1Title: Microbial oxidation of lithospheric organic carbon in rapidly eroding2tropical mountain soils

Authors: Jordon D. Hemingway^{1,2,*,†}, Robert G. Hilton³, Niels Hovius^{4,5}, Timothy I. Eglinton⁶,
 Negar Haghipour⁶, Lukas Wacker⁷, Meng-Chiang Chen⁸, Valier V. Galy¹

5 Affiliations:

- ¹Woods Hole Oceanographic Institution, Department of Marine Chemistry and Geochemistry,
 266 Woods Hole Road, Woods Hole MA 02543, USA.
- ⁸ ²Massachusetts Institute of Technology Woods Hole Oceanographic Institution Joint Program
- 9 in Oceanography and Applied Ocean Science and Engineering, 77 Massachusetts Avenue, Cam-
- 10 bridge MA 02139, USA.
- ¹¹ ³Durham University, Department of Geography, South Road, Durham DH1 3LE, UK.
- ¹² ⁴GFZ German Research Center for Geoscience, Telegrafenberg, Potsdam 14473, Germany.
- ¹³ ⁵Department of Earth and Environmental Sciences, University of Potsdam, Karl-
- 14 Liebknechtstraße 24, Golm 14476, Germany.
- ⁶ETH Zürich, Geological Institute, Department of Earth Sciences, Sonneggstrasse 5, Zürich
 8092, Switzerland.
- ¹⁷ ⁷ETH Zürich, Laboratory of Ion Beam Physics, Department of Physics, Otto-Stern-Weg 5,
- 18 Zürich 8092, Switzerland.
- ¹⁹ ⁸Taroko National Park Headquarters, Fu-Su Village, Hualien 972, Taiwan.
- 20 *Correspondence to: jordon_hemingway@fas.harvard.edu.
- ²¹ [†]Current address: Harvard University, Department of Earth and Planetary Sciences, 20 Oxford
- 22 Street, Cambridge MA 02138, USA.
- 23 Abstract: Lithospheric organic carbon ("petrogenic"; OC_{petro}) is oxidized during exhumation and
- subsequent erosion within mountain ranges. This process is a significant source of CO_2 to the
- 25 atmosphere over geologic timescales, but the mechanisms that govern oxidation rates in
- mountain landscapes remain poorly constrained. We demonstrate that, on average, 67 ± 11 % of
- 27 OC_{petro} initially present in bedrock exhumed from the tropical, rapidly eroding Central Range of
- Taiwan is oxidized within soils, leading to CO_2 emissions of 6.1 18.6 t C km⁻² yr⁻¹. The
- 29 molecular and isotopic evolution of bulk OC and lipid biomarkers during soil formation reveals
- that OC_{petro} remineralization is microbially mediated. Rapid oxidation in mountain soils drives
 CO₂ emissions fluxes that increase with erosion rate, thereby counteracting CO₂ drawdown by
- 32 silicate weathering and biospheric OC burial.
- 33 **One Sentence Summary:** Oxidation of lithospheric organic carbon in eroding mountain soils is
- rapid and microbially mediated, and resulting CO₂ emissions counteract CO₂ drawdown by
- 35 silicate weathering and biospheric organic carbon burial.

