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Co-variation of silicate, carbonate and sulfide weathering drives CO₂ release with erosion

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Global climate is thought to be modulated by the supply of minerals to Earth's surface. Whereas silicate weathering removes carbon dioxide (CO₂) from the atmosphere, weathering of accessory carbonate and sulfide minerals is a geologically relevant source of CO₂. Although these weathering pathways commonly operate side by side, we lack quantitative constraints on their co-variation across erosion rate gradients. Here we use stream-water chemistry across an erosion rate gradient of three orders of magnitude in shales and sandstones of southern Taiwan, and find that sulfide and carbonate weathering rates rise with increasing erosion, while silicate weathering rates remain steady. As a result, on timescales shorter than marine sulfide compensation (approximately 10⁶–10⁷ years), weathering in rapidly eroding terrain leads to net CO₂ emission rates that are at least twice as fast as CO₂ sequestration rates in slow-eroding terrain. We propose that these weathering reactions are linked and that sulfuric acid generated from sulfide oxidation boosts carbonate solubility, whereas silicate weathering kinetics remain unaffected, possibly due to efficient buffering of the pH. We expect that these patterns are broadly applicable to many Cenozoic mountain ranges that expose marine metasediments.

Silicate weathering is a key component of Earth's long-term carbon cycle. It neutralizes CO₂ emissions from the crust and mantle by producing alkalinity, thereby driving precipitation of carbonates in the oceans¹. Weathering rates are controlled by temperature, the availability of weathering fluids and the reactive mineral surface area². They can be limited either by the supply of minerals (supply limit)^{3–5}, by the kinetics of weathering reactions (kinetic limit)^{3,4} or by the availability of acidic fluids (equilibrium limit; see Table 1 for definitions)⁵. As a result, an intimate link emerges between chemical weathering, the supply of fresh rocks by physical erosion and climate^{6–9}. The sensitivity of weathering to erosion rates underpins the hypothesis that major orogenic events can lead to increased silicate weathering rates, which decreases atmospheric CO₂ concentrations and cools the planet^{7,10–12}.

Most Cenozoic orogens do not exhume pure silicate rocks, but instead expose lithologies, such as shallow marine metasediments, that contain accessory carbonate and sulfide minerals. Recent studies found that weathering of these accessory minerals complicates the simple link between the supply of fresh rocks by erosion, enhanced weathering rates and drawdown of atmospheric CO₂. Importantly, sulfuric acid that is produced during the oxidation of sulfide minerals, such as pyrite, can titrate alkalinity^{13,14}. Therefore, silicate weathering by sulfuric acid does not contribute to CO₂ drawdown, whereas carbonate weathering by sulfuric acid releases CO₂^{13,14}. Because carbonate and sulfide minerals dissolve up to three orders of magnitude faster than silicates^{2,15,16}, they frequently dominate weathering fluxes, even where they constitute only minor proportions of the exposed lithologies^{12–14,17,18}. Moreover, sulfide weathering rates increase with erosion, and the consequent titration of alkalinity has been proposed to compensate for increases in silicate weathering rates during mountain building^{12–14,19}.

Based on these studies, existing global carbon cycle models assume that the weathering rates of all mineral phases, including sulfides, carbonates and silicates, increase with erosion and proceed independently of each other^{1,9,10,20}. However, where these weathering processes operate side by side, the acidity produced by sulfide oxidation may modulate silicate and carbonate weathering rates, with consequences for the carbon cycle^{13,14,21,22}. Critically, we lack quantitative constraints on silicate, carbonate and sulfide weathering rates in parallel across extensive and continuous erosion rate gradients without accompanying variations in runoff, temperature and lithology. In existing compilations where erosion rate and runoff are correlated^{4,8,18}, unravelling the impact of mineral supply on weathering rates is challenging, because runoff impacts weathering^{23,24}. As a consequence, it remains unclear how weathering of accessory sulfide and carbonate minerals co-evolve across erosion rate gradients, and how the supply of fresh minerals during mountain building impacts the balance of CO₂ emission and sequestration and, ultimately, Earth's climate.

Here we present water chemistry data from streams in southern Taiwan that span an erosion rate gradient of nearly three orders of magnitude in a relatively homogeneous substrate with minimal variations in runoff. These data allow us, for the first time to our knowledge, to quantify the impact of erosion rates—independent of lithology and runoff—on the concomitant variation of carbonate, silicate and sulfide weathering and, thereby, on the sequestration and emission of CO₂ from these weathering reactions.

Water chemistry in southern Taiwan

We present 103 water samples from 40 separate catchments that span a striking gradient of northward-increasing relief, channel steepness and erosion rates^{25–27} (Fig. 1). Erosion rate estimates from cosmogenic nuclide data are available for a subset of the sampled

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Table 1 | Definitions of weathering limits

Terminology	Definition
Equilibrium limit	Weathering rates limited by the saturation of weathering fluids. Terms used for this limit elsewhere include 'thermodynamic limit' or 'transport limit'. The latter describes how weathering is limited by the transport of weathering products away from the reaction front (which leads to saturated solutions).
Kinetic limit	Weathering rates limited by the kinetics of the forward reaction of mineral dissolution (in undersaturated solutions).
(Mineral) supply limit	Weathering rate limited by the supply of fresh mineral surfaces. Some authors have used the term 'transport limit', referring to the transport of minerals to the weathering zone. However, a 'mineral transport limit' and a 'fluid transport limit' (see equilibrium limit above) hold different implications for the relationship between erosion rates and weathering rates and are therefore distinguished here.

catchments^{26,27}. These ¹⁰Be erosion rate data are well correlated with the normalized catchment-averaged steepness index (k_{sn}), and we used a regression to predict the erosion rate for each sample from its catchment-averaged k_{sn} estimate (Extended Data Fig. 1). Results show a northward increase of inferred erosion rates from <0.004 to >2 mm yr⁻¹. In contrast to this erosion rate gradient of three orders of magnitude, average annual rainfall and inferred runoff vary by no more than a factor of three within the study area^{28,29}. The bedrock in the sampled catchments is dominated by mildly metamorphosed calcareous argillites and shales of the Cenozoic slate belt³⁰ (Fig. 1). In the northernmost part of the study area, these rocks unconformably overlie high-grade metapelites, whereas the southernmost tip of the island is characterized by alternations of sandstones and shales of the Mutan formation³⁰. Minor carbonate is found as interstitial cement, clastic grains or in veins, in all formations of the study area, and pyrite occurs as a minor rock building mineral³⁰ (Supplementary Text 1).

For all samples, we measured concentrations of major dissolved cations, anions and silica. Across the erosion rate gradient, we find an increase in the concentrations of Ca²⁺ and SO₄²⁻ ions (Fig. 2a), whereas Na⁺ and dissolved silica show constant concentrations at low erosion rates and a decrease at high erosion rates (Fig. 2b). The concentrations of Mg²⁺ and K⁺ do not change systematically (Fig. 2c). We used an inverse approach to estimate the cation concentrations contributed from silicates ([Cat]_{sil}; Fig. 2d), carbonates ([Cat]_{carb}; Fig. 2e), hot springs and cyclic sources (Extended Data Fig. 2), as well as the fraction of the total cation charge derived from carbonate weathering (f_{carb} ; Fig. 2f). We further estimated the concentration of sulfates contributed from pyrite oxidation ([SO₄]_w; Fig. 2e), the fraction of weathering by sulfuric acid (f_{sulf} ; Fig. 2f) and the concentration of Ca²⁺ lost to secondary precipitation of carbonate (Fig. 2d). Inputs from evaporites and anthropogenic pollution are considered negligible (Methods).

Based on the inversion, weathering inputs dominate the solute load and contribute a median of ~88% (first quartile 74%, third quartile 95%) of the total cation budget, whereas the remaining solutes are related to cyclic and hot spring inputs. We find that silicate and carbonate cation concentrations are similar ($f_{carb} \approx 0.5$) for erosion rates <0.1 mm yr⁻¹ (Fig. 2f). At higher erosion rates, [Cat]_{carb} increases five- to tenfold, whereas [Cat]_{sil} remains constant or even declines, resulting in a dominance of carbonate weathering ($f_{carb} = 0.8-1$; Fig. 2d-f). In turn, sulfate concentrations from sulfuric acid weathering increase monotonically by more

than 40-fold from <50 to $>2,000$ μmol l⁻¹ as erosion increases (Fig. 2e). Consequently, f_{sulf} increases from near 0% to about 50% from low to high erosion rates (Fig. 2f).

Silicate, carbonate and sulfide weathering

Using our new water chemistry data, we investigate the limits to weathering of sulfide, carbonate and silicate phases in parallel across a single erosion rate gradient. Because runoff is inferred to vary by less than two- to threefold across the erosion rate gradient^{28,29}, we interpret the solute concentrations to be proportional to weathering rates (within a factor of two to three). The steady, 40-fold increase in concentrations of sulfates (Fig. 2e) suggests that pyrite weathering is limited by the supply of fresh minerals to the weathering zone (supply limit). Such supply limitation is consistent with previous studies¹⁴ and with the observed depletion of pyrite in the deepest parts of the weathering zone^{31,32}. The supply of sulfuric acid by sulfide oxidation has been hypothesized to either increase both carbonate and silicate weathering rates¹⁴ or to boost only carbonate weathering¹³. Our observations suggest that carbonate weathering rates are closely associated with sulfuric acid addition, whereas silicate weathering rates are decoupled (Fig. 3a).

Carbonate weathering in Taiwan seems to be limited by the saturation of weathering fluids (equilibrium limit), based on several observations that include supersaturation of stream waters with respect to calcium carbonate (Extended Data Fig. 3), secondary carbonate precipitates around springs (see Fig. 2 in ref. ³³) and the presence of detrital carbonate in streams. Yet, despite this equilibrium limit, [Cat]_{carb} increases across the erosion rate gradient (Fig. 2d,f). This key observation implies that the solubility of carbonates must increase with erosion. A northward trend of cooling water temperatures from $\sim 30 \pm 5$ to $\sim 23 \pm 4$ °C can only account for a <1.7 -fold increase in [Ca²⁺], much lower than the observed 5- to 10-fold increase. In contrast, acid addition can increase carbonate solubility by an order of magnitude per unit pH decrease (Supplementary Text 2). Hence, based on the close association of sulfate concentrations with [Cat]_{carb} (Fig. 3a), we propose that northward-increasing acid availability is the main driver of the solubility increase. We estimate the pH at which the sampled supersaturated river waters would be at saturation with calcium carbonate (pH_{eq})—a maximum estimate of the pH in the subsurface (Fig. 3b). This inferred equilibrium pH of the weathering fluids generally decreases with increasing erosion rates, mirroring the pattern of [Cat]_{carb} (Fig. 3b). We find that this observed trend in pH_{eq} can be explained to a large extent by an estimate of pH_{SO₄}, which we define as the pH that results from the addition of sulfuric acid to a carbonate-buffered solution (Fig. 3c). The deviation from the 1:1 line between pH_{eq} and pH_{SO₄} implies that we either overestimate the loss of carbonates to secondary precipitation, or that the supply of another source of acidity, for example carbonic acid, also increases with erosion rates. In either case, carbonate weathering increases substantially with erosion and appears strongly coupled to sulfuric acid supply.

