Mountain glaciation drives rapid oxidation of rock-bound organic carbon

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Over millions of years, the oxidation of organic carbon contained within sedimentary rocks is one of the main sources of carbon dioxide to the atmosphere, yet the controls on this emission remain poorly constrained. We use rhenium to track the oxidation of rock-bound organic carbon in the mountain watersheds of New Zealand, where high rates of physical erosion expose rocks to chemical weathering. Oxidative weathering fluxes are two to three times higher in watersheds dominated by valley glaciers and exposed to frost shattering processes, compared to those with less glacial cover; a feature that we also observe in mountain watersheds globally. Consequently, we show that mountain glaciation can result in an atmospheric carbon dioxide source during weathering and erosion, as fresh minerals are exposed for weathering in an environment with high oxygen availability. This provides a counter mechanism against global cooling over geological time scales.

INTRODUCTION

The exposure of organic matter in rocks to oxidative weathering at Earth’s surface releases carbon dioxide (CO₂) to the atmosphere from long-term (>10⁶ years) storage in the lithosphere and consumes atmospheric oxygen (O₂) (1–3). The global CO₂ emissions from the oxidation of rock-derived organic carbon [petrogenic OC (OCpetro)] are estimated to be 40 to 100 × 10⁷ metric tons of carbon (tC) per year (1). Over million-year (geological) time scales, this emission represents the main source of atmospheric CO₂ alongside volcanism and metamorphism (1–4) and plays a role in setting atmospheric O₂ concentrations (2, 5). Geological CO₂ emissions are removed by the atmosphere through chemical weathering of silicate minerals by carbonic acid coupled to carbonate precipitation (4, 6) and the burial of recently photosynthesized OC (3, 7). These atmospheric CO₂ drawdown mechanisms are regulated by erosion, temperature, and runoff and are thought to stabilize CO₂ concentrations and global climate (6). However, we have little understanding of the factors controlling OCpetro oxidation rates (1, 8, 9) and hence how millennial-scale changes in climate (10) might modify this major CO₂ emission.

Chemical weathering of OCpetro proceeds as surficial gases and fluids permeate through sedimentary rocks, oxidizing organic matter and releasing CO₂ (1). The kinetics of OCpetro weathering appear to be ~10 times faster than the kinetics of silicate mineral weathering (11). Faster reaction kinetics imply that shorter fluid residence times are required to reach chemical equilibrium and maximize weathering fluxes during OCpetro oxidation (6), compared to acid hydrolysis silicate weathering (6, 12). When considered together with the high concentrations of O₂ in the present-day atmosphere, weathering models suggest that the OCpetro oxidation rate is set by the mineral supply rate in most locations globally (13). Microorganisms may also be important facilitators of OCpetro oxidation (14). On the basis of these observations, we propose that mountain glaciation could significantly enhance OCpetro oxidation rates due to a combination of physical and biogeochemical factors: (i) frost cracking and abrasive glacial grinding processes, which produce fine sediment with more surface area in an environment with high water availability (15, 16); (ii) lower vegetation and soil cover that can increase the availability of O₂ to exposed bedrock and in deeper soil horizons; and (iii) the activity of microorganisms catalyzing weathering, both subglacially and during primary ecological succession on moraines (14, 17–19). Previous work has suggested that sulfide oxidation is enhanced subglacially (18, 20). If OCpetro oxidation rates also increase, CO₂ release may be highest during periods of repeated mountain glaciation over millennia (21) and may provide a mechanism for countering cooling trends in Earth’s climate over time scales of 100 thousand years to millions of years.

Here, we examine the potential for mountain glaciation to increase OCpetro oxidation rates. We focus on the mountain watersheds of the western Southern Alps, New Zealand (Fig. 1A), where lithological contrasts are relatively small along strike of the Alpine Fault (22), but glacial coverage is variable (23). Previous work has suggested that high silicate weathering rates (20, 24) are facilitated by rapid soil production (25) and mineral supply by bedrock landslides (26). Here, we assess the rates of oxidative weathering and the role of mountain glaciers using river water and sediment samples from 13 watersheds (see Materials and Methods). We also collected samples from two watersheds in the eastern Southern Alps that host glaciers but have lower physical erosion rates and from the Waipaoa River in North Island, which has a high erosion rate but no glaciers (24). In addition, we compile measurements from mountain watersheds draining OCpetro-bearing sedimentary rocks in North America (Yukon and Mackenzie) and Asia (Taiwan rivers, Ganges, and Brahmaputra) (9, 27).

To assess and quantify OCpetro oxidation rates, we measured the concentration of the redox-sensitive trace element rhenium (Re) in river waters ([Re]diss), river bed materials ([Re]BM), weathered colluvium and surface soils on hillslopes ([Re]CLOL) using isotope dilution and inductively coupled plasma mass spectrometry (ICP-MS) (see Materials and Methods). The close association of Re and OCpetro in sedimentary rocks (28) and the solubility of Re upon oxidation during weathering (29) (present as the soluble perrenthan oxyanion, ReO₄⁻, in soils and rivers with pH values between 5.5 and 9.5) have led previous studies to suggest that Re can trace OCpetro oxidation (1, 9, 30, 31). For river watersheds with similar runoff and bedrock composition, the dissolved Re concentration, [Re]diss has been shown to reflect the relative rate of oxidative weathering (9). The dissolved Re discharge (mol year⁻¹) or dissolved Re yield (mol km⁻² year⁻¹) provides a more direct means to quantify the oxidative weathering yield (1, 9, 30, 31). The dissolved Re yield has been...
used to estimate the associated CO₂ emissions by OCpetro oxidation when the Re to OCpetro ratio of the rocks undergoing weathering has been characterized (9). The main uncertainties in the use of the Re proxy derive from the following: (i) As a soluble element, Re may be mobilized more effectively during weathering than the CO₂ derived from OCpetro oxidation (30); (ii) Re may be hosted in silicate and sulfide minerals (27); and (iii) graphitic OCpetro is less susceptible to oxidation (32).

