Was the Late Paleocene-Early Eocene Hot Because Earth Was Flat?
An Ocean Lithium Isotope View of Mountain Building, Continental Weathering, Carbon Dioxide, and Earth’s Cenozoic Climate

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**ABSTRACT.** Hothouse climates in Earth’s geologic past, such as the Eocene epoch, are thought to have been caused by the release of large amounts of carbon dioxide and/or methane, which had been stored as carbon in biogenic gases and organic matter in sediments, to the ocean-atmosphere system. However, to avoid runaway temperatures, there must be long-term negative feedbacks that consume CO₂ on time scales longer than the ~ 100,000 years generally ascribed to ocean uptake of CO₂ and burial of marine organic carbon. Here, we argue that continental chemical weathering of silicate rocks, the ultimate long-term (multi-million year) sink for CO₂, must have been almost dormant during the late Paleocene and early Eocene, allowing buildup of atmospheric CO₂ to levels exceeding 1,000 ppm. This reduction in the strength of the CO₂ sink was the result of minimal global tectonic uplift of silicate rocks that did not produce mountains susceptible to physical and chemical weathering, an inversion of the Uplift-Weathering Hypothesis. There is lack of terrestrial evidence for absence of uplift; however, the δ²⁷Li chemistry of the Paleogene ocean indicates that continental relief during this period of the Early Cenozoic was one of peneplained (flat) continents characterized by high chemical weathering intensity and slow physical and chemical weathering rates, yielding low river fluxes of suspended solids, dissolved cations, and clays delivered to the sea. Only upon re-initiation of mountain building in the Oligocene-Miocene (Himalayas, Andes, Rockies) and drifting of these continental blocks to low-latitude locations near the Inter-Tropical Convergence Zone and monsoonal climate belts did continental weathering take on modern characteristics of rivers with high suspended loads and incongruent weathering, with much of the cations released during weathering being sequestered into secondary clay minerals. The δ²⁷Li record of the Cenozoic ocean provides another piece of circumstantial evidence in support of the Late Cenozoic Uplift-Weathering Hypothesis.

**INTRODUCTION**

Earth’s climate and the chemistry of its global ocean are intricately linked via (1) the chemical weathering of igneous silicate rocks on land, which consumes CO₂ and releases cations and alkalinity to the sea, and (2) the re-formation and uplift of igneous rocks during plate collision and subduction, which releases CO₂ (Urey and Korff, 1952; Holland, 1978; Walker et al., 1981; Berner et al., 1983; Raymo et al., 1988; Raymo, 1991; Raymo and Ruddiman, 1992; Ruddiman et al., 1997). This Urey-Walker-Berner rock chemistry cycle (see Box 1) is dependent on four processes: the planet’s system of continental drift, seafloor spreading, and plate tectonics; mountain building by continental-continental and continental-oceanic crust collisions; subduction; and arc volcanism. The cycle is generally thought to form the basis for the long-term (multi-million year) feedbacks that have kept Earth’s climate from running away to either permanent hothouse or icehouse conditions. The details of these up-kinetic-climate linkages can be discovered by unraveling the past histories of climate change and ocean chemistry that were markedly different from today over long periods of Earth’s past.

The Eocene epoch, about 56 to 34 million years ago (Ma), is one such example of an extreme climate, with very warm temperatures, up to 5°C (low latitude) and 10°C (high latitude) warmer than today (Zachos et al., 2001, 2005, 2008; Sluijs et al., 2006). Low-latitude surface ocean temperatures may have approached 40°C and bottom water temperatures were as warm as 12°C, apparently lacking high-latitude cold, dense bottom water sources. There were no high-latitude ice caps on the poles, and, as a result, sea level was almost 100 m higher than today. The continental shelves were swamped. Inland epicontinental seas were prevalent. Evolution was still recovering from the Cretaceous-Paleogene bolide impact and the Deccan Trap eruptions (KPg Boundary, 66 Ma) that killed off the dinosaurs, destroyed over 75% of all extant genera, and left the surface ocean and continental land masses with a depaupered ecological diversity incapable of producing massive amounts of organic carbon (Alvarez et al., 1980; Courtillot and Renne, 2003).