Main Text: Erosion-induced weathering in collisional mountain belts is a major carbon-cycle 36 regulator over million-year timescales and provides a link between tectonics and climate (1, 2). 37 Atmospheric CO₂ is consumed by the export and burial in marine sediments of biospheric 38 organic carbon (OC_{bio}) and carbonate minerals precipitated following silicate rock weathering 39 (1). The CO₂ drawdown flux associated with both processes increases with erosion rate (3, 4), 40 highlighting the importance of steep, erosive orogens in driving CO₂ drawdown. By comparison, 41 42 CO₂ release during exhumation and erosion has received considerably less attention despite its potential to partially or fully negate the effects of geological CO_2 consumption (1, 5, 6). 43 Oxidative weathering of both sulfide minerals (coupled with carbonate dissolution) and 44 petrogenic organic carbon (OC_{petro}) contained in exhumed rocks can increase atmospheric CO₂ 45 and decrease O_2 concentrations over geologic timescales (1, 7-9). Still, the mechanisms that 46 47 govern oxidation rates and efficiencies in mountain belts remain under-constrained (5, 8, 9). To better constrain orogenic CO_2 emissions, we assess the controls on OC_{petro} oxidation 48 and export within the Central Range of Taiwan, one of the fastest exhuming and eroding 49 mountain belts on Earth (10). Steep relief (11), frequent typhoon landfall (10), and high bedrock 50 landslide rates (11) lead to long-term erosion rates of 3 - 6 mm yr⁻¹ across the range (10). While 51 supplemental contributions from deeper in the exhumation path are likely, weathering in such 52 mountain landscapes occurs primarily on hillslopes and in colluvial deposits (12, 13). We 53 54 therefore assess OC molecular and isotopic evolution within multiple hillslope soil profiles located in the LiWu and WuLu River basins (Fig. S1) and verify these observations at the 55 catchment scale using LiWu River suspended sediments (14). Soils at our study sites are ≤ 1 m 56 thick, including mineral (A+E) and saprolite (C) layers (15), experience residence times on the 57 order of centuries (14), and overlay bedrock ranging from Mesozoic greenschist and amphibolite 58 at low elevations (Tananao schists) to Cenozoic slate and phyllite near the Lishan Fault (Pilushan 59 and Lushan formations) (16). All lithologies are carbonaceous, with bedrock outcrops containing 60 0.2 - 0.7 % OC_{petro} (Table S1) (17). 61 Significant OC_{petro} loss is observed in all soil profiles, as evidenced by the relationship

62 Significant OC_{petro} loss is observed in all soil profiles, as evidenced by the relationship 63 between soil OC content (% OC_{soil}) and ¹⁴C activity (expressed as "fraction modern" or Fm) 64 (14). To account for differences in % OC between bedrock lithologies (17), % OC_{soil} is expressed 65 as

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$$\Delta$$
%OC = % OC_{soil} - % OC_{bedrock},

(1)

67 where % OC_{bedrock} is the OC content of bedrock immediately underlying each soil sample. The 68 average fraction of bedrock OC that is oxidized during soil formation, f_{ox} , can then be quantified 69 by utilizing the fact that OC_{petro} is inherently ¹⁴C-free (Fm_{petro} = 0.0) and setting Fm_{bio} = 1.045 ± 70 0.079, the measured ¹⁴C activity of vascular plant leaf-wax fatty acids extracted form A+E 71 horizon soils (**Table S2**) (14). Soil OC is treated as a mixture of OC_{bio} and residual OC_{petro}, 72 leading to the equation (14):

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$$\operatorname{Fm}_{\text{soil}} = \operatorname{Fm}_{\text{bio}} \left[\frac{\Delta\% \operatorname{OC} + (f_{\text{ox}})(\% \operatorname{OC}_{\text{bedrock}})}{\Delta\% \operatorname{OC} + \% \operatorname{OC}_{\text{bedrock}}} \right].$$
(2)

Fm_{soil} is a hyperbolic function of Δ %OC with curvature that is defined by both %OC_{bedrock} and

- f_{ox} , as shown in <u>Fig 1</u>. We simultaneously solve <u>Eq. 2</u> for the best-fit % OC_{bedrock} and f_{ox} values
- ⁷⁶ using orthogonal distance regression and account for uncertainty using Monte Carlo resampling
- 77 *(14)*.