**RESULTS**

**River bed materials and weathered colluvium**

The mean organic carbon (OC) weight percentage in river bed materials, [OC]BM, from the western Southern Alps is 0.13 ± 0.01 weight % (wt %) \( (n = 31, ±2 SE) \), and the mean stable carbon isotope composition is \( δ^{13}C = -22.5 ± 0.6\%_o \) \( (n = 31, ±2 SE) \). For comparison, the [OC]BM values are lower than those in Taiwan \( (9) \) but more than double those measured in the Himalaya \( (32) \). The mean values of river bed materials are similar to previous measurements from this location \( (22, 33) \) and to the mean of bedrock values in the western Southern Alps, [OC] = 0.15 ± 0.05 wt % and \( δ^{13}C = -21.1 ± 1.1\%_o \) \( (n = 11, ±2 SE) \) \( (22) \), suggesting that they are dominated by OCpetro with minor inputs from biospheric OC generated by recent photosynthesis by C₃ plants \( (δ^{13}C = -28\%_o) \). The river bed materials have less variability in their average composition when compared to bedrock, which likely reflects the integration of OCpetro-bearing sediment from landslides and mass wasting processes, which can erode OCpetro from large areas of the watershed \( (34, 35) \). The river sediments downstream are a mixture of these inputs \( (22) \). Although the river bed materials have a slightly lower OC concentration than bedrock in the mountain belt, they are indistinguishable within the variability in the means and can therefore provide a robust method to assess the watershed-averaged bedrock composition \( (22, 32, 36) \). This is consistent with findings in other erosive settings where bedrock landslides excavate deep into the landscape, tap into unweathered rock, and supply it to rivers in the sand-silt-clay fraction of river sediments \( (37) \).

The mean Re concentration in river bed materials, [Re]BM, is 118 ± 21 parts per trillion (ppt) \( (n = 31, ±2 SE) \) \( (table S1) \). Although this Re concentration is low for sedimentary rocks \( (38) \), it is in line with the low [OC]BM in the western Southern Alps and consistent with organic matter being a dominant host of Re in sediments \( (28) \). The mean [Re]BM/[OC]BM ratio is 0.9 ± 0.2 \( × 10^{-7} \) g⁻¹ \( (n = 31, ±2 SE) \), and there is no systematic variability along strike of the Alpine Fault (that is, no variability with latitude and longitude).

We examine samples of weathered colluvium collected from hillslopes in the western Southern Alps for loss of OCpetro and Re \( (table S2) \). In the Docherty Creek watershed, which neighbors the Waibao River, weathered colluvium has a mean [Re]COL = 29 ± 6 ppt \( (n = 4, ±2 SE) \), compared to local river bed materials downstream at Docherty Creek \( (NZ14-90; [Re]BM = 112 ppt) \). Soil litter samples are also depleted in Re \( (table S2) \). This is expected if Re loss occurs during oxidative weathering. The mean OC content of the colluvium, [OC]COL = 1.2 ± 0.3 wt % \( (n = 4, ±2 SE) \), is higher than that of the local river bed materials \( (NZ14-90; [OC]BM = 0.21 wt \%) \). However, organic matter in the colluvium has a stable isotopic composition \( (mean δ^{13}C = -25.9 ± 0.4\%) \) and radiocarbon activity \( (reported as fraction modern, F_{mod} = 0.80 ± 0.09) \) that are very different from the expected rock inputs \( (NZ14-90; δ^{13}C = -21.2\%_o \) and expected \( F_{mod} = 0) \). The values suggest an important contribution from biospheric OC in these samples \( (22) \).

To assess the OCpetro content of the colluvium and to account for biospheric OC, we use \( δ^{13}C \) and \( F_{mod} \) values in a mixing analysis \( (see Materials and Methods) \). The high [OC]COL of these samples and the relatively low OCpetro content of rocks in this mountain range \( (32) \), together with the observation that century-aged biospheric OC is important in these New Zealand soils \( (fig. S1) \), result in uncertainty on the absolute OCpetro contents. The mean [OCpetro] of the weathered colluvium is calculated as 0.15 ± 0.06 \% \( (n = 4, ±2 SE) \), which is lower than that of the river bed material. The data are consistent with OCpetro loss during weathering on hillslopes. The coupled loss of Re and OCpetro during chemical weathering supports previous measurements on soil from the Ohio Shale \( (table S2) \), the Himalaya \( (39) \), and Taiwan \( (9) \). The weathered colluvium from New Zealand and published soil data show that Re is generally more mobile during OCpetro, weathering, and so we account for this when estimating the OCpetro weathering rate using the dissolved Re flux \( (see Materials and Methods) \).

**River waters**

The major dissolved ions in rivers draining the western Southern Alps \( (Ca^{2+}, Mg^{2+}, Na^+, \text{and} K^+) \) characterize the overall weathering processes.
and reflect a source rock comprising metasedimentary silicate rocks hosting trace carbonate minerals (fig. S2A). The water measurements from this study in 2014 are consistent with sampling campaigns in 1998, 1999, and 2000 (20, 24). Notably, all sampling campaigns find higher relative HCO$_3^-$ and Ca$^{2+}$ concentrations in the heavily glaciated Fox and Waiho rivers (fig. S2A), which probably reflects the higher susceptibility of carbonate minerals to acid hydrolysis reactions in these watersheds. The overlap of the data sets collected from different years, seasons, and flow regimes (fig. S2, A and B) suggests that spatial patterns in dissolved ion composition are retained despite the potential for seasonal and flood–event scale variability (40).

The [Re]$\text{diss}$ values in western Southern Alps rivers range from 0.81 to 11.55 pmol liter$^{-1}$, with a mean $= 3.05 \pm 0.69$ pmol liter$^{-1}$ ($n = 51$, ±2 SE) (table S3). There is a distinct variability between different watersheds (fig. 1A), with the Waiho and Fox watersheds having the highest mean [Re]$\text{diss}$ values throughout the sampling period. Mean [Re]$\text{diss}$ is not correlated with [Re]$\text{BM}$ in the western Southern Alps, suggesting that the bedrock geology does not set the spatial pattern in [Re]$\text{diss}$. When all the data are considered together, the watershed-averaged [Re]$\text{diss}$ is correlated with the proportion of area covered by glaciers upstream (fig. 1B; $r^2 = 0.87$, $P < 0.001$, $n = 13$).