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The beginning of this “Hot Eocene” was the Paleocene-Eocene Thermal Maximum (PETM, 56 Ma), a short period when atmospheric CO₂ concentrations peaked at over 1,000 ppm, almost four times that of our current pre-Anthropocene atmosphere (Figure 1; Zachos et al., 2001, 2005; Beerling and Royer, 2011). The ocean carbonate compensation depth (CCD) was about 1,500 m shallower than today at the onset of the Early Eocene Climatic Optimum (EECO, 52 Ma) (Pälike et al., 2012), reflecting high levels of seawater CO₂ and ocean acidification (Zachos et al., 2005). Most of the literature dealing with this period—the PETM, the EECO, and the periodic hyperthermals—has emphasized the supply side of the CO₂ balance (Zachos et al., 2008) to explain not only high persistent CO₂ but also sudden carbon isotope excursions (CIE) during orbitally paced hyperthermals that each seem to have lasted about 100,000 years. These CIE events have been variously interpreted as “burps” and “bleeds” releasing (1) isotopically light methane from warmed sedimentary gas hydrate reservoirs (Zeebe, 2013, and references therein), (2) thermogenic CO₂ and CH₄ from cooking of sedimentary organic carbon by magmatic intrusions into organic-rich sediments (Svensen et al., 2004), (3) CO₂ during subduction of carbon-rich upwellings (Reagan et al., 2013), (4) CO₂ from burning of peatlands (Kurtz et al., 2003), and (5) CO₂ during subduction of carbonate-laden oceanic crust upon collision and closure of the low-latitude Tethys Seaway and initiation of the western Pacific subduction zones (Bouihol et al., 2013). A decrease in the burial of organic matter, not a source but rather a “missing sink” that would leave light carbon dioxide behind in the atmosphere-ocean system (e.g., Komar et al., 2013), has also been argued. Change in the ultimate carbon sink (continental weathering of silicates) is often invoked to explain high CO₂ during the Eocene, but it is generally modeled with only CO₂ and temperature feedbacks on weathering rates, neglecting the importance of uplift tectonics on erosion and weathering (West et al., 2005, 2012). Evidence of absence of uplift is lacking, yet absence of evidence of uplift is abundant, leading to the apparent illusion of no connection between mountain building and climate (Hay et al., 2002).

Many models of this epoch have been incapable of reproducing the magnitude of the Eocene excursions in δ¹⁸O and δ¹³C or the increase in CO₂ and consequent shoaling in the depth of the oceanic CCD (see below) (Sluijs et al., 2013; Komar et al., 2013, and references therein). The general long-term cooling trend from the middle Eocene to the Eocene-Oligocene boundary (Oi-1) and the initiation of Oligocene Antarctic glaciation and sea level drop (Figure 1) has been attributed to a slow drawdown of CO₂, but again, this is most recently ascribed to a reduction in the volcanic CO₂ source rather than an increase in the weathering of continental silicates (Lefebvre et al., 2013). Yet, one of the most important recorders of change in ocean chemistry related to continental delivery of cations to the ocean, strontium isotopes, clearly show a major change in the input of continental radiogenic ⁸⁷Sr to the ocean shortly after 40 Ma (Figure 2), coincident with the final collision of India with Southeast Asia (Bouihal et al., 2013) and weathering of the uplifted Himalayas (Raymo et al., 1988; Palmer and Edmond, 1989, 1992; Edmond, 1992).

In the following sections, we briefly describe how rates of chemical weathering of the continents (river fluxes of cations produced by silicate weathering destined for the ocean) and the texture of weathering intensity (the ratio of

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**BOX 1. THE GLOBAL UPLIFT-WEATHERING-CLIMATE CONNECTION***

\[
\text{CaSiO}_3 \xrightarrow{Tectonics + Atmospheric CO}_3 \xrightarrow{Mountain Building + Weathering} \text{CaCO}_3
\]

\[
\text{CO}_2 \leftarrow \text{Ocean (Ecology) + Subduction} \xrightarrow{Tectonics (+ Life)} \text{SiO}_2
\]

The net global reactions reflects the long-term, multi-million-year controls on atmospheric CO₂, climate, and ocean chemistry.

river-borne cations carried to the sea in dissolved form vs. that in clays) might explain the overall hothouse climate of the early Eocene. The details are contained both in the ocean chemical history of cations and in the complexities of weathering of the continents.

The basis of our argument is summarized in the feedback loops in Box 1 (Walker et al., 1981): *The long term (multi-million year) CO₂ content of the atmosphere must adjust (up or down) to the level required to achieve a quasi-steady state so that the amount of CO₂ consumed (and Ca²⁺ and Si released) during weathering of continental silicates is exactly balanced by the amount of CO₂ produced (and CaCO₃ + opal destroyed) during subduction.* The CO₂ content of the atmosphere-ocean system controls climate and, ultimately, global tectonics controls CO₂ concentrations. The stable isotopes of lithium (⁶Li and ⁷Li) are important to this theory because Li is one of the few cations that is hosted almost exclusively in silicate minerals and aluminosilicate clays and thus is directly linked to continental weathering of igneous rocks and reverse weathering in marine sediments (Aller, 2014, and references therein).