In contrast to carbonate weathering, cation concentrations from silicate weathering do not increase or may even decline at higher erosion rates (Fig. 2e). An equilibrium limit whereby silicates are saturated is unlikely, because, similar to carbonates, the decrease of the solution pH by at least one unit (Fig. 3b) is expected to increase the solubility of silicates by a factor of three to ten³⁴. Hence, silicate weathering in the study area appears limited across almost the entire erosion rate gradient of three orders of magnitude wide by the slow kinetics of silica dissolution³⁴. This interpretation implies that silicate weathering kinetics are constant or declining from south to north. Northward-decreasing temperatures may explain a slowing of weathering kinetics by a factor of $\sim 1.5-2.5$ (Supplementary Text 2). In addition, the potential two- to threefold increase in runoff could dilute major cations, although the dilution is probably less than a factor of two to three (see Supplementary Text 2). In contrast,

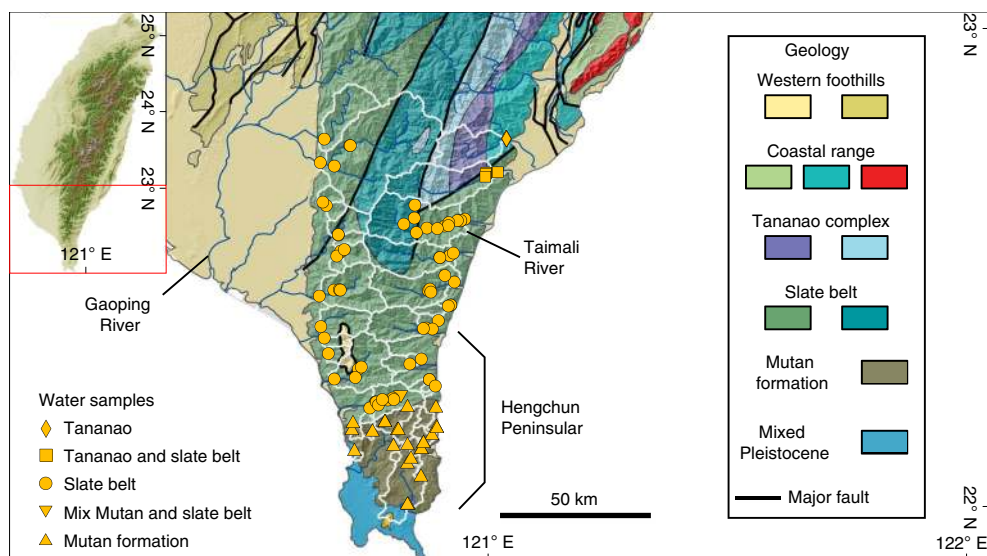


Fig. 1 | Lithologic map of southern Taiwan and sampling locations. Map of the southern tip of Taiwan showing all sampling locations. Major lithological units and structures are modified after refs. ^{30,50}. Symbols for water samples indicate dominant lithology outcropping in the catchment. Light grey lines mark the 40 unique catchments sampled here. Two rivers mentioned in the main text are marked.

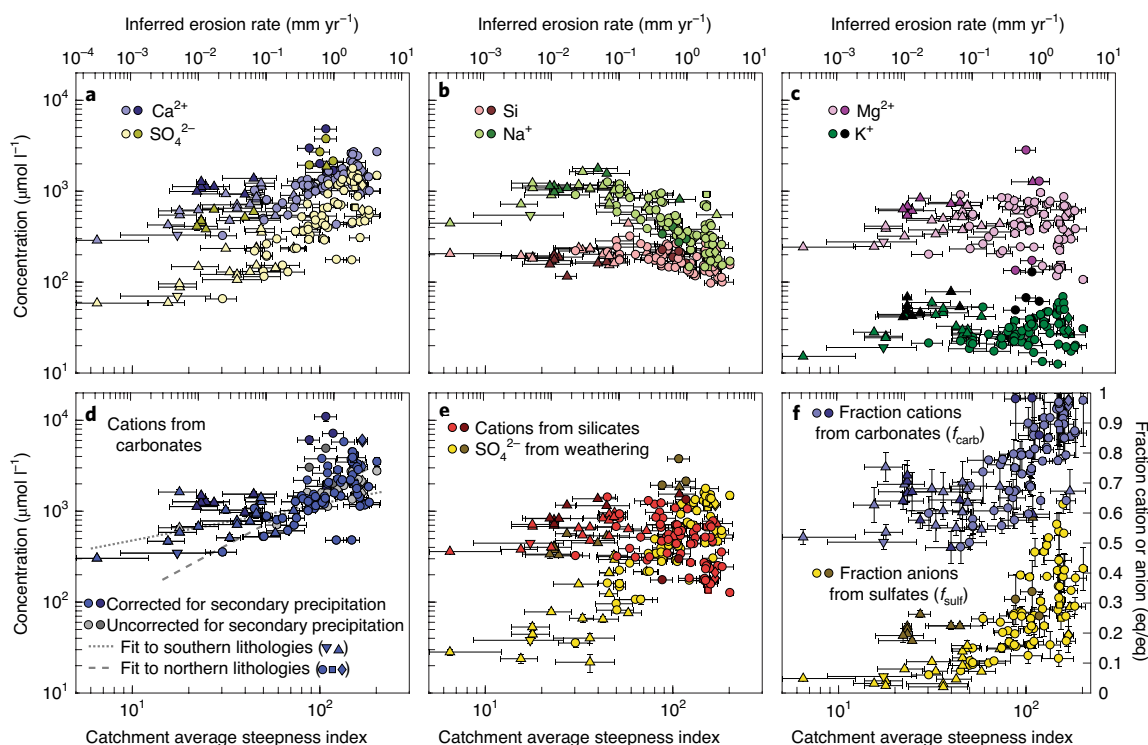


Fig. 2 | Stream-water chemistry across the erosion rate gradient. **a–e.** Concentrations of calcium and sulfate (**a**), silica and sodium (**b**), magnesium and potassium (**c**), cations from carbonate weathering (**d**) and cations from silicate weathering and sulfate from sulfide oxidation (**e**). **f.** Fraction of cation charge contributed from carbonate weathering and fraction of cation charge balanced by sulfate. All results are plotted against the catchment-averaged steepness index (lower x axis) and inferred erosion rates (upper x axis). Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval. Samples with anomalously high sulfate concentrations are marked as darker points.

silicate weathering kinetics can be strongly modulated by the solution pH and increase by up to an order of magnitude per unit of pH change, both with increasing acidity (at pH <5–6) and with increasing alkalinity (at pH >8–9)^{3,35}. Such changes would more than compensate for the potential impact of temperatures or runoff.

However, at circumneutral pH values of ~6–8 the dissolution rate of feldspars is at a minimum and insensitive to pH changes³⁵. The observed invariance or even slight decline of silicate weathering kinetics despite a 40-fold increase in the supply of sulfuric acid (Fig. 2e) is, therefore, notable. The subsurface pH appears to be

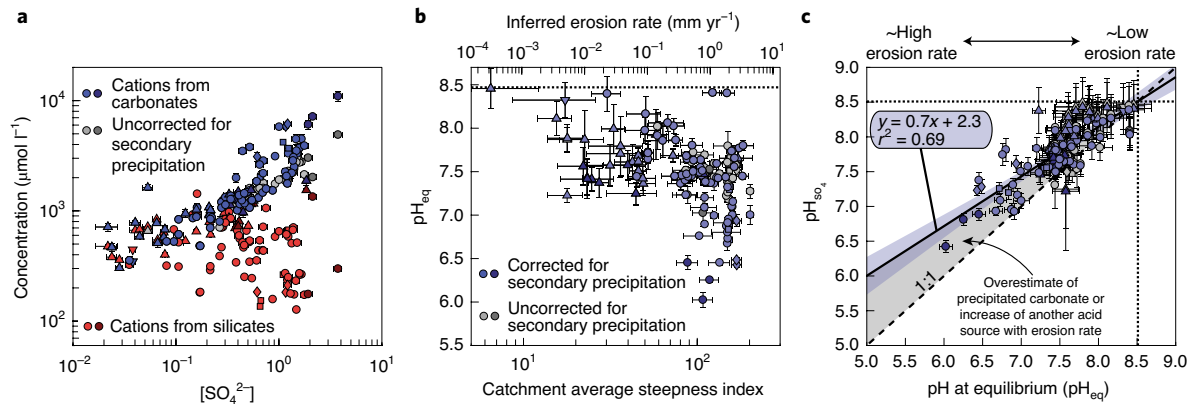


Fig. 3 | Link between sulfuric acid, weathering and pH. **a**, Co-variation of cation concentrations with sulfate. **b**, Predicted maximum pH in the weathering zone (pH_{eq}) against catchment-averaged steepness index (lower x axis) and inferred erosion rates (upper x axis). **c**, Predicted pH for a carbonate-buffered solution after addition of sulfuric acid (pH_{SO_4}) against the equilibrium pH. The solid line shows regression with 95% confidence bands; the dotted lines show initial pH before addition of sulfuric acid. Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval. Samples with anomalously high sulfate concentrations are marked as darker points.

sufficiently buffered to within circumneutral values to prevent any substantial increase in silicate weathering kinetics by the addition of sulfuric acid. On the contrary, addition of sulfuric acid may even slightly decrease weathering rates by pushing the pH from alkaline ($\text{pH} \sim 8.5$) to circumneutral. Such buffering of the pH is consistent with subsurface pH values inferred from carbonate equilibria (Fig. 3b), and it provides a compelling explanation for the differential coupling of silicate and carbonate weathering to sulfide oxidation (Fig. 3a).

By observing the weathering of sulfide, silicate and carbonate across a single erosion rate gradient, we find that rock mass weathering does not have a universal limit. Rather, across a large range of erosion rates in the mountains of Taiwan, silicate weathering is kinetically limited, carbonate weathering is equilibrium limited and sulfide weathering is supply limited. Moreover, our data suggest that the supply of sulfuric acid does not increase weathering rates of both silicates and carbonates as previously hypothesized¹⁴, even though sulfuric acid probably contributes to weathering of both phases²². Instead, we propose that in metasediments with sufficient buffering capacity, the supply of sulfuric acid increases the equilibrium limit of carbonates, whereas buffering of weathering fluids at circumneutral pH prevents an increase in silicate weathering rates. The details of this coupling depend on the relative positions of weathering fronts and the distribution of mineral phases in the subsurface³², but the dominance of carbonate weathering, especially at high erosion rates (Fig. 2), implies that most sulfuric acid must be buffered by carbonates, as previously described for shale weathering³⁶.