78 On average, $67 \pm 11 \% (\pm 1\sigma)$ of bedrock OC is lost during soil formation, a minimum estimate since deep weathering has likely already removed OC from initial bedrock (18). To test 79 80 if observed % OC trends simply reflect mobile element losses during weathering and not oxidation per se, we solve Eq. 2 for a subset of samples after normalizing OC content to the 81 immobile element titanium (Table S1) (14). Calculated f_{ox} values using normalized and un-82 normalized data are identical within uncertainty, indicating no appreciable mobility effect on our 83 84 results (Fig. S2). Assuming that all OC lost is oxidized to $CO_2(8)$, f_{ox} can be used to estimate the steady-85

state CO₂ emission flux from soils due to OC_{petro} oxidation, termed Φ_{ox} , according to

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$$\Phi_{\text{ox}} = \frac{(f_{\text{ox}})(\% \text{ OC}_{\text{bedrock}})(\rho_{\text{soil}})(z_{\text{soil}})}{\tau_{\text{soil}}},$$
(3)

where ρ_{soil} is the soil density, z_{soil} is the soil thickness (15), and τ_{soil} is the soil residence time on 88 hillslopes. We estimate τ_{soil} using three independent methods (landslide rates, catchment-average 89 denudation rates, OCbio erosion rates) and incorporate uncertainty for each variable in Eq. 3 90 using Monte Carlo resampling across the range of observed values (14), resulting in a median 91 Φ_{ox} range of 6.1 – 18.6 t C km⁻² yr⁻¹ for conditions prevalent across the Central Range (Fig. 92 S3A) (14). We emphasize that Φ_{ox} is a minimum estimate of total CO₂ emissions by OC_{petro} 93 oxidation due to the potential for OC losses occurring during deep weathering (18). Still, this 94 flux is statistically identical to two independent, catchment-integrated OC_{petro} oxidation estimates 95 for Taiwanese rivers based on fluvial OC_{petro} export ($\leq 12 \text{ t C km}^{-2} \text{ vr}^{-1}$) (19) and dissolved 96 rhenium yield $(7 - 13 \text{ t C km}^{-2} \text{ yr}^{-1}; \text{ Fig. S3B})$ (5) and is 2- to 6-fold higher than estimates of CO₂ 97 98 drawdown by silicate weathering in the LiWu catchment (Fig. S3C) (18). The observation that 99 Φ_{ox} matches catchment-integrated emissions implies that OC_{petro} oxidation in Taiwan occurs 100 predominantly within rapidly eroding hillslope soils.

101 A saprolite depth profile collected from the WuLu catchment indicates that bedrock OC can be oxidized and replaced with OC_{bio} before A+E horizons have fully developed. Two 102 samples collected at 0.5 m and 0.2 m depth contain similar OC concentrations (0.20 %, 0.28 %, 103 respectively) but drastically different Fm values (0.108, 0.839, respectively; Table S1). Rapid 104 OC_{petro} oxidation can occur (i) abiotically without chemical alteration, (ii) abiotically with 105 chemical alteration, (iii) biotically without chemical alteration, or (iv) biotically with chemical 106 alteration and ¹⁴C-depleted biomass production (20-22). To assess alteration and to track 107 108 multiple OC sources within a single sample, we utilize Ramped PyrOx (RPO) serial combustion (23). This technique heats each sample at a constant ramp rate to separate OC based on thermal 109 lability and determines Fm values for specific temperature intervals (termed RPO fractions) (14). 110 To quantitatively compare OC chemical structure, we determine the underlying thermal 111 activation energy (E) distribution for each sample, termed p(0,E), as this is an intrinsic property 112 of carbon bond strength and thus a proxy for chemical composition (23). Unlike ¹⁴C activity, 113 end-member mixing does not shift OC in E space. Mixing OCbio with unaltered OCpetro will thus 114 result in a bimodial p(0,E) distribution, whereas chemical alteration is required to explain the 115 presence of intermediate E values (14, 23). 116 We constrain bedrock E using particulate OC (POC) from 27 suspended sediment 117

samples, including isolated ≥ 2 mm clasts, collected from the LiWu River during four typhoon events (14). Because sediment exported during typhoons is dominated by material sourced from