The [Re]$\text{diss}$ values are generally low compared to those measured in rivers globally (27) and in rivers draining metasedimentary rocks in Taiwan, which have values ranging from ~5 to 25 pmol liter$^{-1}$ (9). However, when [Re]$\text{diss}$ values are normalized to the concentration of Re in river bed materials, [Re]$\text{BM}$ (9, 32, 36), the values are more similar to those in Taiwan. In the western Southern Alps, watersheds dominated by river erosion and bedrock landslides (that is, not by glacial erosion processes) have a mean [Re]$\text{diss}$ = 1.92 ± 0.76 pmol liter$^{-1}$ and mean [Re]$\text{BM}$ = 105 ± 20 ppt, giving a [Re]$\text{diss}$/[Re]$\text{BM}$ = 3.4 ± 1.4 × 10$^{-3}$ (pg g$^{-1}$/pg g$^{-1}$), which is slightly higher than that measured in Taiwan, where the average [Re]$\text{diss}$/[Re]$\text{BM}$ is 3.3 ± 0.5 × 10$^{-3}$. The two mountain belts have comparable physical erosion rates (9, 24), suggesting that Re mobility is similar between sites despite the contrasts in bedrock geology. The [Re]$\text{diss}$/[Re]$\text{BM}$ values are more than double in the Waiho and Fox watersheds that host valley glaciers and have extensive glacial coverage.

To estimate oxidative weathering yields, we quantify the dissolved Re yield (mol km$^{-2}$ year$^{-1}$) in watersheds where we have annual water discharge estimates (41). These are in the Hokitika, Whataroa, Haast, and Waiho rivers (tables S4 and S5). Our river water samples from 2014 cover a relatively narrow dynamic range in water flow especially in the glacial watershed (0.5 to 1.5 times mean flow values) but do not show significant dilution at a higher flow (fig. S2C and table S5). This suggests that the annual water discharge ($Q_{\text{annual}},$ m$^3$ year$^{-1}$ (41) and our mean [Re]$\text{diss}$ for each large watershed can together provide a reliable estimate of dissolved Re flux. However, the average ion concentrations collected at lower $Q_m$ may overestimate the dissolved ion flux, if ions are diluted at a high flow (40). We find that using an average concentration does not systematically overestimate or underestimate the dissolved Re yield (see Materials and Methods). To assess the role of seasonal and/or interannual variability, we plot our major ion data (for example, Ca$^{2+}$) alongside daily water discharge ($Q_{\text{w}},$ m$^3$ s$^{-1}$) for the Whataroa and Hokitika watersheds (fig. S1B) and compare this to published data from 1998 to 1999 (24) and 2000 to 2001 (fig. S2B) (20). The broad consistency in ion concentrations and ion ratios suggest that annual and seasonal variability may be of second-order importance when compared to contrasts between different watersheds. Rainwater and hydrothermal water samples (table S3) have very low [Re]$\text{diss}$ (<0.2 pmol liter$^{-1}$), so no correction is made to the Re flux from these inputs.

**Global compilation**

Published data are compiled from watersheds around the world that allow for a comparison to our findings in New Zealand (table S6). The required data are as follows: (i) dissolved Re concentration ([Re]$\text{diss}$); (ii) watershed-averaged bedrock Re composition, an indication of which is provided by [Re]$\text{BM}$; (iii) annual water discharge to estimate dissolved Re yield; and (iv) suspended sediment yield as a proxy of physical erosion rate (9, 20, 22, 27, 41–45). We also add new measurements of river water and bed material samples from the Jolly and Hooker watersheds draining the eastern Southern Alps, the Waipaoa River in North Island, New Zealand, and the Yukon River and Mackenzie River in Canada, which we collected using similar methods. We estimate the dissolved Re yield for each of these sites and account for variability in the bedrock geology by normalizing the Re yield to the measured [Re]$\text{BM}$. We quantify the upstream area covered by glaciers from published work or by using the World Glacier Inventory as we do for the western Southern Alps (see Materials and Methods). We find dissolved Re yield (normalized by [Re]$\text{BM}$) increases with increasing annual suspended sediment yields across the data set (fig. 2), supporting previous work from Taiwan (9). However, watersheds with mountain glaciation upstream of the sampling locations (where glaciers cover >1% of the area) have a higher dissolved Re yield for a given suspended sediment yield (fig. 2).

**DISCUSSION**

The erosion rates in the western Southern Alps are high (4000 to 10,000 t km$^{-2}$ year$^{-1}$) and comparable to those in Taiwan (9, 24). The similarity in Re mobility between these settings agrees with the notion that high erosion rates can enhance oxidative weathering of OC$_\text{petro}$ and the release of Re to the dissolved load of rivers (9). This probably reflects the rapid soil formation in the western Southern Alps (25) and is consistent with an important role of bedrock landslides for exposing bedrock clasts in landslide deposits and focusing hydrological pathways in landslide scars (26). In the global data compilation, we find that suspended sediment yield (a proxy for physical erosion rate) is correlated with dissolved Re yield (fig. 2). This suggests that OC$_\text{petro}$ oxidation is supply-limited in many locations, as predicted by the relatively fast kinetics of OC$_\text{petro}$ oxidation and the high atmospheric O$_2$ concentrations at present (11, 13). This contrasts with acid hydrolysis weathering of silicate minerals, which is thought to be kinetically limited at the high erosion rates experienced in mountain belts (6, 12). The implication is that for watersheds underlain by sedimentary rocks, increased erosion may result in a less effective CO$_2$ drawdown by silicate weathering, whereas CO$_2$ release by oxidative weathering of OC$_\text{petro}$ continues to increase (fig. 2).

In the western Southern Alps, we find that glacier cover is a major control on the average [Re]$\text{diss}$ measured in watersheds (fig. 1B). This is not only the case for the watersheds with large valley glaciers (the Waiho and Fox; fig. 1A): Glacial processes appear to enhance oxidative weathering to some degree in all watersheds. Although this may seem in conflict with the idea that OC$_\text{petro}$ oxidation in mountains is already supply-limited (9), there are characteristics of glacial watersheds, such as (i) physical mechanisms that increase effective surface area (15, 16) and (ii) biogeochemical mechanisms that increase O$_2$ availability and the competitiveness of microbial communities (17), which mean that for the same physical erosion rate, oxidation rate may be further enhanced. Physical mechanisms include glacial abrasion (16), which can supply large quantities of fine material for weathering within the glacial system and in the deposited moraines. At higher elevations on steep rock walls,
freeze/thaw cycles and frost cracking driven by sustained subzero temperatures and water availability in the porous bedrock can also increase the supply of fresh, fine material to O2 in the air and water (15). Biogeochemical factors work in parallel, with limited vegetation and soil development in glacial watersheds resulting in less demand for O2 by heterotrophic respiration (17). Oxygen could therefore penetrate deeper into exposed rock surfaces (11, 13). Microbial communities also facilitate OCpetro oxidation (1, 14) and are active both subglacially (18, 19) and in moraines colonized by organisms during primary succession (17).