The observed trends continue across the lithologic boundary between the more sand-rich Mutan formation and the more shale-rich slate belt (Fig. 1). Lithology may cause a change in the trend between carbonate weathering rates and erosion rates (Fig. 2d), and local lithologic variations may lead to anomalously high sulfate concentrations of some samples. Nevertheless, our observations also hold when only catchments in the slate belt are considered (circles in all figures; spanning a ~ 200 -fold range of estimated erosion rates). The similarity of samples collected under different hydrological conditions across two field seasons—a dry winter and a wet spring (Extended Data Fig. 4)—suggests that the broad patterns we observe hold across variations in discharge. Moreover, the chemistry of many streams is relatively insensitive to short-term variations in runoff^{24,37}, and Taiwanese catchments typically show dilution by factors of less than five across discharge ranges of several orders of magnitude^{38,39}. In apparent contrast to our findings,

existing global compilations of river chemistry often find a positive link between silicate weathering fluxes (that is, the concentration multiplied by runoff) and erosion^{4,8}. However, these positive correlations are frequently dominated by a strong co-variation of runoff and erosion rate. Because runoff can directly affect weathering rates²³, this co-variation obscures the link between mineral supply and chemical weathering. Hence, a direct comparison between these data and our observations is difficult.

Impacts for Earth's carbon cycle

Existing carbon cycle models typically assume that the reactivities of all mineral phases increase with rates of erosion and mineral supply, and evolve independently of each other^{1,9,10,20}. As a consequence, decreasing atmospheric CO_2 concentrations during the Cenozoic era have been linked to increased silicate weathering rates with uplift and erosion of the Alpine–Himalayan system^{7,10,11}. In contrast, our data suggest that in the absence of a strong co-variation of mineral supply and runoff, increased erosion of marine siliciclastic sediment sequences may lead to a constant or even decreasing reactivity of silicate minerals, whereas carbonate and sulfide reaction rates increase (Fig. 2). We postulate that this co-variation of silicate, carbonate and sulfide weathering rates applies to erosion rate gradients in many active mountain ranges. Catchments in southern Taiwan are underlain by (meta)sedimentary rocks of an inverted passive margin sequence that is typical of rocks uplifted along active orogens. Moreover, solutes from carbonates and sulfides dominate total weathering budgets from freshly exposed metasediments in the Southern Alps of New Zealand¹⁸, the Himalaya¹⁷, the Rocky Mountains¹⁴, the Andes¹³ and even in slower-eroding continental interior settings such as northern Texas⁴⁰. Finally, it is likely that sulfuric acid addition acts to increase silicate weathering rates only in rare cases, where acid production exceeds the buffering capacity of the sediment, for example in acid mine drainage^{41,42}, or in headwater streams draining pure silicate lithologies⁴³.

Relative changes in sulfide, silicate and carbonate weathering rates modulate the sequestration and release of CO_2 on millennial to multi-million-year timescales¹³. In particular, the increase of sulfide weathering relative to silicate dissolution as erosion rates increase (Figs. 2 and 3) results in a clear shift from reactions that sequester CO_2 to reactions that release CO_2 (Fig. 4). In southern Taiwan, this shift occurs at erosion rates of ~ 0.3 – 0.8 mm yr^{-1} (Fig. 4). Importantly, CO_2 release rates at erosion rates $> 1 \text{ mm yr}^{-1}$ are substantially higher than CO_2 sequestration rates at erosion

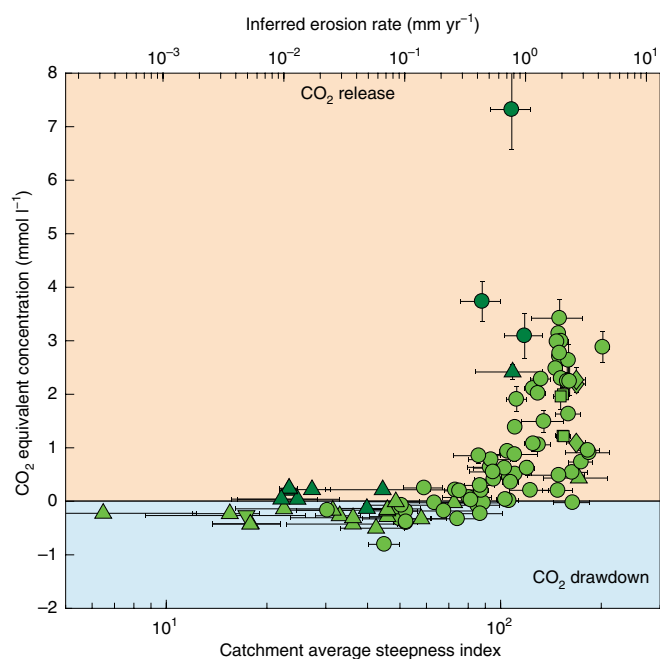


Fig. 4 | Impact of erosion on the long-term carbon cycle. Moles of CO₂ emitted or sequestered per volume of water against catchment-averaged steepness index (lower x axis) and inferred erosion rates (upper x axis). Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval. Samples with anomalously high sulfate concentrations are marked as darker points.

rates of $<0.1 \text{ mm yr}^{-1}$ (Fig. 4). This contrast between CO₂ release and sequestration may be even more pronounced where runoff increases in parallel with erosion rates^{4,8,44}. As a result, the total CO₂ release from mountain building may overwhelm CO₂ drawdown from more slowly eroding landscapes, even if areas of high erosion rates do not cover the majority of exposed land area⁴⁵.

Ultimately, sulfuric acid generation by sulfide oxidation is balanced by marine sulfate reduction, although the timescale for achieving such a balance probably exceeds 10^6 years⁴⁶. Thus, the shift from CO₂ sequestration to CO₂ emission with increasing erosion rates (Fig. 4) applies to timescales between those of carbonate compensation ($\sim 10^3$ – 10^4 years) and pyrite neof ormation in marine sediments ($\sim 10^6$ – 10^7 years). These are the timescales of long-term climate change and mountain building. Even beyond the timescales of sulfide compensation in the ocean, the rate of CO₂ sequestration from silicate weathering may be largely unaffected by mineral supply (Extended Data Fig. 5), because the concentration of solutes from silicate weathering does not increase across a large range of erosion rates (Fig. 2e). If our findings extend to metasediments globally, the Cenozoic decrease of atmospheric CO₂ concentrations cannot be associated with an increased supply of minerals within inverted passive margins and must, instead, be linked to increased organic carbon burial⁴⁷, increased relief and orographic precipitation^{33,44}, alkalinity generation in floodplains⁴⁸ or weathering of fertile igneous silicates⁴⁹. Over the course of Earth history, the growing accumulation of carbonaceous shelf sediments on continental crust may thus have shifted orogenesis from a net sink of CO₂ to a net source of CO₂.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of

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Methods

Notations and constants. We use the following notations (Extended Data Table 1): $[i]$ is the concentration of ion i , v_i is the valence of ion i , $[i]^{eq} = v_i [i]$ is the charge equivalent concentration of ion i , γ_i is the activity coefficient of ion i and a_i is the activity of ion i with $a_i = \gamma_i [i]$. Subscripts carb, sil and sulf denote quantities calculated for the ions derived from carbonate, silicate and sulfide weathering, respectively, and the subscripts hs, cy, sc, m and wtot refer to contributions from hotspots, cyclic sources, silicate and carbonate weathering, and to the total measured concentrations and the solutes from weathering reactions, respectively. We use the equilibrium constants and standard enthalpies as specified in Extended Data Table 2.

Water sampling and measurements. We collected 119 water samples across southern Taiwan for chemical analysis in 1 litre high-density polyethylene bottles from river banks with access to the main flow. The transition from colluvial to fluvial processes typically occurs around ~ 1 km² of drainage area in Taiwanese mountain catchments⁵¹. Because a scaling of erosion rates and catchment-averaged steepness indices is only sensible within the fluvial regime, we limit our analysis to 104 samples with catchment areas of > 2 km². All samples were filtered the evening after sampling into a set of smaller high-density polyethylene bottles (for anion and cation measurements) using 0.22 μ m Merck Express Plus membrane filters. We rinsed each bottle with filtered water before filling, and we acidified all bottles for cation analyses with concentrated ultrapure nitric acid (HNO₃). We measured pH and temperature of the river in the field at the sample location using a WTW Multi 3430 multimeter and a WTW SensoLyt 900 pH probe. Electrical conductivity and dissolved oxygen content of the water were measured with a WTW TetraCon 925 and a WTW FDO 925 probe, respectively. After dilution with a caesium-spiked solution, cations (Ba²⁺, Ca²⁺, Fe²⁺, K⁺, Li⁺, Mg²⁺, Mn²⁺, Na⁺ and Sr²⁺) and dissolved silica were measured on a Varian 720 ICP-OES at the GFZ. Every ten samples, a quality control sample mixed at the GFZ was measured to monitor machine drift. All accepted runs had a drift of $< 5\%$. We measured external river water standards SLRS-5 or SLRS-6 and USGS-T187 and M212 for quality control. The measurements were calibrated using a set of 11 in-house standards, and any standards that did not fall within 10% of a linear fit through all standards were discarded. We used measurements only within the range of accepted standards, and estimated the uncertainty in the cation analysis from the maximum deviation of the calibration standards from the calibration line. We measured major anions (F⁻, Cl⁻, NO₃⁻ and SO₄²⁻) on a Dionex ICS-1100 chromatograph using a six-point linear calibration and USGS-206 and USGS-212 standards for quality control. Uncertainty estimates were based on the standard deviation from three repeat measurements. Bicarbonate concentrations (HCO₃⁻) were estimated by charge balance, which assumes that the contribution of bicarbonate from organic sources is negligible (Supplementary Text 4). Uncertainties were propagated from the analytical uncertainties of the measured concentrations. All raw measurements are reported in Supplementary Data 1.

Erosion rate– k_n scaling. Catchment-averaged erosion rates are commonly well correlated with channel steepness, but the relationship has to be calibrated locally⁵². We used a regression between existing cosmogenic nuclide erosion rate data^{26,27} (Extended Data Fig. 1 and Supplementary Data 2) and the catchment-averaged steepness indices for all catchments with drainage areas of > 2 km² to predict catchment-averaged erosion rates for each of our water samples. First, we calculated mean steepness indices upstream of each cosmogenic nuclide sample and upstream of each water sample using TopoToolbox v2.2⁵³. Cosmogenic nuclide concentrations in southern Taiwan decrease northward, and concentrations in repeat samples collected in 2006, 2015 and 2016 generally agree^{26,27}. We used a regression through the catchment-averaged normalized steepness and catchment-averaged erosion rates to predict, from the steepness index, a catchment-averaged erosion rate for each of the water samples collected in this study (Extended Data Fig. 1). Uncertainties in erosion rates were estimated from the confidence band of the regression. ¹⁰Be concentrations in samples collected in 2012 tend to be relatively lower and more scattered than during other years²⁷. The decrease in ¹⁰Be concentrations of the 2012 samples was interpreted to result from extensive landsliding triggered by Typhoon Morakot in 2009²⁷, and we excluded these data from the analysis (blue points in Extended Data Fig. 1). Our water samples extend farther south than existing cosmogenic nuclide data. Therefore, all erosion rates < 0.06 mm yr⁻¹ were extrapolated from the regression (Supplementary Text 4).