bedrock incision, distributed runoff erosion, and landsliding throughout the basin (11, 12), we 120 expect this sample set to integrate outcropped bedrock lithologies that contain relatively 121 unweathered OC_{petro}. This is supported by bulk POC ¹³C content (expressed as δ^{13} C values) and 122 total nitrogen to POC ratios (Table S3), which span the range of Tananao schist, Lushan 123 formation, and Pilushan formation values (17). Fig. 2A shows that bedrock OC is exclusively 124 associated with $E \ge 185$ kJ mol⁻¹ (termed high-E; Fig. S3A) (14), consistent with the observed 125 partial graphitization of this material (16). We additionally constrain vascular-plant OC p(0,E)126 using two organic-rich (\geq 5 %) surface soils characterized by bulk Fm values similar to those of 127 plant-wax fatty acids (14). For both samples, ≥ 90 % of OC is associated with E < 150 kJ mol⁻¹ 128 (termed low-*E*), indicating that OC_{bio} and OC_{petro} are effectively separated in *E* space. 129 Energy distributions and ¹⁴C activity in soil and saprolite materials provide strong 130 evidence for OC_{petro} chemical alteration during weathering. Up to 51 % of OC contained in 131 saprolites and deep A+E horizons lies between 150 and 185 kJ mol⁻¹ (termed mid-*E*; **Table S4**; 132 **Fig. S4B-C**); higher than that corresponding to vascular-plant OC ($\leq 150 \text{ kJ mol}^{-1}$) yet lower 133 than bedrock OC (\geq 185 kJ mol⁻¹). This observation can result from either (i) increasing 134 vascular-plant OC E by stabilization during aging in soils (24) or (ii) decreasing residual OC_{petro} 135 *E* during oxidative weathering (20, 21). We assess the relative importance of these mechanisms 136 using the ¹⁴C activity of each RPO fraction (Table S5). As shown in Fig. 2B, low-*E* Fm values 137 cluster near those of vascular-plant fatty acids, whereas high-E material approaches Fm of zero. 138 Meanwhile, mid-E OC spans an Fm range from 0.083 ± 0.002 to 0.912 ± 0.008 . We rule out the 139 possibility that ¹⁴C-depleted mid-E OC exclusively reflects OC_{bio} aging because (i) this would 140 require a biospheric component that has aged up to 20,000 ¹⁴C yr, significantly longer than the 141 centennial soil residence times in Taiwan (14), and (ii) plant-wax fatty acids are not detected in 142 some saprolite samples (Table S6). Thus, mid-E material must reflect a mixture of weathered 143 OCpetro and moderately aged OCbio. 144 We treat OC_{petro} that has been chemically altered during weathering as a unique end 145 member described by Fm = 0.0 and a value of f_{mid} , the fraction of p(0,E) contained within the 146 mid-E range, greater than the highest observed saprolite value of 0.51 (14). Fig. 3A shows that 147 all hillslope samples, with the exception of one unweathered saprolite, are adequately explained 148

by a mixture of OC_{bio} and chemically altered OC_{petro}. This end member is also observed in LiWu

150 River POC collected during typhoon floods, as evidenced by the divergence from a vertical

mixing line between OC_{petro} and OC_{bio} in **Fig. 3A**. Therefore, along with unweathered bedrock OC (19) sourced from deep incision and landsliding (11), we detect catchment-scale export of

152 chemically altered OC_{petro} from Central Range hillslopes during typhoon flood events. Because

154 calculated f_{mid} depends on our choice of mid-*E* range (here, 150 to 185 kJ mol⁻¹), it is possible

that mixing trends and end-member compositions are sensitive to changes in *E* boundary values.

156 We test this sensitivity by allowing these boundary values to vary by ± 10 kJ mol⁻¹ (14).

Although quantitative differences exist (**Fig. S5**), resulting mixing trends are qualitatively robust, indicating that the importance of chemically altered OC_{petro} is insensitive to our choice of mid-*E* boundary values.