**CONTINENTAL PHYSICAL AND CONGRUENT CHEMICAL WEATHERING: GLOBAL THERMOSTAT PART 1**

Uplift of igneous rocks by tectonics and denudation by physical erosion provide both new mass and new surface area of reactant (silicate rocks) for chemical weathering, provided there is sufficient CO₂ to generate carbonic acid (H₂CO₃) and enough water to provide solvent. These reactions were first synthesized as acid-base neutralizations by Urey and Korff (1952), conceived as congruent dissolution reactions:

\[
\text{CaSiO}_3 + 2 \text{CO}_2 + 3 \text{H}_2\text{O} \rightarrow \text{Ca}^{2+} + 2 \text{HCO}_3^- + \text{Si(OH)}_4^- \quad (1)
\]

CaSiO₃ is “granite,” here simplified as wollastonite. The carbon dioxide is carbonic acid in solution in rainwater, soil water, groundwater, and rivers. This CO₂ is ultimately derived from the atmosphere even though it may have been recycled through organic matter in the terrestrial regolith. The calcium silicate “granite” represents all the igneous cation-silicate minerals that contain Mg, K, Na, Sr, and Li and are presumed to weather “congruently” (i.e., homogeneously: the cations weathered and appearing in solution in rivers are the same as the cations in the igneous minerals). Two moles of CO₂ are removed from the atmosphere for every two units of alkalinity (two equivalents of cation charge) released to the sea. The dissolved calcium and other cations, bicarbonate, and dissolved silica (silicic acid; Si(OH)₄) are eventually washed out of soils and groundwaters and carried to

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**Figure 1.** Multi-proxy view of Cenozoic atmospheric CO₂. Ten million years after the Cretaceous-Tertiary bolide impact at 66 million years ago (KPg, the Cretaceous-Paleogene boundary) destroyed most of the global continental and marine ecosystems, the atmospheric CO₂ rose to high values about three to four times the pre-Anthropocene (Holocene) values, perhaps to over 1,000 ppm. The Early Eocene Climatic Optimum (EECO) marks the warmest period. Oi-1 marks the first Antarctic permanent glaciation at the beginning of the Oligocene/ end of the “Hot Eocene.” Ma = millions of years ago. Modified from Figure 1 of Beerling and Royer (2011)
the ocean in their dissolved forms by rivers and submarine groundwater discharge. Thus, mountains are destroyed, sea salt is created, and atmospheric CO$_2$ is consumed. Concepts and models of long-term climate modulation (Berner et al., 1983; Berner, 1994; Berner and Kothavla, 2001) generally take the dissolved cation fluxes in rivers as the rates of cations released by igneous silicate mineral reactions and set this equivalent to the CO$_2$ consumed from the atmosphere. There are, of course, additional physical and chemical weathering reactions on the continents (sedimentary conglomerates, limestones, dolomites, shales, evaporites), but as Walker et al. (1981) pointed out, it’s the igneous silicate minerals that matter for long-term consumption of CO$_2$ on global tectonic time scales. We contend that uplift and erosion-denudation of igneous rocks set the long-term rates (see Chamberlin, 1899, for origin of this idea, Holland, 1978, for its modern explication, and Hay et al., 2002, and West et al., 2012, for a more complete discussion of the ideas).

One problem with the above reactions in CO$_2$ models is the assumption of congruency, that all cations released by chemical weathering end up in the dissolved load delivered to the ocean as alkalinity, that there are no solid phase products of chemical weathering. The production of secondary clay minerals carrying cations in particulate matter is generally ignored—incongruent weathering. We return to this problem below in the concept of chemical weathering intensity.

OCEANIC CALCIFICATION AND SILICIFICATION: GLOBAL THERMOSTAT PART 2

\[ \text{Ca}^{2+} + 2 \text{HCO}_3^- \rightarrow \text{CaCO}_3 + \text{H}_2\text{O} + \text{CO}_2 \]  (2)

\[ \text{Si(OH)}_4 \rightarrow \text{SiO}_2 + 2 \text{H}_2\text{O} \]  (3)

Calcifying and silicifying plankton in the surface ocean take up dissolved calcium and silica that are delivered to the sea by rivers and use them to produce shells that are then delivered to the seabed where they accumulate as carbonate- and silica-rich sediments (limestone and opal). Note that one mole of the two moles of CO$_2$ taken up in reaction (1) are released by marine carbonate production in reaction (2).