Unmixing major contributions to the cation solute budget. We consider carbonates, silicates, cyclic water and hotspots as major contributors to the cation budget (see endmembers in Extended Data Table 3 and Supplementary Text 3). No evaporite deposits have been reported in Taiwan^{30,54}, and stream waters were collected upstream of major anthropogenically managed areas. In our analysis, we considered only the major cations Ca²⁺, Mg²⁺, K⁺ and Na⁺. The concentrations of cations derived from carbonate weathering, $[\text{Cat}]_{\text{carb}}$, silicate weathering, $[\text{Cat}]_{\text{sil}}$

the sum of silicate and carbonate weathering $[\text{Cat}]_{\text{wtot}}$, cyclic contributions $[\text{Cat}]_{\text{cy}}$ and hotspots $[\text{Cat}]_{\text{hs}}$ are:

$$\begin{aligned} [\text{Cat}]_{\text{carb}} &= \sum_i [i]_{\text{carb}}; [\text{Cat}]_{\text{sil}} = \sum_i [i]_{\text{sil}}; [\text{Cat}]_{\text{wtot}} \\ &= [\text{Cat}]_{\text{sil}} + [\text{Cat}]_{\text{carb}}; [\text{Cat}]_{\text{cy}} = \sum_i [i]_{\text{cy}}; [\text{Cat}]_{\text{hs}} \\ &= \sum_i [i]_{\text{hs}}; \end{aligned} \quad (1)$$

We define corresponding charge equivalent concentrations, $[\text{Cat}]^{\text{eq}}$, as:

$$\begin{aligned} [\text{Cat}]_{\text{carb}}^{\text{eq}} &= \sum_i v_i [i]_{\text{carb}}; [\text{Cat}]_{\text{sil}}^{\text{eq}} = \sum_i v_i [i]_{\text{sil}}; [\text{Cat}]_{\text{wtot}}^{\text{eq}} \\ &= [\text{Cat}]_{\text{sil}}^{\text{eq}} + [\text{Cat}]_{\text{carb}}^{\text{eq}}; [\text{Cat}]_{\text{cy}}^{\text{eq}} = \sum_i v_i [i]_{\text{cy}}; [\text{Cat}]_{\text{hs}}^{\text{eq}} \\ &= \sum_i v_i [i]_{\text{hs}}; \end{aligned} \quad (2)$$

The fraction of cation charge related to carbonate weathering, f_{carb} , and the fraction of cation charge balanced by sulfate, f_{sulf} are defined as:

$$f_{\text{carb}} = \frac{[\text{Cat}]_{\text{carb}}^{\text{eq}}}{[\text{Cat}]_{\text{wtot}}^{\text{eq}}}; f_{\text{sulf}} = \frac{[\text{SO}_4^{2-}]_{\text{sulf}}^{\text{eq}}}{[\text{Cat}]_{\text{wtot}}^{\text{eq}}} \quad (3)$$

We used an inverse approach to estimate the contribution of carbonate and silicate minerals to the dissolved solids. Following previous authors^{6,8,33,37,55}, this estimate was based on modelling the relative concentrations of three major soluble cations, Na⁺, Ca²⁺ and Mg²⁺, as a mix of a silicate and a carbonate endmember (Extended Data Table 3). In addition, here we include the concentration of Cl⁻ to allow correction of the cyclic and hotspots contribution (Extended Data Table 3). Based on these endmembers, we solved the following equations using the linear least squares solver lsqin in MATLAB:

$$\begin{aligned} \left[\frac{\text{Na}^+}{\text{Ca}^{2+}} \right]_{\text{m}} &= \alpha_{\text{Ca,sil}} \left[\frac{\text{Na}^+}{\text{Ca}^{2+}} \right]_{\text{sil}} + \alpha_{\text{Ca,carb}} \left[\frac{\text{Na}^+}{\text{Ca}^{2+}} \right]_{\text{carb}} + \alpha_{\text{Ca,cy}} \left[\frac{\text{Na}^+}{\text{Ca}^{2+}} \right]_{\text{cy}} \\ &+ \alpha_{\text{Ca,hs}} \left[\frac{\text{Na}^+}{\text{Ca}^{2+}} \right]_{\text{hs}} \end{aligned} \quad (4)$$

$$\begin{aligned} \left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{m}} &= \alpha_{\text{Ca,sil}} \left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{sil}} + \alpha_{\text{Ca,carb}} \left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{carb}} + \alpha_{\text{Ca,cy}} \left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{cy}} \\ &+ \alpha_{\text{Ca,hs}} \left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{hs}} \end{aligned} \quad (5)$$

$$\begin{aligned} \left[\frac{\text{Cl}^-}{\text{Ca}^{2+}} \right]_{\text{m}} &= \alpha_{\text{Ca,sil}} \left[\frac{\text{Cl}^-}{\text{Ca}^{2+}} \right]_{\text{sil}} + \alpha_{\text{Ca,carb}} \left[\frac{\text{Cl}^-}{\text{Ca}^{2+}} \right]_{\text{carb}} + \alpha_{\text{Ca,cy}} \left[\frac{\text{Cl}^-}{\text{Ca}^{2+}} \right]_{\text{cy}} \\ &+ \alpha_{\text{Ca,hs}} \left[\frac{\text{Cl}^-}{\text{Ca}^{2+}} \right]_{\text{hs}} \end{aligned} \quad (6)$$

under the conditions that:

$$\alpha_{\text{Ca,sil}} + \alpha_{\text{Ca,carb}} + \alpha_{\text{Ca,cy}} + \alpha_{\text{Ca,hs}} = 1 \quad (7)$$

and

$$0 \leq \alpha_{\text{Ca,sil}} \leq 1; 0 \leq \alpha_{\text{Ca,carb}} \leq 1; 0 \leq \alpha_{\text{Ca,cy}} \leq 1; 0 \leq \alpha_{\text{Ca,hs}} \leq 1 \quad (8)$$

where $\alpha_{\text{Ca,sil}}$, $\alpha_{\text{Ca,carb}}$, $\alpha_{\text{Ca,cy}}$ and $\alpha_{\text{Ca,hs}}$ are the fractions of calcium sourced from silicate weathering, carbonate weathering, cyclic input and hotspots input, respectively. Following previous authors³⁷, we estimated sets of ‘best fit’ endmembers for each sample with 10,000 Monte Carlo runs. In each run, we randomly picked each endmember from normal distributions defined by the mean and the standard deviation of the local endmember estimates (Extended Data Table 3). For each sample, we then obtained an individual endmember estimate from the mean and standard deviation of all Monte Carlo runs that were weighted by the inverse chi-squared misfit between model and data of each run (Extended Data Fig. 6). One sample with an anomalously high ratio of $\left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}} \right]_{\text{m}}$ was discarded from the inversion because it does not fall within the bounds of the silicate–carbonate–hotspots–cyclic endmember (Supplementary Fig. 2a).

Estimate of sulfuric acid contribution to weathering. The contribution of weathering by sulfuric acid was estimated from the total sulfate concentrations corrected for cyclic and hotspots inputs:

$$\left[\text{SO}_4^{2-} \right]_{\text{sulf}} = \left[\text{SO}_4^{2-} \right]_{\text{m}} - \left[\text{SO}_4^{2-} \right]_{\text{cy}} - \left[\text{SO}_4^{2-} \right]_{\text{hs}} \quad (9)$$

with

$$\left[\text{SO}_4^{2-} \right]_{\text{cy}} = \alpha_{\text{Ca,cy}} \left[\frac{\text{SO}_4^{2-}}{\text{Ca}^{2+}} \right]_{\text{cy}}; \left[\text{SO}_4^{2-} \right]_{\text{hs}} = \alpha_{\text{Ca,hs}} \left[\frac{\text{SO}_4^{2-}}{\text{Ca}^{2+}} \right]_{\text{hs}} \quad (10)$$

This calculation assumes negligible anthropogenic and evaporite inputs of sulfate to Taiwanese rivers and an absence of major sulfate sinks (Supplementary Text 4).

Corrections for secondary precipitation of calcite. Supersaturation of almost all sampled waters (Extended Data Fig. 3) and the presence of calcium carbonate precipitates near springs in the field (see Fig. 2 in ref. ³³) attest to the importance of secondary calcite precipitation in the studied catchments. Supersaturation can occur where subsurface waters equilibrate at a high partial pressure of CO₂ (that is, at a low pH) in soils and then degas CO₂ in contact with the atmosphere. Degassing leads to secondary precipitation of CaCO₃, but where secondary precipitation is incomplete, waters remain supersaturated. The enrichment of waters in Sr²⁺ relative to Ca²⁺ and Na⁺ can be used as an indicator for secondary precipitation of calcite^{33,56}. Assuming that all measured Sr is contributed from chemical weathering, solutes from a mixed silicate and carbonate source that have not experienced re-precipitation of calcite should fall on a mixing line between a pure calcite endmember at $\left[\frac{\text{Na}^+}{\text{Ca}^{2+}}\right]$ close to zero and $1,000 \times \left[\frac{\text{Sr}^{2+}}{\text{Ca}^{2+}}\right] = 1 - 2$, and a silicate endmember with both a higher $\left[\frac{\text{Na}^+}{\text{Ca}^{2+}}\right]$ and a higher $\left[\frac{\text{Sr}^{2+}}{\text{Ca}^{2+}}\right]$ ratio³³. We assumed that the published elemental analyses of suspended sediments sampled in the Chenyoulan River³³ approximate the endmember mixing line between silicates and carbonates, and we found the best fit line through these data that is described by:

$$\frac{1,000 \times [\text{Sr}^{2+}]}{[\text{Ca}^{2+}]} = a + b \frac{[\text{Na}^+]}{[\text{Ca}^{2+}]} \quad (11)$$

where a and b are the parameter of the linear fit (Extended Data Fig. 7). With precipitation of calcium, the Sr/Ca ratio increases according to:

$$\frac{[\text{Sr}^{2+}]_f}{[\text{Ca}^{2+}]_f} = \frac{[\text{Sr}^{2+}]_0}{[\text{Ca}^{2+}]_0} \beta^{(kd-1)} \quad (12)$$

where subscripts 0 and f denote the initial and final concentrations respectively, kd is the partition coefficient for Sr into abiotic calcite, and $\beta = \left[\frac{\text{Ca}^{2+}}{\text{Ca}^{2+}}\right]_f$ is the fraction of original calcite remaining in the water⁵⁶. The partition coefficient has been shown to vary with precipitation rates and temperature between ~0.02 and 0.2 (refs. ^{57–59}). The influence of this value is relatively small (Supplementary Text 4), so we follow refs. ^{33,56} and use a value of $kd = 0.05$. Moreover, we neglect incorporation of Mg into calcite, because Mg incorporation into calcite at concentrations of $\left[\frac{\text{Mg}^{2+}}{\text{Ca}^{2+}}\right] < 1$ (Extended Data Fig. 6) is expected to be <5%^{60,61}. To estimate β (and thereby $[\text{Ca}^{2+}]_0$) we note that:

$$\frac{[\text{Sr}^{2+}]_0}{[\text{Ca}^{2+}]_0} = \frac{[\text{Sr}^{2+}]_f}{[\text{Ca}^{2+}]_f} \beta^{(1-kd)} \quad (13)$$

and because the concentration of sodium is unchanged:

$$[\text{Na}^+]_0 = [\text{Na}^+]_f \quad (14)$$

From Eqs. (11) and (14) and the definition of β , we get:

$$\frac{1,000 \times [\text{Sr}^{2+}]_0}{[\text{Ca}^{2+}]_0} = a + b \frac{[\text{Na}^+]_0}{[\text{Ca}^{2+}]_0} = a + b \frac{[\text{Na}^+]_f}{[\text{Ca}^{2+}]_f} \beta \quad (15)$$