The mechanisms described here are not unique to watersheds of the western Southern Alps but should operate wherever mountain glaciation occurs on OCpetro-bearing rocks. In the glaciated eastern Southern Alps watersheds, erosion rates are lower than those in the western Southern Alps (15, 20, 24), and so lower OCpetro oxidation rates may be expected (9). However, when compared to watersheds with similar erosion rates in unglaciated Taiwan, the glaciated watersheds have higher dissolved Re yields (Fig. 2). The glacier-free Waipaoa River data are consistent with the data from Taiwan. When we examine larger rivers draining OCpetro-bearing sedimentary rocks (table S6), we find that a global pattern starts to emerge (Fig. 2). Erosion rate is a first-order control on oxidative weathering rate (dissolved Re yield), but watersheds hosting glaciers (>1% of the watershed area; for example, Yukon, Brahmputra, Ganges, and Southern Alps) have dissolved Re yields that are up to three times greater for a given erosion rate than watersheds with very low glacial coverage, regardless of the basin area.

The dissolved Re yield can be used to estimate the CO2 oxidation yield, \( J_{\text{CO}_2} \) (g C km\(^{-2}\) year\(^{-1}\)), by OCpetro oxidation (1, 9, 30, 31). First, there must be good constraint on the Re-to-OC ratio of the sedimentary rocks and the behavior of Re and OCpetro during weathering. In the western Southern Alps, the river bed materials provide an estimate of watershed-averaged \([\text{Re}]\) and \([\text{OCpetro}]\), and their compositions are similar to measured bedrocks. The weathered colluvium confirms coupled Re and OCpetro loss (fig. S2 and table S2) (1, 9, 31). To quantify the CO2 release, the dissolved Re yield in grams (\( J_{\text{Re}} \) g km\(^{-2}\) year\(^{-1}\)) is combined with the \([\text{OC}]_{\text{BM}}/[\text{Re}]_{\text{BM}}\) ratio (Eq. 1).

\[
J_{\text{CO}_2} = J_{\text{Re}} \times \left( [\text{OC}]_{\text{BM}}/[\text{Re}]_{\text{BM}} \right) \times f_C \times (1 - f_{\text{graphite}}) \tag{1}
\]

We correct the estimated CO2 release to account for the relative mobility of Re and OCpetro, during weathering in soils, with \( f_C \) being the ratio between percentage loss of OCpetro in soil versus percentage loss of Re in soil. This factor also accounts for the role of sulfide and silicate minerals as trace sources of dissolved Re (9, 27). On the basis of the published data from soils (9, 30, 39) and our measurements from the western Southern Alps (table S3), \( f_C \) is expected to be <1 but >0.5. To account for the presence of graphite, which may not be oxidized, we vary the fraction of OCpetro as graphite (\( f_{\text{graphite}} \)) from 0.5 to 0, informed by measurements from the study location (33). The CO2 oxidation flux, \( J_{\text{CO}_2} \) (g C km\(^{-2}\) year\(^{-1}\)), is calculated using a Monte Carlo simulation to account for these uncertainties (see Materials and Methods).

In the western Southern Alps, watersheds with limited glacial coverage are estimated to release 14 \(^{+10/-2}\) t C km\(^{-2}\) year\(^{-1}\) (Whataroa) by OCpetro oxidation using the Re proxy (table S4). These are similar to OCpetro oxidation yields estimated in Taiwan where erosion rates are similar (9). This suggests that the Re proxy is producing consistent results at the watershed scale. The Re-derived estimate of CO2 flux from OCpetro oxidation for the glaciated Waiho watershed is approximately double, at 30 \(^{+20/-11}\) t C km\(^{-2}\) year\(^{-1}\). These values are similar to seasonal measurements of soil respiration in a primary succession on sedimentary rocks exposed by recent glacial retreat in Svalbard (~10 to 24 t C km\(^{-2}\) year\(^{-1}\), based on monthly averaged data) (46) but lower than typical rates of soil respiration in mineral soils that contain non-rock-derived organic matter (47). Although the uncertainties on the CO2 fluxes are relatively large on the basis of our current understanding of Re and OCpetro mobility, the difference between the glaciated watersheds and the other watersheds is larger than these uncertainties (table S4).

In watersheds where mountain glaciers are confined to headwaters, the Re-derived estimates of CO2 release do not negate CO2 drawdown by silicate weathering [-2 to 10 t C km\(^{-2}\) year\(^{-1}\) (24)] and by erosion and sedimentary burial of biospheric OC [\( J_{\text{CO}_2} \) Fig. 3; ~40 t C km\(^{-2}\) year\(^{-1}\) (22)]. In stark contrast, the doubling of the OCpetro oxidation rate in the Waiho watershed converts it into a net CO2 source during erosion and weathering (Fig. 3). At present, the Waiho and Fox rivers drain less than 5% of the sampled area (Fig. 1A), so the enhanced glacial contribution to CO2 fluxes from OCpetro oxidation has a modest influence across the mountain belt. However, under more heavily glaciated conditions, the Southern Alps would be primed to act as a CO2 source. Accelerated OCpetro oxidation driven by the physical and biogeochemical mechanisms that we have identified may have increased CO2 emissions (Fig. 2). The heavily glaciated western Southern Alps watersheds indicate that biospheric OC erosion and burial will also decrease as glacial cover increases. In addition, at high erosion rates and high rates of mineral supply, silicate weathering rates are very sensitive to temperature and
runoff and cooler temperatures may decrease rates of CO₂ drawdown (Fig. 3).