PLATE TECTONICS AND SUBDUCTION: GLOBAL THERMOSTAT PART 3

\[ \text{CaCO}_3 + \text{SiO}_2 \rightarrow \text{CaSiO}_3 + \text{CO}_2 \]  (4)

Reaction (4) reflects the transport by plate tectonics of sediment-laden oceanic crust to deep subduction zones where high heat and pressure fuse the limestone and opal to create new “granite” (wollastonite) magmas that are brought back to Earth’s surface via volcanic activity, a process that releases a mole of CO$_2$. The ocean (reactions 2 and 3) provides the shuttle that connects the uplift and weathering of the continents with the re-formation of continental rocks during
subduction. The rates of reactions (1) and (4) compete to reach a steady state of atmospheric CO₂ concentrations. These rates are dependent on plate tectonics (uplift and subduction) and on the CO₂, temperature, and erosion feedbacks on weathering (Walker et al., 1981; Berner et al., 1983; West et al., 2012). While these are simplified versions of complicated processes, they suffice to illustrate the long-term tectonic control of CO₂ that sets the planet’s thermostat. The net global reaction (Box 1) reflects the long-term, million-year controls on atmospheric CO₂, climate, and ocean chemistry.

**INCONGRUENT WEATHERING, CLAYS, AND LITHIUM ISOTOPES**

Igneous minerals seldom weather congruently to deliver only dissolved cations and silica; intermediate cation-bearing clay minerals are also produced. These clays are silica deficient and alumina enriched—the secondary aluminosilicate residues in soils. They are ubiquitous, complex, generally of mixed fine phases, and virtually impossible to separate and identify from the mass of other fine phase primary minerals and oxides accumulating in soils and being transported by rivers. For example, the incongruent weathering of K-feldspar (orthoclase) to form a potassium mica clay (biotite) is written:

\[
\frac{3}{2} \text{KAlSi}_3\text{O}_8 + \text{CO}_2 + 7 \text{H}_2\text{O} \rightarrow \frac{1}{2} \text{KAlSi}_3\text{O}_{10}(\text{OH})_2 + \text{K}^+ + 3 \text{Si(OH)}_4 + \text{HCO}_3^{-} \\
\text{(K-feldspar + carbon dioxide} \rightarrow \text{K-mica + dissolved potassium, silica, and bicarbonate)}
\]

The potassium that ends up in the clay k-mica has been weathered from orthoclase and has consumed CO₂, and if it is delivered to the ocean intact, then it represents a potassium weathered from an igneous mineral but not delivered to the sea as alkalinity (i.e., dissolved K⁺). If the potassium is weathered from the mica prior to particulate removal,

\[
2 \text{KAlSi}_3\text{O}_{10}(\text{OH})_2 + 2 \text{CO}_2 + 5 \text{H}_2\text{O} \rightarrow 3 \text{AlSi}_2\text{O}_5(\text{OH})_4 + 2 \text{K}^+ + 2 \text{HCO}_3^{-} \\
(\text{K-mica + carbon dioxide} \rightarrow \text{kaolinite + dissolved potassium and bicarbonate})
\]

The requisite CO₂ is consumed to balance the potassium released from weathering the clay. The new clay mineral produced, kaolinite, contains no cations.

\[
\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + 5 \text{H}_2\text{O} \rightarrow 2 \text{Al(OH)}_3 + 2 \text{Si(OH)}_4
\]

(kaolinite → gibbsite + dissolved silica)

Kaolinite dissolution does not consume CO₂ as there are no cations released—it is not an acid-base reaction but rather simple dissolution. It is congruent with respect to Si (all Si in kaolinite is released to solution) but incongruent with respect to Al (which ends up as gibbsite). Although experimental data are lacking that might demonstrate this reaction, octahedral Mg (and Li) in kaolinite are presumably released to solution during its dissolution.

**RIVERS AND LITHIUM ISOTOPES**

The importance of these clay mineral reactions is that Mg commonly substitutes for Al in the structural octahedral sites of many secondary clays formed by weathering—kaolinites, smectites, and illites. Lithium, because it has the same ionic size, easily substitutes for octahedral Mg (Decarreau et al., 2012), especially in mixed-layer smectites that dominate secondary mixed-layer clays. Li produced by the weathering of igneous minerals then has two fates: (1) it remains in the dissolved phase of rivers and groundwaters and is delivered to the sea, or (2) it is incorporated into secondary aluminosilicate clays. This clay-Li is isotopically very light compared to the Li in mother solutions (Huh et al., 1998, 2001; Kisakurek et al., 2005; Vigier et al., 2006, 2008, 2009; Pogge von Strandmann et al., 2006, 2008; Millot et al., 2010; Wimpenny et al., 2010). In addition, virtually all (95%) of the Li cycled on Earth’s surface resides in silicates and aluminosilicates (Stoffyn-Egły and Mackenzie, 1984; Misra and Froelich, 2012). There is little lithium in limestones, dolomites, biosiliceous deposits, or evaporites. The trace LiCO₃ in evaporites (precipitated from Li in seawater) is easily identified in rivers draining these deposits because of the correlations with chloride and sulfate. Like the evaporite- and limestone-derived Na, Ca, Mg, Cl, and SO₄ in rivers, the Li flux in rivers due to recycled marine evaporites and carbonates is easily subtracted from the fluvial Li burden in order to obtain the silicate-weathered component only.