Now, we have two estimates for $\frac{[\text{Sr}^{2+}]_0}{[\text{Ca}^{2+}]_0}$ from equations (13) and (15), and we minimized the following equation numerically:

$$\frac{1,000 \times [\text{Sr}^{2+}]_f}{[\text{Ca}^{2+}]_f} \beta^{1-kd} - a - b \frac{[\text{Na}^+]_0}{[\text{Ca}^{2+}]_f} \beta = 0 \quad (16)$$

Uncertainties in the fraction of original calcite remaining in the water, β , were estimated from uncertainties in the regression parameters and the measured concentrations using a Monte Carlo routine. To this end, we repeated the minimization 10,000 times, randomly picking (1) the regression parameters from a bivariate normal distribution defined by the coefficients and variance-covariance matrix of the regression, and (2) $[\text{Sr}^{2+}]$, $[\text{Ca}^{2+}]$ and $[\text{Na}^+]$ from the normal distribution defined by the measurement and uncertainty. We then estimated the uncertainty of β from the standard deviation of β from all Monte Carlo runs. The final uncertainty in the corrected calcium concentrations was estimated from this standard deviation and the analytical uncertainty of the original calcium measurement.

CO₂ budget from weathering reactions. We estimated the impact of erosion rates and the associated weathering patterns measured in Taiwan on the emission or drawdown of CO₂ by considering the balance of alkalinity and dissolved inorganic

carbon produced by weathering (see Table A.1 and equations A11 and A12 in ref. ¹³). Importantly, this CO₂ balance does not depend on whether sulfuric acid reacts with carbonates or silicates. The short-term effect of weathering on the inorganic CO₂ balance, $[\text{CO}_2]_{\text{st}}$, is (Extended Data Fig. 5a):

$$[\text{CO}_2]_{\text{st}} = f_{\text{sulf}} 0.5 [\text{Cat}]_{\text{carb}}^{\text{eq}} + (1 - f_{\text{sulf}}) 0.5 [\text{Cat}]_{\text{carb}}^{\text{eq}} - (1 - f_{\text{sulf}}) ([\text{Cat}]_{\text{carb}}^{\text{eq}} + [\text{Cat}]_{\text{sil}}^{\text{eq}}) \quad (17)$$

where $[\text{CO}_2]$ is the moles of CO₂ produced (positive $[\text{CO}_2]$) or sequestered (negative $[\text{CO}_2]$) per unit volume of weathering fluid. Remembering that:

$$f_{\text{sulf}} = \frac{[\text{SO}_4^{2-}]^{\text{eq}}}{[\text{Cat}]_{\text{carb}}^{\text{eq}} + [\text{Cat}]_{\text{sil}}^{\text{eq}}} \quad (18)$$

this equation can be simplified to:

$$[\text{CO}_2]_{\text{st}} = [\text{SO}_4^{2-}]^{\text{eq}} - 0.5 [\text{Cat}]_{\text{carb}}^{\text{eq}} - [\text{Cat}]_{\text{sil}}^{\text{eq}} \quad (19)$$

Beyond the calcium carbonate compensation time (~10 kyr; ref. ⁶²), the production of alkalinity and dissolved cations by chemical weathering is balanced by the precipitation of (mostly) marine calcium carbonate. Then, the medium-term effect of weathering on the inorganic CO₂ balance, $[\text{CO}_2]_{\text{mt}}$, is (Extended Data Fig. 5b):

$$[\text{CO}_2]_{\text{mt}} = f_{\text{sulf}} 0.5 [\text{Cat}]_{\text{carb}}^{\text{eq}} + (1 - f_{\text{sulf}}) 0.5 [\text{Cat}]_{\text{carb}}^{\text{eq}} - 0.5 (1 - f_{\text{sulf}}) ([\text{Cat}]_{\text{carb}}^{\text{eq}} + [\text{Cat}]_{\text{sil}}^{\text{eq}}) \quad (20)$$

This equation can be simplified to:

$$[\text{CO}_2]_{\text{mt}} = [\text{SO}_4^{2-}]^{\text{eq}} - 0.5 [\text{Cat}]_{\text{sil}}^{\text{eq}} \quad (21)$$

We stress that equations (19) and (21) yield idealized estimates for the CO₂ balance of weathering rates on timescales shorter and longer than the precipitation of calcium carbonate. These scenarios do not consider alkalinity consumption by reverse-weathering reactions⁶³ or alkalinity production by sulfide reduction⁶⁴, and they do not distinguish between alkalinity produced by weathering sodium, potassium or calcium silicate⁶⁵. We also note that the timescale of ~10 kyr for carbonate precipitation in the ocean is only an estimate (although probably a maximum estimate). As argued above, some carbonate evidently precipitates once the water exits the subsurface. The location of precipitation is not directly relevant for the CO₂ balance, but it will influence the timescales over which equations (19) and (21) apply. On multi-million-year timescales, for a system under constant boundary conditions and a carbon cycle in steady state, the long-term effect of weathering on the inorganic CO₂ balance, $[\text{CO}_2]_{\text{lt}}$, may depend only on the calcium produced by silicate weathering⁶ (Extended Data Fig. 5c):

$$[\text{CO}_2]_{\text{lt}} = -0.5 [\text{Ca}^{2+}]_{\text{sil}}^{\text{eq}} \quad (22)$$

where $[\text{Ca}^{2+}]_{\text{sil}}^{\text{eq}}$ is the equivalent calcium concentration from silicate weathering.

Carbonate equilibria. The saturation state of calcium carbonate in water (Ω_{cal}) is defined as:

$$\Omega_{\text{cal}} = \frac{a_{\text{Ca}^{2+}} a_{\text{CO}_3^{2-}}}{K_{\text{sp}}} \quad (23)$$

where K_{sp} is the equilibrium constant for the dissolution/precipitation reaction of calcium carbonate in water (Extended Data Table 2). By definition, the second dissociation constant of carbonic acid (K_2):

$$K_2 = \frac{a_{\text{H}^+} a_{\text{CO}_3^{2-}}}{a_{\text{HCO}_3^-}} \quad (24)$$

It follows that

$$\Omega_{\text{cal}} = \frac{K_2 a_{\text{Ca}^{2+}} a_{\text{HCO}_3^-}}{K_{\text{sp}} a_{\text{H}^+}} \quad (25)$$

In order to convert the measured concentrations into activities, we calculated the activity coefficients γ_i for both calcium and bicarbonate using the Davies equation⁶⁶:

$$\log_{10} \gamma_i = -A v_i^2 \left(\frac{\sqrt{I}}{1 + \sqrt{I}} - 0.3I \right) \quad (26)$$

where $A = 1.82 \times 10^6 (\epsilon T)^{-\frac{3}{2}}$ is an empirical constant, T is the temperature, ϵ is the dielectric constant and I is the ionic strength of the solution given by:

$$I = \frac{1}{2} \sum_i v_i^2 [i] \quad (27)$$

We estimated the dielectric constant using an empirical fit to measurements of the dielectric constant at atmospheric pressures and temperatures between 273.15 and 373.15 K (Table 4 in ref. 67):

$$\epsilon = 308.6464e^{(-0.00459717 T)} \quad (28)$$

The overall temperature dependence of the factor A in equation (26) is small and $A \approx 0.51$.

We also estimated the equilibrium constant for reaction j at water temperature T ($K_{j,T}$) using the van't Hoff equation⁶⁸:

$$\ln \left(\frac{K_{j,T}}{K_{j,298}} \right) = -\frac{\Delta H_j^\circ}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \quad (29)$$

where $K_{j,298}$ is the equilibrium constant of reaction j at 298.15 K, ΔH_j° is the standard enthalpy of reaction j and R is the ideal gas constant. The saturation index for calcium carbonate in water SI_{cal} at temperature T (Extended Data Fig. 3) is then:

$$\begin{aligned} SI_{\text{cal}} &= \log_{10} (\Omega_{\text{cal}}) \\ &= \log_{10} (K_{2,T}) - \log_{10} (K_{\text{sp},T}) \\ &\quad + \log_{10} (\gamma_{\text{Ca}^{2+}} [\text{Ca}^{2+}]) + \log_{10} (\gamma_{\text{HCO}_3^-} [\text{HCO}_3^-]) + \text{pH} \end{aligned} \quad (30)$$

In addition to the saturation index, we estimated the pH at equilibrium (pH_{eq}) using:

$$\begin{aligned} \text{pH}_{\text{eq}} &= \log_{10} (K_{\text{sp},T}) - \log_{10} (K_{2,T}) \\ &\quad - \log_{10} (\gamma_{\text{Ca}^{2+}} [\text{Ca}^{2+}]) - \log_{10} (\gamma_{\text{HCO}_3^-} [\text{HCO}_3^-]) \end{aligned} \quad (31)$$

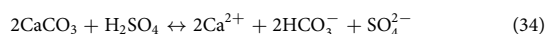
This pH_{eq} (Fig. 3b) represents the pH of a solution that is exactly saturated with respect to the measured solutes. The value is a maximum estimate for the pH in the subsurface waters and the pH could, in principle, be lower. The presence of detrital carbonate in streams⁶⁹ and carbonate precipitation around natural springs in the field suggests that carbonates are not strongly undersaturated in the subsurface. For all steps described above, uncertainties were propagated from the analytical uncertainties of measured concentrations. In these calculations, we considered the contributions of solutes only from weathering, cyclic sources and the secondary carbonate lost to precipitation. This procedure assumes that any hot spring input occurs in the stream, downstream of the weathering zone, and does not contribute substantially to the dissolved Sr^{2+} (Supplementary Text 4).