The global CO₂ emissions by OCpetro oxidation remain to be better quantified. Nevertheless, they are probably as large as those from volcanic degassing (1). To evaluate the strength of this feedback in glaciated mountain belts for counteracting global cooling and for potentially having the capacity to end a glaciation, both the OCpetro presence and abundance in glaciated mountain ranges must be taken into account. By way of example, the Himalaya has the potential to be a key site for CO₂ fluxes by OCpetro oxidation because it hosts OCpetro-bearing Tethyan Sedimentary Series shales at high altitudes. Furthermore, around the world, rocks containing OCpetro have been subject to repeated mountain glaciation throughout the Late Cenozoic. Enhanced OCpetro oxidation in locations such as the Rockies, the Andes, and the European Alps (21) could have driven these sites to operate as important CO₂ sources and O₂ sinks during sustained periods of global cooling and glaciation over 10⁵ to 10⁹ years (5). OCpetro oxidation could act in tandem with transient CO₂ emissions from enhanced sulfide oxidation and weathering of carbonate minerals by sulfuric acid, during glaciation (48). Although the global fluxes are difficult to quantify from the available data (Fig. 2), enhanced OCpetro oxidation associated with more extensive glacial erosion processes is consistent with the ~2% decline in atmospheric O₂ recorded in ice cores over the past 800,000 years (5). If the decline in atmospheric O₂ results from changes in OC burial versus OCpetro oxidation alone, it implies a net CO₂ release of ~3 × 10¹³ mol year⁻¹ or ~3 to 4 × 10¹⁶ tC year⁻¹ (5). This corresponds to only a modest (~6%) increase in global OCpetro oxidation rates (1) over this period in which the duration and intensity of glaciation increased (21). We propose that the link between OCpetro weathering and mountain glaciation offers a previously unrecognized feedback between climate and the carbon cycle, where increased CO₂ emissions act to counter further global cooling during the Late Cenozoic.

**MATERIALS AND METHODS**

**Sample collection**

Samples were collected from 13 watersheds in the western Southern Alps, which drain to the west of the main divide, two draining the eastern Southern Alps to the east of the main divide and the Waipaoa River in North Island, New Zealand (tables S1 to S3). Together, these watersheds allowed us to examine the dual roles of physical erosion rate (9) and glacial coverage on OCpetro weathering. The Southern Alps is a steep mountain belt built by transpression along the Alpine Fault. The western flank has a temperate climate, with a high erosion rate driven by orographic precipitation (exceeding 8 m year⁻¹), steep slopes, and bedrock landslides, which expose OCpetro in metasedimentary rocks (34). Previous work has documented high rates of silicate and carbonate weathering in the western Southern Alps (20, 24). Erosion dominates the denudation, with the largest percentage of chemical denudation occurring in the Haast watershed (5%) (24). Along the western Southern Alps, the metamorphic grade varies perpendicular to the Alpine Fault strike, but the sedimentary protolith is similar in all watersheds, and the OCpetro content ranges from ~0.1 to ~0.2% (22, 33). In contrast, there are significant differences in glacial coverage (Fig. 1A), including two watersheds with large valley glaciers (Waipaoa and Fox) and some frontal watersheds with very minimal glacial coverage (for example, Waitangitaoa). The eastern Southern Alps is also dominated by glacial processes but experiences lower precipitation (<2 m year⁻¹), lower rates of bedrock landsliding (15), and slower long-term exhumation rates (49). As a result, the chemical denudation rates are also lower (50). The Waipaoa River in North Island, New Zealand drains lower metamorphic grade OCpetro-bearing sedimentary rocks at high erosion rates but lacks glacial influence (51, 52).

To assess OCpetro oxidation in the western Southern Alps, we sampled the dissolved products of chemical weathering, weathered colluvium on hillslopes, and relatively unweathered river bed materials of sand and finer grade. River waters (n = 51) were collected from the center of river channels at their surface. Watersheds were sampled two to six times over 1 month (14 September 2014 to 03 October 2014) at variable flow to examine the hydrological variability of the dissolved ions released from chemical weathering (40). Water samples were decanted to sterile plastic containers before filtration through 0.2-μm polyethersulfone filters with a diameter of 142 mm within a day of collection and stored in acid-cleaned low-density polyethylene bottles. Alkalinity measurements were made by Gran titration on an aliquot of filtered water. All water samples intended for cation and Re analysis were acidified in the field to pH ~2 (9, 31), with an unacidified aliquot for anion analyses. Two 250-ml rainwater samples were collected over 10-hour periods (table S1).

In the western Southern Alps, soils are thin and weathering profiles are often poorly developed (25). Landslide-derived colluvium is an important locus of weathering (26). Therefore, to characterize Re and OCpetro behavior in the weathering zone, we collected ~500-cm⁻²-sized bulk samples of weathered colluvium at discrete depths between 10 and 70 cm below the soil surface at three sites on the forested hillslopes of Alex Knob, which drains to the Docherty Creek watershed. We also collected surface soil samples from the upper 3 cm that comprised a mixture of litter and mineral soil (n = 5) using a metal trowel and transferred samples to sterile plastic bags (table S3).

River bed material samples (n = 31) were also collected to help constrain the composition of unweathered materials (tables S2 and S3) (9). Samples were taken from channel edges or bank deposits that were taken to represent the sand-to-silt fraction deposited on the river bed during recent flow regimes and transferred to sterile plastic...
bags. River water samples and bed materials were also collected from the eastern Southern Alps and Waipaoa River using these methods (table S2).

**Geochemical analyses**

Dissolved Re concentrations in river water samples ([Re]_{Diss} pmol liter\(^{-1}\)) were measured by isotope dilution quadrupole ICP-MS (Q-ICP-MS) in conjunction with anion exchange column chemistry to preconcentrate and purify Re. Between 30 and 500 ml of water samples was doped with a known amount of tracer solution consisting of enriched \(^{185}\)Re and evaporated to dryness, with the dried sample dissolved in HNO\(_3\) before anion exchange column chemistry.

Solid samples (surface soil, weathered colluvium, and river bed materials) were ground to a fine powder before acid digestion to generate an integrated bulk sample. Homogenizing bedload samples permitted assessment of the average Re composition of the rocks at the watershed scale for each river because fluvial transit times are short. A known weight of powder (0.5 g) was doped with a known amount of \(^{185}\)Re spike and digested in a 63 HF-HNO\(_3\) mix (9 ml) for 24 hours at 120°C and then evaporated. The dried sample was further digested in a 2:1 mix of HNO\(_3\)-HCl (3 ml) for 24 hours at 120°C and then evaporated. Re was isolated and purified using a NaOH-acetone solvent extraction methodology (53).