Thus, the isotopic offset between source (Li in igneous minerals) and product (Li in clays) results in the dissolved Li in rivers and groundwaters being isotopically very heavy (Huh et al., 2001). The δ⁷Li values of igneous minerals in granites and basalts do not vary greatly (values generally lie between 0‰ and +4‰). The average global river dissolved δ⁷Li values are about 21‰ heavier than their rock-Li sources (Figure 3). Today, this reflects a global partitioning between dissolved-Li and clay-Li of about 1:4. In other words, the difference between the heavy dissolved river Li and the source rocks requires about 80% of the Li derived from igneous rock dissolution to end up...
in clays regardless of whether these clays remain behind in soils or are transported undissolved to the sea. The systematics of river-dissolved $\delta^7$Li are complicated by enormous heterogeneity because the complementary solid phase secondary clays cannot be separated from the detrital fine phases in rivers, and because the distribution and transport of Li in secondary clays (particulates) and dissolved Li (solution) are discontinuous in both space and time. As a result, analyses of river grab samples with suspended material containing secondary clays and dissolved Li do not reflect in situ weathering processes except in the broadest, whole drainage basin, sense.

For example, in Figure 3, the differences between dissolved and suspended Li in rivers are queued up in rank order of increasing $\delta^7$Li offsets from left to right (see Figure 3 legend for details of this plotting protocol). If rivers were laboratory bottle weathering experiments, we would expect a simple relationship—as dissolved-Li $\delta^7$Li increases (from left to right across the plot), the corresponding clay-Li (particulate $\delta^7$Li) should decrease from left to right. This latter effect for the particulates is barely discernible in the plot. The reasons are threefold. First, the particulates in the fine phase include random mixtures of secondary aluminosilicate and primary igneous minerals—the latter dilutes the isotopic signal we would wish to identify in the clay minerals. Second, the aluminosilicate clay $\delta^7$Li that represents the dissolved $\delta^7$Li was formed somewhere and sometime else than where the river sample was grabbed. Third, mixing of river waters and particulates downstream from the rock weathering locations (soils? groundwaters?) precludes identifying the primary isotopic sources for either phase.

Nevertheless, Figure 3 does contain important information. The data with the highest offsets (the right side of each river grouping with the heaviest $\delta^7$Li) come from portions of the drainages with the highest relief and igneous silicate lithologies. The data with the lowest offsets and lowest dissolved $\delta^7$Li typically arise from more intensely weathered...
low-relief regions of each drainage basin. We interpret this as the dissolved-Li δ⁷Li signature of weathering intensity in today’s world (see Box 2). Low weathering intensity regimes (weathering limited) with steep relief, fast uplift, and incongruent weathering generally tend to display heavy dissolved δ⁷Li with clays carrying light δ⁷Li in their octahedral sites (e.g., cation analogs to reactions 5 and 6). The ratio of clay-Li to dissolved-Li is high—most of the weathered cation-Li ends up in secondary clays. High weathering intensity (transport limited) rivers with flat relief (peneplains) tend to display light δ⁷Li with clays carrying heavy δ⁷Li in their octahedral sites (e.g., cation analogs to reactions 5 and 6). The ratio of clay-Li to dissolved-Li is low and all of the weathered cations end up in the dissolved load.

Although high weathering intensity terrains are poorly represented today in the global river δ⁷Li data set (and on the present-day globe in general), we argue below that the intensely weathered low latitudes of our Eocene planet were dominated by these flat-land, water-saturated drainages. Eocene global chemical weathering texture was much more congruent and highly intense than during the Late Cenozoic and delivered a very different δ⁷Li to the sea that dramatically affected the δ⁷Li of the ocean (Figure 4).