Estimate of pH change due to addition of sulfuric acid. We compared the inferred subsurface pH, pH_{eq} , with the pH obtained by adding sulfuric acid to a carbonate-buffered solution, pH_{SO_4} . For simplicity, this particular calculation assumes that all sulfuric acid reacts with calcium carbonate, which is reasonable, particularly for the northern rivers in which carbonate weathering constitutes much more than half and up to >90% of the weathering budget. We define pH_{SO_4} as the pH of a solution that results from the addition of sulfuric acid with concentration $[\text{SO}_4^{2-}]$ to a calcium-carbonate-buffered solution with an initial pH noted as pH_0 . From equation (25), we recall that for a calcium-carbonate-buffered solution at equilibrium (that is, at $\Omega_{\text{cal}} = 1$), we have:

$$a_{\text{H}^+,0} = \frac{K_2}{K_{\text{sp}}} a_{\text{Ca}^{2+},0} a_{\text{HCO}_3^-,0} \quad (32)$$

$$a_{\text{H}^+,0} = \frac{K_2}{K_{\text{sp}}} \gamma_{\text{Ca}^{2+},0} [\text{Ca}^{2+}]_0 \gamma_{\text{HCO}_3^-,0} [\text{HCO}_3^-]_0 \quad (33)$$

where the subscript 0 denotes the quantities before the addition of sulfuric acid. We further assume that the following reaction of sulfuric acid with calcium carbonate dominates:



and that HCO_3^- remains the dominant dissolved carbonate species (which is reasonable for the range of pH between 6 and 8). After addition of the acid, the new activity of protons at equilibrium with the acid is:

$$\begin{aligned} a_{\text{H}^+,f} &= \frac{K_2}{K_{\text{sp}}} \gamma_{\text{Ca}^{2+},f} [\text{Ca}^{2+}]_f \gamma_{\text{HCO}_3^-,f} [\text{HCO}_3^-]_f \\ &= \frac{K_2}{K_{\text{sp}}} \gamma_{\text{Ca}^{2+},f} \left([\text{Ca}^{2+}]_0 + 2 [\text{SO}_4^{2-}] \right) \gamma_{\text{HCO}_3^-,f} \left([\text{HCO}_3^-]_0 + 2 [\text{SO}_4^{2-}] \right) \end{aligned} \quad (35)$$

where the subscript f denotes the final quantities after equilibration with the acid. Hence:

$$\begin{aligned} a_{\text{H}^+,f} &= \frac{K_2}{K_{\text{sp}}} \gamma_{\text{Ca}^{2+},f} \gamma_{\text{HCO}_3^-,f} \left([\text{Ca}^{2+}]_0 [\text{HCO}_3^-]_0 + 2 [\text{SO}_4^{2-}] \right. \\ &\quad \left. [\text{Ca}^{2+}]_0 + 2 [\text{SO}_4^{2-}] [\text{HCO}_3^-]_0 + 4 [\text{SO}_4^{2-}]^2 \right) \end{aligned} \quad (36)$$

Next, we link the initial concentration of calcium, $[\text{Ca}^{2+}]_0$ and bicarbonate, $[\text{HCO}_3^-]_0$, to the initial pH, pH_0 (or initial activity of protons $a_{\text{H}^+,0}$). Let $[\text{CC}]_0 = [\text{Ca}^{2+}]_0 [\text{HCO}_3^-]_0$. Rearranging equation (33) we get:

$$[\text{CC}]_0 = [\text{Ca}^{2+}]_0 [\text{HCO}_3^-]_0 = \frac{K_{\text{sp}}}{K_2} \frac{a_{\text{H}^+,0}}{\gamma_{\text{Ca}^{2+},0} \gamma_{\text{HCO}_3^-,0}} \quad (37)$$

Further, we posit that we can express the concentration of calcium before the addition of sulfuric acid as a function of the concentration of bicarbonate and vice versa:

$$[\text{Ca}^{2+}]_0 = \frac{X_{\text{Ca}^{2+},0}}{2} [\text{HCO}_3^-]_0 \quad (38)$$

and

$$[\text{HCO}_3^-]_0 = \frac{2}{X_{\text{Ca}^{2+},0}} [\text{Ca}^{2+}]_0 \quad (39)$$

where $X_{\text{Ca}^{2+},0}$ is the fraction of the total cation charge that is represented by calcium:

$$X_{\text{Ca}^{2+},0} = \frac{2 [\text{Ca}^{2+}]_0}{\sum v_i [i]_0} \quad (40)$$

with

$$[\text{Ca}^{2+}]_0 = [\text{Ca}^{2+}]_f - 2 [\text{SO}_4^{2-}] \quad (41)$$

and

$$\sum v_i [i]_0 = \sum v_i [i]_f - 4 [\text{SO}_4^{2-}] \quad (42)$$

Equations (38)–(42) allow us to express the concentrations of calcium and bicarbonate by their total concentration $[\text{CC}]_0$:

$$[\text{Ca}^{2+}]_0 = \sqrt{\frac{X_{\text{Ca}^{2+}}}{2} [\text{CC}]_0} \quad (43)$$

and

$$[\text{HCO}_3^-]_0 = \sqrt{\frac{2}{X_{\text{Ca}^{2+}}} [\text{CC}]_0} \quad (44)$$

Now, the combination of equations (36), (43) and (44) yields:

$$\begin{aligned} a_{\text{H}^+,f} &= \frac{K_2}{K_{\text{sp}}} \gamma_{\text{Ca}^{2+},f} \gamma_{\text{HCO}_3^-,f} \\ &\quad \left([\text{CC}]_0 + 2 [\text{SO}_4^{2-}] \sqrt{[\text{CC}]_0} \left(\sqrt{\frac{X_{\text{Ca}^{2+},0}}{2} + \frac{2}{X_{\text{Ca}^{2+},0}}} \right) + 4 [\text{SO}_4^{2-}]^2 \right) \end{aligned} \quad (45)$$

and the final pH after addition of sulfuric acid is:

$$\text{pH}_{\text{SO}_4} = -\log (a_{\text{H}^+,f}) \quad (46)$$

In our data analysis, we used equation (37) and an estimate of $\text{pH}_0 = 8.51$ (see below for reasoning) to calculate $[\text{CC}]_0$. Then, we calculated the pH change that is expected from pH_0 due to the addition of sulfuric acid. We discarded 37 datapoints for which sulfate concentrations are larger than or within $10 \mu\text{mol l}^{-1}$ of the calcium concentrations ($[\text{Ca}^{2+}]_f - 2 [\text{SO}_4^{2-}] < 10 \mu\text{mol l}^{-1}$). In these cases, the assumption that all sulfuric acid was buffered entirely by calcium carbonate dissolution is

clearly inapplicable. Note that the factor of $\sqrt{\frac{X_{\text{Ca}^{2+},0}}{2} + \frac{2}{X_{\text{Ca}^{2+},0}}}$ and, thus, the

measured calcium $[\text{Ca}^{2+}]_f$ and inferred bicarbonate $[\text{HCO}_3^-]_f$ values have a very small effect on the final pH_{SO_4} . The main controls on pH_{SO_4} are the initial pH and the concentration of sulfate. Thus, pH_{SO_4} (equation (46)) is mostly independent of pH_{eq} (equation (31)), except for the shared activity coefficients and equilibrium constants. We define the initial pH, $\text{pH}_0 = 8.51$ ($\text{pH}_0 = 8.58$ without secondary precipitation of calcite), as the unique value at which a regression through the data intersects the point $\text{pH}_{\text{SO}_4} = \text{pH}_{\text{eq}} = \text{pH}_0$. This point represents the necessity

that the subsurface pH before addition of sulfuric acid is equal to the initial pH. All uncertainties were propagated from the analytical uncertainties of measured concentrations. We note that bicarbonate contributions (used to calculate pH_{eq}) were derived from a charge balance and are, therefore, negatively correlated with sulfate concentrations. In the case that we inadvertently ignore a major charge contributor (Supplementary Text 4), the correlation between the two pH estimates could be weakened.

Data availability

All data analysed in this study are available at <https://doi.org/10.5880/GFZ.4.6.2021.001> (ref. ⁷⁰).

Code availability

The MATLAB code used to implement the inversion of the geochemical data is available on request from the corresponding author.

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Author contributions

N.H., R.E. and A.B. conceived the study; A.B., R.E., H.J.H.-G. and N.H. contributed samples and laboratory analyses; A.B. led the data analysis with contributions from J.K.C.R., N.H., R.E. and A.G., and the writing of the manuscript with contributions from all authors.

Competing interests

The authors declare no competing interests.

Additional information

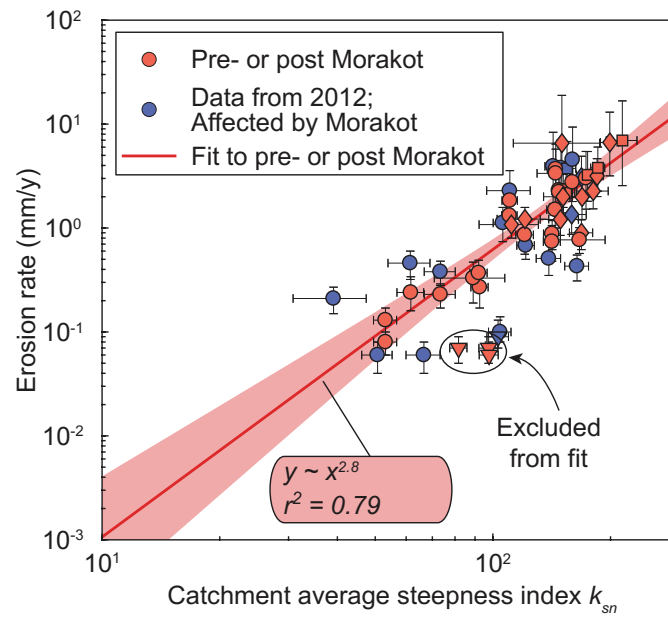
Extended data is available for this paper at <https://doi.org/10.1038/s41561-021-00714-3>.

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41561-021-00714-3>.