The Re isotope composition of the purified Re aliquots were determined in a 0.8 N HNO\(_3\) medium using a Thermo Fisher Scientific X-Series Q-ICP-MS at Durham University. The procedural blank was \(<1%\) of the lowest concentration samples. Uncertainties in the Re abundance were determined by error propagation of uncertainties in Re mass spectrometry measurements, blank abundance and isotopic compositions, spike calibrations, and reproducibility of standard Re isotopic values. Repeat analyses of [Re]_{Diss} in a river water standard, SLRS-5, gave a concentration of 59.8 ± 1.7 ppt (n = 12, ±2 SE), in agreement with the previously reported value of 66 ± 12 ppt (54).

Major ion concentrations in water samples were analyzed by ion chromatography. Cation and anion standards and a certified reference standard (LETHBRIDGE-03) were run to validate the analytical results. The HCO\(_3^−\) concentration was estimated using total alkalinity, temperature, and pH (measured in the field) data inputted to CO2SYS (55). The charge balance of dissolved cations ([TZ′] = Na\(^+\) + K\(^+\), 2Mg\(^{2+}\) + 2Ca\(^{2+}\)) and dissolved anions ([TZ′] = Cl\(^−\) + HCO\(_3^−\) + 2SO\(_4^{2−}\)) was determined ([TZ′ − TZ])/([TZ′ + TZ]) as a measure of data quality (20). This was 11% across all samples, within the combined uncertainty and similar to previous work in this location (20).

In river bed materials, surface soils, and weathered colluvium, the OC concentration ([OC], %) was measured following a 0.2 N HCl leach protocol (56), which was tested on samples from this location to ensure full removal of detrital carbonates. Aliquots of samples were combusted, protocol (57), and similar to previous work in this location ([OC]_{Det} %) was measured following a 0.2 N HCl leach protocol (56). The HCO\(_3^−\) concentration was measured by ion chromatography. Cation and anion standards and a certified reference standard (LETHBRIDGE-03) were run to validate the analytical results. The procedural blank was \(<1%\) of the lowest concentration samples. Uncertainties in the Re abundance were determined by error propagation of uncertainties in Re mass spectrometry measurements, blank abundance and isotopic compositions, spike calibrations, and reproducibility of standard Re isotopic values. Repeat analyses of Re and [Ca\(^{2+}\)]_a and [SO\(_4^{2−}\)]_a were determined using a Costech elemental analyzer coupled to a Thermo Fisher Scientific Delta V Advantage isotope ratio mass spectrometer at Durham University. The procedural blank was \(<1%\) of the lowest concentration samples. Uncertainties in the Re abundance were determined by spike calibrations and reproducibility of standard Re isotopic values. Repeat analyses of [Re]_{Diss} in a river water standard, SLRS-5, gave a concentration of 59.8 ± 1.7 ppt (n = 12, ±2 SE), in agreement with the previously reported value of 66 ± 12 ppt (54).

**OCpetro content of weathered colluvium**

To assess the OCpetro content of the weathered colluvium samples and to account for OC derived from recent productivity (biospheric OC), we adopted a mixing analysis based on observations of \(^{14}\)C and F_{Mod} values. The \(^{14}\)C activity of sedimentary rocks is generally considered to be below the analytical background, that is, F_{Mod} = 0, and thus distinct from modern biospheric OC (F_{Mod} ~ 1) and degraded soil of ~1000 years old (F_{Mod} ~ 0.9). The \(^{13}\)C of OCpetro in the Southern Alps was \(^{13}\)C-enriched (\(^{13}\)C ~ 21 to –22‰) compared to the terrestrial biosphere (dominated by C\(_3\) plants) (22).

The stable isotope composition of an element shown by the reciprocal of its concentration can reveal mixing trends or processes that alter the concentration and fractionate isotopes. Surface soil litters are OC-rich and \(^{13}\)C-depleted, and they describe a linear trend between \(^{13}\)Corg and 1/[OC], albeit one that only describes ~40% of the variability in the data. This is consistent with the degradation of plant-derived OC in surface soils, loss of OC, and enrichment in \(^{13}\)C (fig. S2A). In contrast, the weathered colluvium samples define a different linear trend, which we interpret as a mixture of degraded biospheric organic material (originally derived from the surface soil) with \(^{13}\)C-enriched, OC-poor material from the sedimentary rocks (fig. S2B). The intercept of these two trends implies that the degraded soil OC has a value of ~26.8 ± 0.8‰ (propagating the 95% uncertainty bounds on the linear trends).

The weathered colluvium samples are also \(^{14}\)C-depleted (fig. S2B), which is consistent with OCpetro addition. The samples can be described by a linear trend between F_{Mod} and \(^{13}\)Corg that intercepts biospheric and petrogenic OC (fig. 2B). Using the “degraded soil” \(^{13}\)Corg value (fig. S2A) and the trend defined by the samples, we estimated the F_{Mod} of the biospheric OC in these samples as F_{Mod} = 0.93 ± 0.36. These values and their uncertainties were used in a two-component end member mixing model (51) to quantify the fraction of OCpetro (f_{petro}) and the corresponding [OC]petro (f_{petro} × [OC]) (table S3). This approach does not consider additional aging of biospheric OC, which would act to reduce the F_{Mod} of the biospheric OC. We therefore calculated an upper bound on the [OC]petro and a lower bound on the loss of OCpetro during weathering. The weathered colluvium samples had an average [OC]petro = 0.15 ± 0.06% (n = 4, ±2 SE), which is lower than the local river bed materials in this watershed ([OC]_{Det} = 0.21%) (table S3). Coupled loss of Re and OCpetro during chemical weathering supports measurements on soils from the Ohio Shale (table S1) (30), the Himalaya (39), and Taiwan (9). These data sets generally show that Re is more mobile during weathering, and so the dissolved Re flux must be corrected when estimating an OCpetro weathering rate (9).