THE GLOBAL LITHIUM ISOTOPE CYCLE—TODAY

Today, global rivers contribute about 10 × 10⁹ moles per year of very heavy dissolved Li to the sea (δ⁷LiRivers = +23‰), much heavier than average upper continental crustal granites (δ⁷LiUCC = +1.7‰) (Figure 5a). Hydrothermal Li fluxes via seawater recirculating through the global mid-ocean ridge system contribute another 13 × 10⁹ moles per year (δ⁷LiHT = +8.3‰), slightly heavier than basalts (δ⁷LiBasalts = +3.7‰). Both input fluxes are similar in magnitude and both of the dissolved fluxes are heavier than their source rock, reflecting removal of light ⁶Li into authigenic clays (Misra and Froelich, 2012). Thus, weathering of continental and oceanic crusts today provides a dissolved Li flux to the sea that is much heavier than all rock sources (flux-weighted average δ⁷LiInput = +15‰). Note that these heavy inputs cannot explain the even heavier value of dissolved lithium in seawater (δ⁷LiSW = +31‰) (Figure 3). These isotopic offsets are enormous, the result of mass-dependent fractionation due to the large relative difference in ⁶Li and ⁷Li masses (16%). The secondary mineral clay-Li fraction carried down rivers to the sea (presumably without dissolution), or left behind temporarily on the continents.

Low Weathering Intensity (weathering limited)

- Steep relief (steep grinding rivers, tectonics, uplift, mountains)
- Rapid physical weathering (high suspended solids in rivers)
- Incongruent weathering ([Cations]clay/[Cations]dissolved is high)
- Characteristic (today) of Himalayas, Andes (monsoons, orographic rain)
- Most of “weathered” lithium delivered to the sea is in 2° clays – [Li]clay
- Global average river lithium ratio today is 4 : 1 :: [Li]clay : [Li+]dissolved

High Weathering Intensity (transport limited; “supply”-limited of West et al., 2012)

- Peneplained continents (flat, sluggish rivers; no mountains in the rain belts)
- Low physical weathering (low suspended solids in rivers; very thick soils)
- Intense and congruent chemical weathering (little secondary clay formation; rock dissolution slow but homogeneous; residual alumina (gibbsite) and laterite (FeOx soils); [Cations]clay/[Cations]dissolved is low)
- Characteristic (today) of shields in hot, wet, humid tropics (equatorial rain belt, Inter-Tropical Convergence Zone [ITCZ] and Orinoco basin “Right Bank Guyana Shield” Rio Negro; very few examples)
- Most of “weathered” lithium delivered to sea is dissolved – [Li]dissolved
- Global average river ratio at 60 Ma was about 1 : 4 :: [Li]clay : [Li+]dissolved

Weathering intensity says nothing about river fluxes, which are dependent on orography (uplift) and ITCZ/monsoon runoff and rainfall. What matters is where the uplift and tectonics occur with respect to rainfall—the tropics.

in weathered horizons of soils, is isotopically light. The clay-Li portion delivered to the seabed is mixed with other solid components (products of physical weathering) carrying near-crustal isotope values (Δ7Li UCC = +1.7‰) that swamp the light clay-Li isotope signal in sediments. These inputs to the sea must be balanced on time scales of the Li residence time in the sea (~ 1 million years). Because Li is greatly enriched in secondary clays, and this enrichment carries light 6Li, most studies of the lithium isotope cycle have concluded that marine removal must occur via “reverse weathering” in the ocean (Mackenzie and Garrels, 1966; Aller, 2014)—the burial of Li via the re-formation of authigenic aluminosilicate clays on and in the seabed from diagenetic alteration of silica, continental clays, and detrital silicate debris (Al) and cations (especially Mg and Li) (Sayles, 1979, 1981; Mackin, 1986; Mackin and Aller, 1989; Michalopoulos and Aller, 1995, 2004; Michalopoulos et al., 2000; Aller, 2014). This removal must balance the inputs. Misra and Froelich (2012) lumped the two “known” sinks of reverse weathering into one sink: marine authigenic aluminosilicate clays (MAAC – the formation of authigenic clays in marine sediments) and altered oceanic crust (AOC – the “weathering” alteration of oceanic basalts by seawater circulation through the “low temperature” crust off axis). In both cases, the source of Li entering the MAAC and AOC diagenetic pathways is from seawater. These two processes are known to enrich light 6Li in authigenic aluminosilicates (Chan et al., 1992, 2006; Chan and Kastner, 2000). The steady-state balance requires a net removal of about 23 × 10⁹ moles per year at the Δ7Li Input value of +15‰ (Figure 5a; note that we ignore the small subduction reflux in Misra and Froelich [2012] for simplicity because it has little effect on the overall long-term balance and is very poorly understood). This removal (Δ7Li Output = +15‰) is 16‰ lighter than the seawater Li source (Δ7Li SW = +31‰). This 16‰ offset between seawater-Li and the diagenetic-Li sink pushes seawater Δ7Li SW to isotopically heavy values. Thus, about half of the present-day seawater Δ7Li SW offset from rocks (+31‰) is due to heavy inputs to the ocean from crustal sources (+15‰), and about half is due to light outputs from the ocean into reverse weathering sinks (~16‰). Changes in
input fluxes (weathering of rocks) or output fluxes (formation marine authigenic clays) or shifts in their isotopic values will force the seawater $\delta^7\text{Li}_{\text{SW}}$ to change. Such $\delta^7\text{Li}_{\text{SW}}$ changes provide clues to changes in the weathering and reverse weathering dynamics of the planet and its ocean.