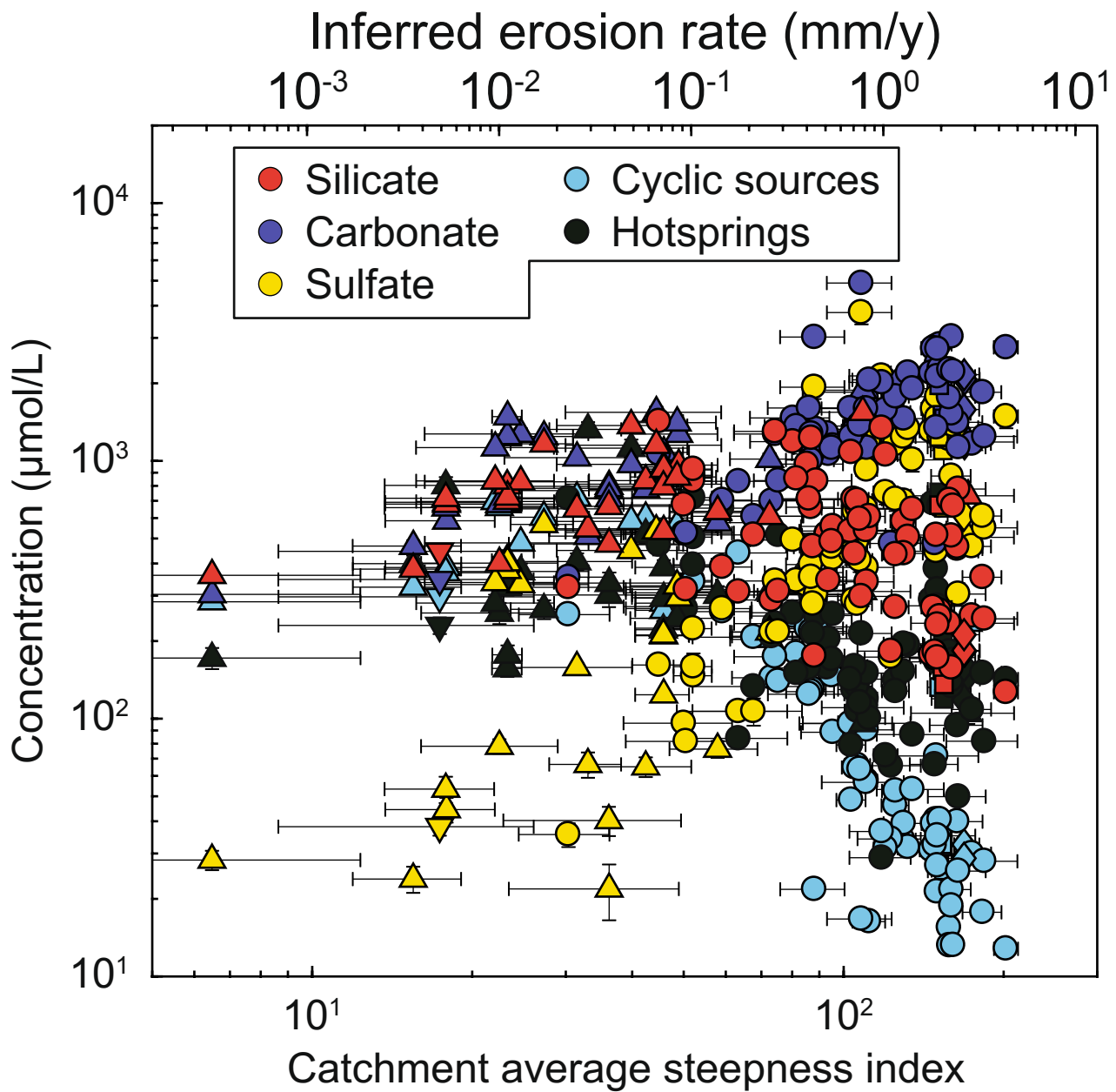
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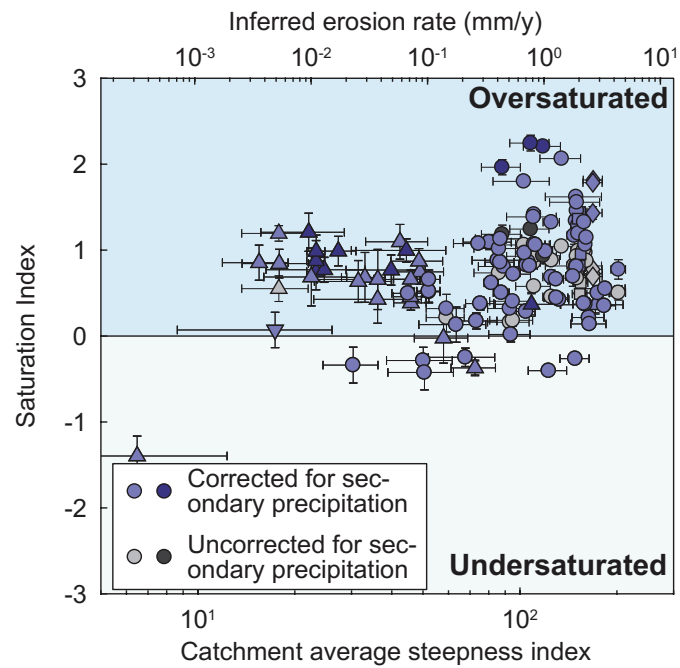
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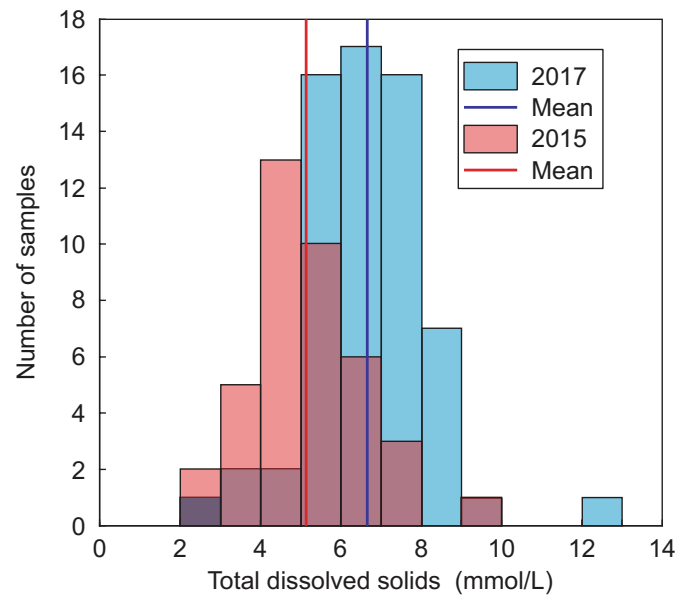
Extended Data Fig. 1 | Normalized steepness index and erosion rates. Scaling between catchment-averaged steepness index and catchment-averaged erosion rates from published ^{10}Be cosmogenic nuclide data. Red line shows regression through data from before and from at least 6 years after Typhoon Morakot (red points) and 95% confidence band. Three outliers, all located in the Mutan formation (see Fig. 1) were excluded. If they were included, the inferred erosion rate gradient across the study area would be even wider. Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval.



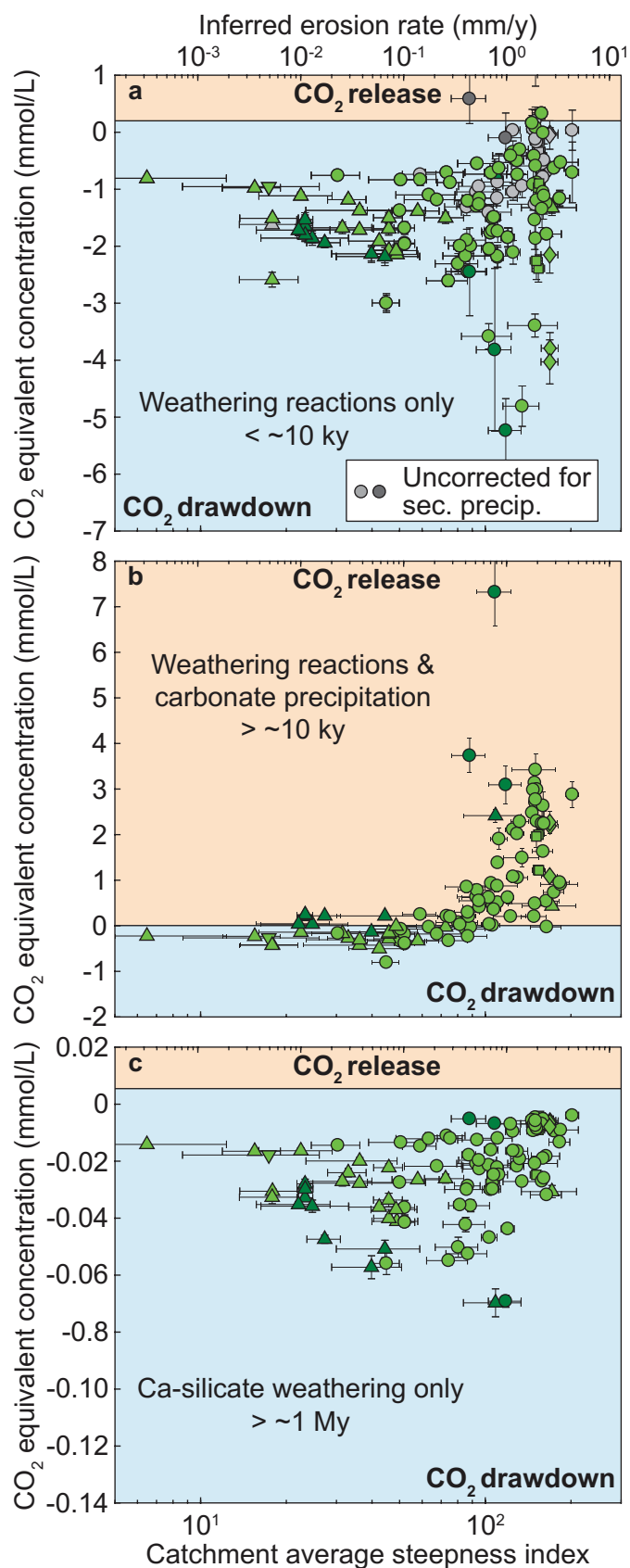
Extended Data Fig. 2 | Inversion results including hotspring and cyclic contributions. Symbology is equivalent to Fig. 2. Error bars are equivalent to a 95% confidence interval.



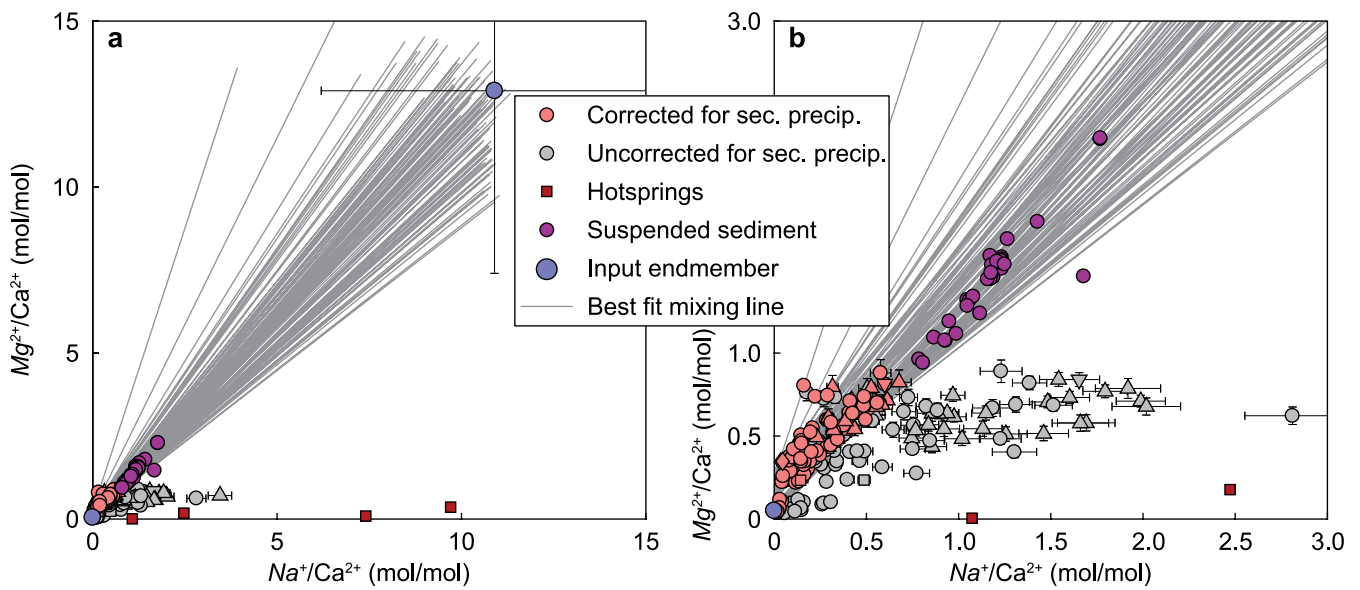
Extended Data Fig. 3 | Saturation index. Saturation index of samples corrected for secondary precipitation (blue) and uncorrected for secondary precipitation (grey). Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval. Samples with anomalously high sulfate concentrations are marked as darker points.