**Dissolved Re flux**

To assess whether the average of the ion concentrations collected at low Q_{w} may overestimate the dissolved ion flux, we compared the flux calculated using an average concentration method to that calculated by a rating curve method (taking into account dilution) using published data sets from the study location (24). For these data, the average measured [Ca\(^{2+}\)] multiplied by the mean annual runoff returns a Ca\(^{2+}\) flux of \(4.21 \times 10^8\) mol year\(^{-1}\) for the Hokiitaka River. If we model the [Ca\(^{2+}\)]-Q_{w} relationship (fig. S1B) as a power law rating curve (57) and apply it to the daily Q_{w} data from 1971 to 2015, the annual Ca\(^{2+}\) flux was \(4.39 \times 10^8\) mol year\(^{-1}\). The methods agree within 4%, with the average concentration method slightly underestimating flux. In comparison, for the SO\(_4^{2−}\) flux, which also shows dilution with Q_{w} in published data (24), the difference in estimated fluxes is 3%. Assuming similar dilution trends for [Ca\(^{2+}\)] as for [Ca\(^{2+}\)] and [SO\(_4^{2−}\)], which is suggested on the basis of

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the available data (fig. S2C), results in a <5% underestimation of flux. Although a longer time series sampling would be informative for tracking the dissolved ion source and linking it to hydrological pathways (40, 57), an average concentration with an accuracy of within ~5% is adequate for calculating the flux.

Quantification of OCpetro oxidation rate and its uncertainty

To convert the dissolved Re flux to an estimate of CO2 release, there are known uncertainties related to the behavior of Re and OCpetro during weathering (1, 9). The main uncertainties in this setting are as follows: (i) Because Re is a soluble element, oxidative weathering may mobilize Re more effectively during early soil formation, meaning that some Re measured in river waters does not correspond to release of CO2 at the weathering site (30); (ii), partly related to (i), some dissolved Re may come from sulfide and silicate minerals (9, 27); and (iii) graphitic OCpetro may not be susceptible to oxidation and remain in the soil (32, 58). Regarding point (i) and (ii), soils from Taiwan suggest congruent dissolution of Re, and loss of OCpetro can occur during weathering of young, thin soils (9). On a soil profile developed on OCpetro-rich shale from Ohio (30), Re depletion reaches 100% in the most highly weathered soil, whereas OCpetro loss is ~70% (table S1). The data from weathered colluvium in the western Southern Alps are more consistent with the Ohio Shale than Taiwan. For point (iii), graphite is present in rocks within a kilometer of the Alpine Fault, making up almost 50% of the OC (33), and its measured abundance decreases to small amounts ~10 to 20 km from the Alpine Fault (59).

To account for and quantify the uncertainties on the CO2 flux estimate we used a Monte Carlo simulation. This includes uncertainty on the dissolved Re flux, the measured variability in [OC]BM/[Re]BM, and the assumptions (i), (ii), and (ii) above. For each watershed (area = A km2), we used ±2 SE on the mean for the [Re]diss and the [OC]BM/[Re]BM ratio. To account for the relative mobility of Re and OCpetro during weathering (30), we defined f2 as the ratio between the percentage loss of OCpetro in soil compared to bedrock and the percentage loss of Re in soil compared to bedrock. On the basis of published work, we varied f2 from 0.5 to 1 (9, 30). To account for the presence of graphite, which may not be oxidized, we varied the fraction of OCpetro as graphite (fgraphite) from 0.5 to 0, informed by measurements from the study location (33). The CO2 oxidation flux, fCCO2 (g C km−2 year−1), was then calculated using Eq. 1.

The Monte Carlo simulation was run 100,000 times for each watershed, with a “full probability” distribution for each variable. We reported the median value ± 1 SD range. This reflects the present state of knowledge of the Re proxy for OCpetro oxidation. Future work should seek to refine this approach and reduce the uncertainties, for a better understanding of f2 and fgraphite. By doing so, it may be possible to quantify watershed-scale fluxes using Re measurements more widely (1)

Quantification of glacier cover using the World Glacier Inventory

The relative importance of glacial processes in the watersheds of the western Southern Alps was quantified using glacier locations and areas from the World Glacier Inventory (23). The total sum of glacier area (km2) was quantified for each watershed using ArcGIS, with flow routing algorithms to isolate drainage areas (table S4). In the global data set, published estimates were used (table S6).

Biospheric OC erosion and burial

Previous work has estimated the erosion rate of biospheric OC in watersheds of the western Southern Alps (22). To estimate how this contributes to CO2 drawdown, we required an estimate of the burial efficiency of OC. On the basis of a recent global compilation (44), the sediment yield to an offshore basin plays an important role in setting the burial efficiency of OC. This is because sediment accumulation rate is a first-order control on OC burial efficiency in many marine environments (60). Using the sediment yield of ~6000 to 10,000 t km−2 year−1 for the western Southern Alps (22), the burial efficiency would be predicted to be from ~40% to 100% based on the global data set (44). A high burial efficiency would be consistent with the very high preservation potential of terrestrial palynomorphs offshore from the western Southern Alps (61). To provide a conservative estimate of biospheric OC burial, we used the lowest value in this range (40%) and multiplied the erosional export flux (22) by the OC burial efficiency to estimate CO2 drawdown by biospheric OC erosion as shown in Fig. 3.
18. M. Tranter, M. J. Sharp, H. R. Lamb, G. H. Brown, B. P. Hubbard, I. C. Willis, Geochemical weathering at the bed of Haut Glacier d’Arolla, Switzerland—A new model. Hydrol. Process. 16, 959–993 (2002).

19. J. L. Wadham, S. Bottrell, M. Tranter, R. Raiswell, Stable isotope evidence for microbial sulphate reduction at the bed of a polythermal high Arctic glacier. Earth Planet. Sci. Lett. 219, 341–355 (2004).

20. A. D. Jacobson, J. D. Blum, C. P. Chamberlain, D. Craw, P. O. Koons, Climatic and tectonic controls on chemical weathering in the New Zealand Southern Alps. Geochim. Cosmochim. Acta 67, 29–46 (2003).

21. F. Herman, D. Seward, P. G. Valla, C. A. Carter, B. Kohn, S. D. Willett, T. A. Ehlers, Worldwide acceleration of mountain erosion under a cooling climate. Nature 504, 423–426 (2013).

22. R. G. Hilton, A. Galy, N. Hovius, Riverine particulate organic carbon from an active mountain belt: Importance of landslides. Global Biogeochim. Cycles 22, GB1017 (2008).

23. National Snow and Ice Data Center, “World Glacier Inventory” (2016); http://nsidc.org/.

24. W. B. Lyons, A. E. Carey, D. M. Hicks, A. A. Nezat, Chemical weathering in high-sediment-yielding watersheds, New Zealand. J. Geophys. Res. Earth Surf. 110, F01008 (2005).