**THE CENOZOIC HISTORY OF SEAWATER LITHIUM ISOTOPES**

Lithium incorporated into well-dated, cleaned foraminifera in marine sediments indicates that $\delta^7\text{Li}$ has not been constant through the Cenozoic (Hathorne and James, 2006; Misra and Froelich, 2012; Figure 4). Both of these records document a rapid rise of about 4‰ from about 19 Ma until 6 Ma (middle Miocene), after which values plateaued at today’s value. Misra and Froelich’s (2012) record back to 68 Ma shows Late Cretaceous values that are reminiscent of this 14–6 Ma rise with a drastic 5‰ drop at the 66 Ma KPg boundary at the end of the Cretaceous. The reasons for this KPg drop are not understood, but may be related to some dramatic and fast continental or oceanic reset of weathering or reverse weathering processes that are not discussed here. Within 6 Ma after the KPg boundary, $\delta^7\text{Li}_{\text{SW}}$ values decline to a Cenozoic low of about 22‰ in the late Paleocene and early Eocene (Figure 4). We argue that this post-KPg decrease was due at least in part to a slow (5–10 Ma) Paleocene disappearance of high-relief terrains.

Figure 5. (a) Present-day and (b) late Paleocene (60 Ma) depictions of steady-state lithium and lithium isotope fluxes to and from the ocean (modified after Misra and Froelich, 2012). Note that Li and West (in press) partition the reverse weathering sink into two components, marine authigenic aluminosilicate and altered oceanic crust, with different fractionation factors. See text for their alternative explanation of the Cenozoic rise in seawater $\delta^7\text{Li}$.
The low value at 60 Ma is about 9‰ below today’s seawater and reflects large changes in Li input or outputs, or both, compared to today. After the EECO at 52 Ma, δ7LiSW began rising in sporadic pulses. One upward shift in δ7LiSW at 51 Ma occurred after India (carrying the Deccan Traps) drifted northward across the equatorial Inter-Tropical Convergence Zone (ITCZ; Kent and Muttoni, 2008) and initiation of the collision of India with Southeast Asia. A second, at 40 Ma, occurred near the final closure of this suture and initiation of uplift of the Himalayas (Raymo and Ruddiman, 1992; Ruddiman et al., 1997; Bouihol et al., 2013). The rise of the Andes and the Rockies after about 20 Ma and 15 Ma coincides with the final upstics in the δ7LiSW record.

Coincidence does not prove causality, but the implications of the seawater δ7Li record are that (1) if the seawater δ7LiSW change is being driven by changes in continental weathering, and (2) if the Li isotope fractionation exhibited by clays is a general feature of Li transfer into authigenic aluminosilicates, then the Cenozoic δ7LiSW record is most easily interpreted as one of increasing relief and uplift, decreasing continental weathering intensity leading to more incongruent weathering, more cations in clays, and faster denudation in the Late Cenozoic. The early Eocene must have been largely deficient in these features.

**THE GLOBAL LITHIUM ISOTOPE CYCLE—60 MILLION YEARS AGO**

A possible lithium isotope balance for the Paleocene/Eocene boundary ocean at 60 Ma is one in which the entire δ7LiSW change (9‰) is delegated to the continental weathering source—all else remaining constant, including invariant hydrothermal fluxes (Rowley, 2002) and lithium isotope fractionation factors (Misra and Froelich, 2012; Figure 5b).

In this simple end-member scenario, which is unlikely to be the whole answer, continental rivers must exhibit almost congruent weathering with very high weathering intensity (δ7LiRivers = +3‰), penexplained low relief continents in the tropical rainfall belts, low physical weathering with low suspended solids in rivers, and low dissolved-Li to clay-Li ratios (Box 2). This is consistent with age-, subduction-, and hiatus-loss corrected seafloor drill core records for Cenozoic Deep-Sea Drilling Project and Ocean Drilling Program sites showing that terrestrial sediment accumulation rates on the seafloor, which must reflect global denudation rates of detrital material to the seabed, increased dramatically in the Neogene (Peizhen et al., 2001; Hay et al., 2002; Molnar, 2004; Herman et al., 2013, and references therein). In contrast, the continental delivery of material to the Paleogene sea consisted mostly of dissolved chemicals, with minimal solid detritus. The Eocene seafloor record shows large deposits of cherts (biogenic opal), phosphorites, organics, and a shallow CCD with shallow-water (reefal) carbonates (Brewster, 1980; Pälike et al., 2012). Paleogene ocean Mg/Ca ratios were a factor of three to five times lower than today (Holland, 1978; Coggon et al., 2010; Rausch et al., 2013), suggesting that the continental delivery of seawater ions (Mg, Li) via weathering and their loss to the seafloor via reverse weathering were both reduced compared to today.

