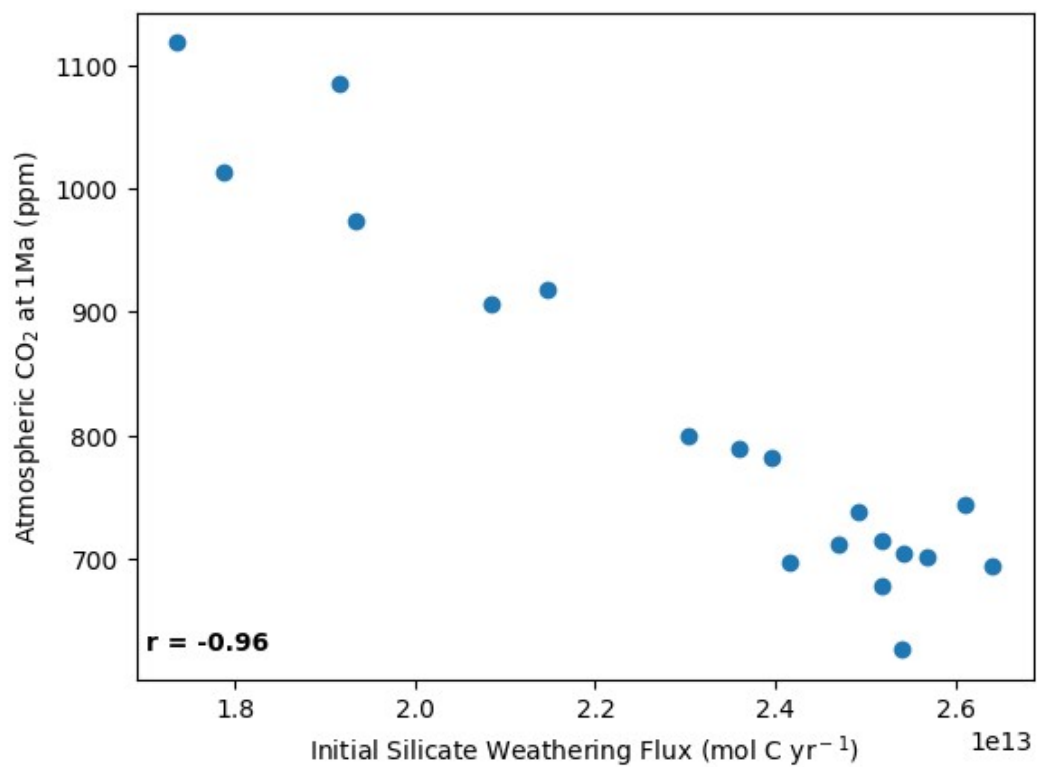
Equation A.2 was used to extrapolate runoff rates (cm yr<sup>-1</sup>),  $R_x$ , at any given CO<sub>2</sub> concentration.  $R_{560}$  represents the runoff rate at 560 ppm, while  $R_{diff}$  is the magnitude difference between runoff values at 1120 and 560 ppm.



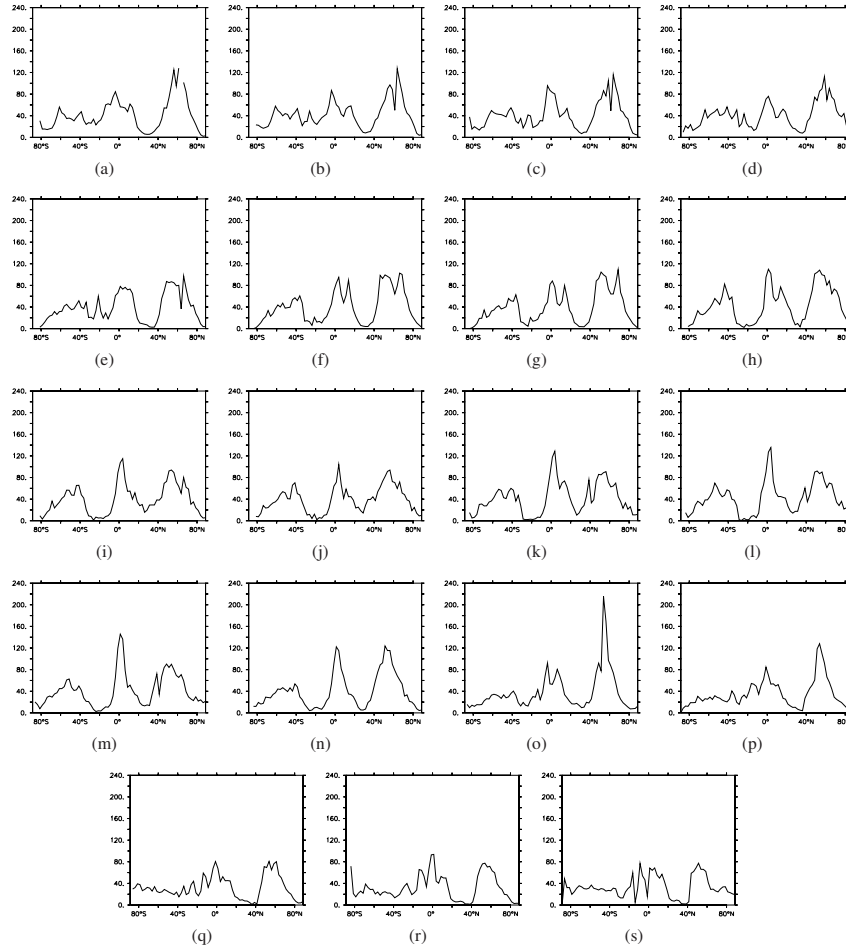
**Figure A1.** Degassing rates from Van Der Meer et al. (2014) (blue) used to provide variable degassing rates for the 19 time slices used in this study (green). Each time slice uses the rate from Van Der Meer et al. (2014) that is nearest in time. Degassing rates are highest in the early to mid Cretaceous (110-130 Ma) and gradually fall into the Paleogene with occasional rises. MC = Miocene, OC = Oligocene, and P = Palaeocene.



**Figure A2.** Scatter plot of mean global runoff and initial silicate weathering fluxes from the 19 modelled timesclies. Global runoff and silicate weathering fluxes show a strong positive correlation ( $R = 0.88$ ).



**Figure A3.** Scatter plot of initial silicate weathering fluxes and steady-state atmospheric CO<sub>2</sub> concentrations from the 19 modelled timeslices. Silicate weathering fluxes and steady-state atmospheric CO<sub>2</sub> concentrations show a very strong negative correlation ( $R = -0.96$ ).



**Figure A4.** Modelled zonal mean runoff ( $\text{cm yr}^{-1}$ ) from the earliest-Cretaceous (s) to the latest-Eocene (A). Zonal runoff means are low in the early-Cretaceous (s-p), but increase significantly into the mid-Cretaceous, especially in the northern mid-latitudes (o-k). Zonal mean runoff falls slightly into the late-Cretaceous, but there is an intensification in equatorial mean zonal runoff (j-h). In the Cenozoic, zonal runoff intensifies again (g-e) but begins to weaken again during the Eocene (d-a).