25. I. J. Larsen, P. C. Almond, A. E. Gey, H. R. D. Montgomery, B. Malcolm, Rapid soil production and weathering in the Southern Alps. New Zealand. Science 334, 637–640 (2011).

26. R. Embrey, N. Hovius, A. Galy, O. Marc, Chemical weathering in active mountain belts controlled by stochastic bedrock landsliding. Nat. Geosci. 9, 42–45 (2016).

27. C. A. Miller, B. Peucker-Ehrenbrink, B. D. Walker, F. Marcattonio, Re-assessing the surface cycling of molybdenum and rhenium. Geochim. Cosmochim. Acta 75, 7146–7179 (2011).

28. D. Selby, R. A. Creaser, Re-Os geochronology of organic rich sediments: An evaluation of organic matter analysis methods. Chem. Geol. 200, 225–240 (2003).

29. D. Colodner, J. Sachs, G. Ravizza, K. Turekian, J. Edmond, E. Boyle, The geochemical cycle of rhenium: A reconnaissance. Earth Planet. Sci. Lett. 117, 205–221 (1993).

30. L. A. Jaffe, B. Peucker-Ehrenbrink, S. T. Petch, Mobility of rhenium, platinum group elements and organic carbon during black shale weathering. Earth Planet. Sci. Lett. 198, 339–353 (2002).

31. T. K. Dalai, S. K. Singh, J. R. Trivedi, S. Krishnaswami, Dissolved rhenium in the Yamuna river system and the Ganga in the Himalaya: Role of black shale weathering on the budgets of Re, Os and U in rivers and CO2 in the atmosphere. Geochim. Cosmochim. Acta 66, 29–43 (2002).

32. V. Galy, O. Bysseck, B. Peucker-Ehrenbrink, T. P. Brabets, B. Wang, R. H. Meade, “Environmental and hydrologic overview of the Yukon River basin, Alaska and Canada,” in Geological and Hydrologic Overview of the Yukon River Basin, Alaska and Canada, edited by T. P. Brabets, B. Wang, R. H. Meade, T. J. Hughes, and N. J. Koerner (USGS Water Resources Investigations Report 07-4230, 2008).

33. L. Nibourel, F. Herman, S. C. Cox, O. Beyssac, J. Lavé, Provenance analysis using Raman spectroscopy of carbonaceous material: A case study in the Southern Alps of New Zealand. J. Geophys. Res. Earth Surf. 120, 2056–2079 (2015).

34. A. Pierson-Wickmann, L. Reisberg, C. France-Lanord, Behavior of Re and Os during weathering of organic rich sediments: An evaluation of CO2 consumption in the New Zealand Southern Alps. Geochim. Cosmochim. Acta 70, 3128–3142 (2006).

35. L. J. Reusser, P. R. Bierman, Using meteoric 10Be to track fluvial sand through the World Glacier Inventory (2016); http://nsidc.org/.

36. M. Dellinger, J. Gaillardet, J. Bouchez, O. Beyssac, J. Gaillardet, C. France-Lanord, L. Maurice, P. Moreira-Turcq, Determination of total organic carbon content and δ13C in carbonate-rich detrital sediments. Geostand. Geoanal. Res. 28, 37–46 (2004).

37. E. R. Lewis and D. W. R. Wallace, Program Developed for CO2 System Calculations (ORNL/CDIAC-105, 1998).

38. V. Galy, J. Bouchet, C. France-Lanord, Determination of total organic carbon content and 13C in carbonate-rich detrital sediments. Geostand. Geoanal. Res. 31, 205–207 (2007).

39. E. T. Tipper, M. J. Bickle, A. Galy, A. J. West, C. Pomies, H. J. Chapman, The short term climatic sensitivity of carbonate and silicate weathering fluxes: Insight from seasonal variations in river chemistry. Geochim. Cosmochim. Acta 70, 2737–2754 (2006).

40. J. Bouchet, O. Bysseck, V. Galy, J. Gaillardet, C. France-Lanord, L. Maurice, P. Moreira-Turcq, Oxidation of petrogenic organic carbon in the Amazon floodplain as a source of atmospheric CO2. Geochim. Cosmochim. Acta 35, 258–259 (2010).

41. O. Bysseck, S. C. Cox, J. Vry, F. Herman, Peak metamorphic temperature and thermal history of the Southern Alps (New Zealand). Tectonophysics 676, 229–249 (2016).

42. N. E. Blair, R. C. Aller, in Annual Review of Marine Science, C. A. Carlson, S. J. Giovanni, Eds. (Annual Reviews, 2012), vol. 4, pp. 401–423.

43. M. T. Ryan, R. M. Newingham, G. B. Dunbar, M. J. Vandergoes, A. B. H. Rees, H. Neil, S. L. Callard, B. V. Alloway, H. Bostock, Q. Hua, B. M. Anderson, Exploring the source-to-sink residence time of terrestrial pollen deposited offshore Westland, New Zealand. Rev. Palaeoecol. Polyso. 230, 37–46 (2016).

44. S. E. Tank, P. A. Raymond, R. G. Sjølie, J. W. Mcclerran, R. M. Holmes, G. Fiske, B. J. Peterson, A land-to-ocean perspective on the magnitude, source and implication of DIC flux from major Arctic rivers to the Arctic Ocean. Global Biogeochim. Cycles 26, GB4018 (2012).

45. W. W. Immerzeel, M. F. P. Bierkens, Seasonal prediction of monsoon rainfall in three Asian river basins: The importance of snow cover on the Tibetan plateau. Int. J. Climatol. 30, 1835–1842 (2010).

46. A.-C. Pierson-Wickmann, L. Reisberg, C. France-Lanord, The Os isotopic composition of Himalayan river bedloads and bedrocks: Importance of black shales. Earth Planet. Sci. Lett. 176, 203–218 (2000).

47. Y. Huh, J.-L. Birck, C. J. Allègre, Osium isotope geochemistry in the Mackenzie River basin. Earth Planet. Sci. Lett. 222, 115–129 (2004).

48. T. P. Brabets, B. Wang, R. H. Meade, “Environmental and hydrologic overview of the Yukon River basin, Alaska and Canada,” in Geological and Hydrologic Overview of the Yukon River Basin, Alaska and Canada, edited by T. P. Brabets, B. Wang, R. H. Meade, T. J. Hughes, and N. J. Koerner (USGS Water Resources Investigations Report 07-4230, 2008).

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