Li and West (in press) make a good case for Cenozoic changes in both continental weathering and in seafloor reverse weathering to explain the Cenozoic δ7LiSW records. In their model, they partition the reverse weathering sink into explicit and separate AOC and MAAC components and apply different temperature-dependent fractionation factors to each. For the MAAC sediment component, they adopt a lithium isotope fractionation offset from seawater of ΔSW-MAAC = 20‰. For the AOC component, they adopt a Li-fractionation factor based on expected warm temperatures in oceanic crust undergoing alteration of ΔSW-AOC = 13‰ (about 80°C; Chan et al., 2002). This 7‰ spread in the two components of ΔRev-Wx is in contrast to the single value for reverse weathering of 16‰ adopted by Misra and Froelich (2012). Then, they scale the continental Li flux to the Cenozoic CO2/ocean dissolved inorganic carbon model of Li and Elderfield (2013) to estimate continental weathering fluxes—these increased fluxes explain about half the oceanic δ7LiSW rise of the past 60 Ma. They can balance the other half of the ocean rise in δ7LiRivers by scaling an increase in the MAAC reverse weathering sink to increased continental weathered detritus (Al-source) delivered to the seafloor by the increased physical denudation of the continents derived from their weathering model. They find that to explain the full change in Cenozoic seawater δ7LiSW (9‰), their model δ7LiRivers must increase from a low of 12.2‰ at 60 Ma to the present-day value of 23.5‰, an increase of about 11‰. This is about half of the change in continental weathering explored by the end-member scenario of Misra and Froelich (2012) and avoids their extremely high weathering intensity at the Paleocene/Eocene boundary. More importantly, the Li and West (in press) result would increase weathering fluxes of continental dissolved lithium fluxes (rivers) by a factor of about 3.4 and the burial flux of MAAC on the seafloor by a factor of about 4.1. These changes reflect...
dramatic shifts in continental weathering and the delivery of detritus and cation dissolved-Li and clay-Li to the sea over the Cenozoic. We estimate from their model that the clay-Li to dissolved-Li ratio of continental rivers increased from 1:2 at 60 Ma ago to 4:1 at present (cf. Box 2), so that if the dissolved Li-flux increased by a factor of 3.4, the clay-Li delivery flux to the ocean rose by a factor of 14, and the total lithium weathered from igneous rocks on the continents rose about a factor of 17.

Clearly, if these changes in continental lithium fluxes scale in any manner to other cations weathered from the continents (especially Mg), then the consumption of CO₂ by silicate weathering and the release of cations to the sea increased dramatically over the Cenozoic, implying the absence of these factors driven by uplift and high relief in the early Eocene.

CONCLUDING REMARKS

The early Cenozoic climatic optimum was a result of increased supply of carbon dioxide to the ocean-atmosphere system as well as diminished removal of carbon dioxide from the atmosphere through weathering of silicate rocks. The scarcity of newly uplifted, fresh, weatherable rocks in the hothouse climate led to a slowdown of the negative feedback mechanism of the Urey-Walker-Berner cycle and promoted a temporary run-away increase in the CO₂ concentration of the ocean-atmosphere system on the post-KPg Earth. The absence of evidence from seawater ⁸⁷/⁸⁶Sr and δ⁷Li records to support increased continental weathering rates in this high-CO₂ hothouse world indicates that the limiting ingredient was igneous silicates undergoing weathering. While this Paleocene-Eocene Earth was probably not flat, evidence from the ocean suggests that despite global high temperatures, high CO₂, and high rainfall in the tropics (all of which should accelerate silicate weathering and promote removal of CO₂), the physical and chemical weathering rates of the continents were subdued because of low uplift rates. CO₂ consumption and cation release by silicate weathering was lower than today. Fluxes of dissolved cations, continental detritus, and secondary detrital clays to the seabed were also significantly lower than today. Formation of authigenic aluminosilicate clays via reverse weathering in marine sediments was limited. This "absence of a CO₂ sink" hypothesis might help explain how the hot Eocene climate persisted for tens of millions of years.

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