Fatty acid molecular distributions and δ^{13} C values imply that rapidly oxidized OC_{petro} in soils is incorporated into microbial biomass, supporting laboratory-based incubation studies (20, 22). We calculate $f_{\text{microbial}}$, the fraction of total fatty acids that are microbial in origin (25, 26), as a proxy for the relative abundance of heterotrophic vs. vascular-plant biomass (14). This

approach excludes fungal contributions and is thus a minimum estimate of heterotrophic 164 biomass. Fig. 3B shows that bulk Fm is negatively correlated with $f_{\text{microbial}}$ across all soil and 165 POC samples. We do not expect this trend to be linear due to fatty acid production biases (25, 166 26). Still, this relationship clearly suggests that heterotrophic biomass is more abundant in 167 samples containing predominantly ¹⁴C-free OC. 168 Sample limitation prevented measurement of microbial fatty acid ¹⁴C activity (14), but 169 their δ^{13} C values imply that bedrock OC is used as substrate (**Table S7**) (26, 27). Bulk OC and 170 plant-wax fatty acid δ^{13} C values correlate strongly in A+E horizons ($r^2 = 0.959$; p-val < 0.001; n 171 = 7), reflecting the predominance of OC_{bio} in these samples, but are uncorrelated in C-horizons 172 173 (p-val > 0.05; n = 4) due to a lack of significant OC_{bio} contribution to saprolites (Fig. S6). Still, if OC_{bio} were the sole substrate for heterotrophs, then microbial and plant-wax fatty acid $\delta^{13}C$ 174 values should correlate strongly with a constant δ^{13} C offset (27) in all samples. This is not 175 observed in either A+E horizon (*p*-val > 0.05; n = 7) or saprolite (*p*-val > 0.05; n = 4) samples, 176 indicating that vascular plant OC cannot be the only substrate. Rather, this lack of correlation 177 requires a secondary microbial carbon source (20-22), namely bedrock OC. We conclude that 178 mid-E, ¹⁴C-free material is a product of microbial bedrock oxidation, produced either directly by 179 extracellular enzymes or indirectly after acid hydrolysis (20), and is manifest as ¹⁴C-depleted 180 living biomass (22) or as residual, chemically altered OC_{petro} (21). 181

182 Substantial bedrock OC replacement within saprolites implies that significant weathering occurs ≤ 1 m below the surface and that microbially mediated OC_{petro} oxidation can proceed at a 183 pace matching the rapid exhumation in Taiwan. We propose that exhumation and hillslope 184 erosion rates exert a first-order control on CO₂ emissions from OC_{petro} oxidation, as faster 185 erosion will increase the rate of bedrock exposure to the weathering front (8). This is further 186 supported by measurements of the dissolved rhenium flux from Taiwanese rivers, a proxy for 187 OC_{petro} oxidation, which increases with erosion rate (5). However, the relationship between 188 OC_{petro} oxidation and physical erosion rate cannot be linear. Large earthquakes and typhoons are 189 known to cause widespread bedrock landsliding (28-30) and elevated export of OC_{petro} by rivers 190 (19). Such events increase catchment-averaged erosion rates (28), but could decrease catchment-191 averaged OC_{petro} oxidation efficiency by bypassing the hillslope soil weathering window. OC_{petro} 192 remineralization in Taiwan is incomplete, as evidenced by the abundance of bedrock OC in 193 194 sediments exported by rivers (19) and deposited in nearby coastal margins (31). We predict a dampened response of OC_{petro} -derived CO_2 emissions to further erosion rate increases, as 195 increasing landslide rates will result in less catchment area that is available for soil formation and 196 weathering. 197

Microbially mediated oxidative weathering in Taiwanese hillslope soils offsets geologic 198 199 CO_2 drawdown and O_2 production by silicate weathering and OC_{bio} burial (1, 5, 8, 22). The Φ_{ox} range calculated here is similar in magnitude to CO_2 source estimates from sulfide oxidation (\geq 200 22.9 ± 1.0 t C km⁻² yr⁻¹; LiWu basin only) (9), as well as CO₂ sinks from silicate weathering (3.1 201 \pm 0.1 t C km⁻² yr⁻¹; LiWu basin only; Fig. S3C) (18) and OC_{bio} burial (21 \pm 10 t C km⁻² yr⁻¹; 202 Taiwan average; Fig. S3D) (14, 32). This process is likely globally significant, as rapid soil 203 formation is observed in other tropical and temperate orogenic settings such as the Southern Alps 204 of New Zealand (33). We therefore hypothesize that CO₂ consumption is not favored within 205 highly erosive mountain belts dominated by OC- and sulfide-rich low- and intermediate-grade 206 207 metasedimentary lithologies. This results from the observation that OCpetro and sulfide mineral