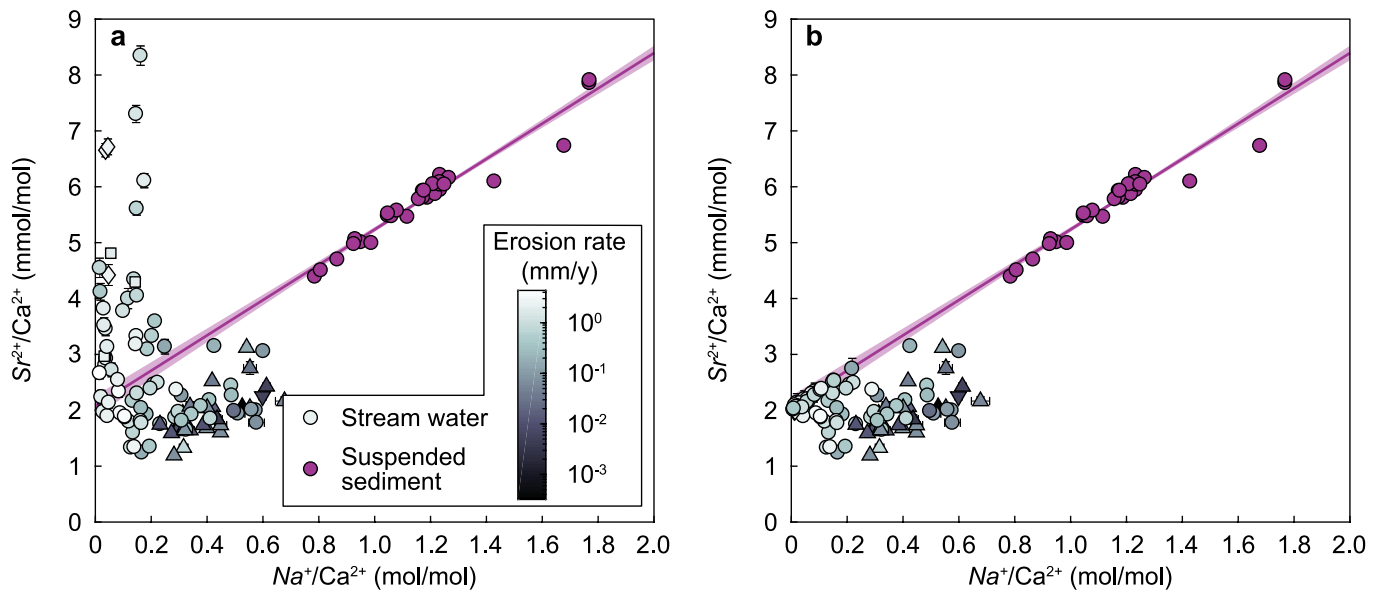
Extended Data Fig. 4 | Variations of total dissolved solid concentrations between sampling seasons. Total dissolved solid values for samples collected in 2017 and 2015. Both sampling seasons include samples from across the erosion rate gradient. Whereas the drier year of 2015 was characterized by somewhat higher concentrations, both distributions are within one standard deviation.



Extended Data Fig. 5 | Impact of weathering on the CO₂-balance. Moles of CO₂ emitted or sequestered per volume of water plotted against catchment-averaged steepness index (lower x-axis) and inferred erosion rates (upper x-axis) for different scenarios and timescales. **a**, Impact of weathering reactions only. **b**, Impact of weathering reactions and carbonate precipitation. **c**, Impact of only Ca-silicate weathering. Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval.



Extended Data Fig. 6 | Mixing diagram for weathering contributions. **a**, Mg^{2+}/Ca^{2+} - Na^+/Ca^{2+} mixing diagram showing raw data (grey) and weathering contributions (red). Blue points mark endmembers with two standard deviations. Dark-red squares are available hot spring data. Purple points are the suspended sediment data from the Chenyulan. The grey lines are the best fit endmember mixing lines found by the inverse approach. **b**, Close-up of panel a. Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval.



Extended Data Fig. 7 | Correction for secondary precipitation. a, Stream water chemistry including contributions from rainwater and weathering coloured by erosion rate. All purple suspended sediment data are used to define the endmember trend (purple line with 95% confidence bands), and all data above that line are projected onto that line. **b**, Stream water chemistry corrected for secondary precipitation of calcite. Only points above the mixing line were corrected. Symbols mark dominant lithology (see Fig. 1). Error bars are equivalent to a 95% confidence interval.

Extended Data Table 1 | Notations

Notation	Definition
A	Empirical constant in Davies equation
a, b	Regression coefficients for endmember mixing in $\frac{1000 \times [Sr^{2+}]}{[Ca^{2+}]} - \frac{[Na^+]}{[Ca^{2+}]}$ space
a_i	Activity of ion i
$\alpha_{Ca,carb/sil}$	Fraction of Ca sourced from carbonates/silicates
β	Ratio of $\frac{[Ca^{2+}]_f}{[Ca^{2+}]_0}$ for secondary precipitation analysis
$[Cat]_{wtot}$	Total concentration of cations from weathering in mol/L
$[Cat]_{carb}$	Concentration of cations from carbonates in mol/L
$[Cat]_{sil}$	Concentration of cations from silicates in mol/L
$[Cat]_{wtot}^{eq}$	Total cation charge from weathering in eq/L
$[Cat]_{carb}^{eq}$	Charge equivalents of cations from carbonates in eq/L
$[Cat]_{sil}^{eq}$	Charge equivalents of cations from silicates in eq/L
γ_i	Activity coefficient of ion i
ΔH_j°	Standard enthalpy of reaction j (see Extended Data Table 2)
E_a	Activation energy
ϵ	dielectric constant
f_{carb}	Fraction of cation charge from carbonate weathering
f_{sulf}	Fraction of cation charge balanced by sulphate
$[i]$	Concentration of ion i in mol/L
$[i]^{eq}$	Concentration of ion i in charge equivalents
K_j	Equilibrium constant of reaction j (see Extended Data Table 2)
k_{sn}	Catchment-averaged normalized steepness index
kd	Sr/Ca partition coefficient for calcite
pH_0	Initial pH before addition of sulphuric acid
pH_{eq}	pH of a solution in equilibrium with calcium carbonate
pH_{SO_4}	pH of a carbonate buffered solution with pH_0 after addition of $[SO_4^{2-}]$
R	Ideal gas constant
SI_{cal}	Saturation index of calcium carbonate in water
T	Temperature
u_x	Atomic mass of x
v_i	Valence of ion i
$X_{Ca,0}$	fraction of the total cation charge that is represented by calcium,
$\langle x \rangle$	Concentration of x in weight percent
Ω_{cal}	Saturation state of calcium carbonate in water

Extended Data Table 2 | Reactions, equilibrium constants, and standard enthalpies

Description	Reaction	Equilibrium constant at 298.15 °K	Standard enthalpy (kJ mol ⁻¹)
Dissolution of CO ₂ in water (Henry's constant)	$CO_{2(g)} \leftrightarrow CO_{2(aq)}$	$K_H = 10^{-1.468}$	$\Delta H_H^\circ = -19.98$
First dissociation of CO ₂ in water	$CO_{2(aq)} + 2H_2O \leftrightarrow HCO_3^- + H_3O^+$	$K_1 = 10^{-6.352}$	$\Delta H_1^\circ = 9.109$
Second dissociation of CO ₂ in water	$HCO_3^- + H_2O \leftrightarrow CO_3^{2-} + H_3O^+$	$K_2 = 10^{-10.329}$	$\Delta H_2^\circ = 14.90$
Dissolution/Precipitation of carbonate in water	$CaCO_3 \leftrightarrow Ca^{2+} + CO_3^{2-}$	$K_{sp} = 10^{-8.480}$	$\Delta H_{sp}^\circ = -9.612$

All parameters are from ref. ⁷¹

Extended Data Table 3 | Endmember compositions

Silicate	Carbonate
Global endmember compositions ^{8,55}	
$\left[\frac{Ca^{2+}}{Na^+}\right]_{sil} = 0.35 \pm 0.25$	$\left[\frac{Ca^{2+}}{Na^+}\right]_{carb} = 60 \pm 30$
$\left[\frac{Mg^{2+}}{Na^+}\right]_{sil} = 0.25 \pm 0.20$	$\left[\frac{Mg^{2+}}{Na^+}\right]_{carb} = 30 \pm 15$
Local endmember compositions: Forward model	
$\left[\frac{Ca^{2+}}{Na^+}\right]_{sil} = 0.10 \pm 0.06$	$\left[\frac{Ca^{2+}}{Na^+}\right]_{carb} = \infty$
$\left[\frac{Mg^{2+}}{Na^+}\right]_{sil} = 0.80 \pm 0.59$	$\left[\frac{Mg^{2+}}{Na^+}\right]_{carb} = \infty$
Local endmember compositions: Inverse model	
$\left[\frac{Na^+}{Ca^{2+}}\right]_{sil} = 10.9 \pm 4.7$	$\left[\frac{Na^+}{Ca^{2+}}\right]_{carb} = 0 \pm 0$
$\left[\frac{Mg^{2+}}{Ca^{2+}}\right]_{sil} = 12.9 \pm 5.5$	$\left[\frac{Mg^{2+}}{Ca^{2+}}\right]_{carb} = 0.05 + 0.24 - 0.05$
Seawater	Hotsprings
$\left[\frac{Na^+}{Ca^{2+}}\right]_{sw} = 45.5$	$\left[\frac{Na^+}{Ca^{2+}}\right]_{hs} = 315 \pm 150$
$\left[\frac{Mg^{2+}}{Ca^{2+}}\right]_{sw} = 5.1$	$\left[\frac{Mg^{2+}}{Ca^{2+}}\right]_{hs} = 0.35 \pm 0.16$
$\left[\frac{Cl^-}{Ca^{2+}}\right]_{sw} = 53$	$\left[\frac{Cl^-}{Ca^{2+}}\right]_{hs} = 5.4 \pm 2.4$
$\left[\frac{SO_4^{2-}}{Ca^{2+}}\right]_{sw} = 11.0$	$\left[\frac{SO_4^{2-}}{Ca^{2+}}\right]_{hs} = 1.7 \pm 0.8$
$\left[\frac{K^+}{Ca^{2+}}\right]_{sw} = 1.0$	$\left[\frac{K^+}{Ca^{2+}}\right]_{hs} = 1.4 \pm 0.6$