- 208 oxidation is not limited by reaction kinetics even at high erosion rates (5, 8, 21), unlike silicate
- 209 weathering and OC_{bio} export (4, 34). Conversely, the magnitude of the net CO_2 sink likely
- 210 increases with physical erosion rate in orogens dominated by high-grade metamorphic and
- 211 igneous rocks due to their lower OC_{petro} and sulfide contents. While the global fluxes and the
- timescales over which they impact atmospheric CO₂ and O₂ concentrations remain to be assessed, our results demonstrate the importance of microbially mediated OC_{petro} oxidation and
- assessed, our results demonstrate the importance of microbially mediated OC_{petro} oxidation and its relationship to tectonic and erosive controls on the global carbon cycle and Earth's long-term
- 215 climate.

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Fig. 1. Evidence for bedrock OC oxidation. Blue line is the solution to **Eq. 2** assuming no OC_{petro} oxidation during soil formation (OC_{bio} addition only; $f_{ox} = 0$). Black line is the orthogonal distance regression best-fit solution that minimizes the residual error between measured (green circles, orange triangles) and predicted Fm_{soil} values. Shaded region for both trends is the propagated $\pm 1\sigma$ uncertainty (14). Best-fit results indicate that 67 ± 11 % of bedrock OC is lost during oxidative weathering. Δ %OC = 0 is shown as a vertical dashed line. Measurement error bars are smaller than marker sizes.



Fig. 2. Evidence for OC_{petro} chemical alteration. (A) Representative p(0,E) distributions 405 highlighting the differences between OC end members: average of LiWu POC exported during 406 typhoon events (n = 27; black), organic-rich A+E horizon topsoil (green), and C horizon 407 saprolite (orange). Each p(0,E) distribution integrates to unity (y-axis values not shown) (14, 23). 408 (B) E vs. Fm relationships for all soils (green circles, orange triangles) and LiWu POC (white 409 diamonds) in which RPO fraction ¹⁴C activity was measured. Marker sizes represent the relative 410 amount of total OC contained in each RPO fraction. Constraints on end-member E and Fm 411 ranges are described in the main text (blue, vascular-plant OC_{bio}; gray, OC_{petro}). Black arrows 412 413 represent theoretical trends for end-member mixing (vertical) and chemical alteration (horizontal) (23) and indicate that alteration is necessary in order to explain the presence of mid-414 E OC. For both panels, dashed lines separate OC into low-E (<150 kJ mol⁻¹), mid-E (150 $\leq E \leq$ 415 185 kJ mol⁻¹), and high-*E* (\geq 185 kJ mol⁻¹) regions. Fm error bars are smaller than marker sizes. 416 417



Fig. 3. Evidence for microbially mediated bedrock OC oxidation. (A) Bulk Fm vs. fmid 419 relationships for soil (green circles, orange triangles) and LiWu POC (white diamonds). All soils, 420 with the exception of the 0.5 m saprolite discussed in the main text, are described by a mixing 421 422 line between vascular-plant OCbio (blue) and chemically altered OCpetro (red) (14). LiWu River POC is dominated by bedrock OC (gray) but does contain detectable chemically altered OC_{petro}, 423 424 as evidenced by the deviation from a vertical mixing line between OC_{bio} and OC_{petro}. (B) Bulk Fm vs. fmicrobial relationships for all samples in which fatty acid concentrations were analyzed 425 (14). The relative abundance of microbial fatty acids increases with decreasing Fm across all 426 samples, suggesting that microbial respiration is the source of chemically altered OC_{petro}. 427 Measurement error bars are smaller than marker sizes. 428