Increased carbon capture by a silicate-treated forested watershed affected by acid deposition

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Abstract. Meeting internationally agreed-upon climate targets requires Carbon Dioxide Removal (CDR) strategies coupled with an urgent phase-down of fossil fuel emissions. However, the efficacy and wider impacts of CDR are poorly understood. Enhanced rock weathering (ERW) is a land-based CDR strategy requiring large-scale field trials. Here we show that a low 3.44 t ha⁻¹ wollastonite treatment in an 11.8-ha acid-rain-impacted forested watershed in New Hampshire, USA led to cumulative carbon capture by carbonic acid weathering of 0.025–0.13 t CO₂ ha⁻¹ over 15 years. Despite a 0.8–2.4 t CO₂ ha⁻¹ logistical carbon penalty from mining, grinding, transportation and spreading, by 2015 weathering together with increased

19 forest productivity led to net CDR of 8.5-11.5 t CO₂ ha⁻¹. Our results demonstrate that ERW may be an effective, scalable

20 CDR strategy for acid-impacted forests but at large-scale requires sustainable sources of silicate rock dust.

21 1 Introduction

22 The Intergovernmental Panel on Climate Change (IPCC)(Rogelj et al., 2018) Special Report on global warming indicates 23 large-scale deployment of Carbon Dioxide Removal (CDR) technologies will be required to avoid warming in excess of 1.5 24 °C by the end of this century. Land-based CDR strategies include enhanced rock weathering (ERW), which aims to accelerate 25 the natural geological process of carbon sequestration by amending soils with crushed reactive calcium (Ca) and magnesium 26 (Mg)-bearing rocks such as basalt (The Royal Society and The Royal Academy of Engineering, 2018;Hartmann et al., 2013). 27 Forests represent potential large-scale deployment opportunities where rock amendments may provide a range of benefits, 28 including amelioration of soil acidification and provisioning of inorganic plant-nutrients to cation-depleted soils (Hartmann et 29 al., 2013;Beerling et al., 2018). Although ERW has not yet been demonstrated as a CDR technique at the catchment scale, a 30 forested watershed experiment at the Hubbard Brook Experimental Forest (HBEF, 43° 56'N, 71° 45'W) in the White

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Mountains of New Hampshire, USA provides an unusual opportunity for assessing proof-of-concept in this priority research
 area.

33 The HBEF watershed experiment, designed to restore soil calcium following decades of leaching by acid rain, involved 34 application of a finely ground rapidly-weathered calcium silicate mineral wollastonite (CaSiO₃; 3.44 t ha⁻¹) on 19 October 35 1999 to an 11.8-ha forested watershed (SI Appendix) (Likens et al., 2004;Peters et al., 2004;Shao et al., 2016). Unlike the 36 carbonate minerals (e.g., CaCO₃) commonly applied to acidified soils (Lundström et al., 2003), wollastonite does not release 37 CO_2 when weathered (Supplementary Information) so is much better suited for CDR (Hartmann et al., 2013). It also has 38 dissolution kinetics comparable to or faster than other calcium-rich silicate minerals such as labradorite found in basalt 39 (Brantley et al., 2008). Thus, the HBEF experiment provides a timely and unparalleled opportunity for investigating the long-40 term (15 years) effects of ERW on CDR potential via forest and stream water chemistry responses.

41 In the case of ERW with wollastonite, CDR follows as Ca cations (Ca²⁺) liberated by weathering consume atmospheric 42 CO₂ through the formation of bicarbonate (HCO₃⁻) by charge balance, as described by the following reaction:

43
$$CaSiO_3 + 3H_2O + 2CO_2 \rightarrow Ca^{2+} + H_4SiO_4 + 2HCO_3^{-},$$
 (1)

However, forests in the northeastern USA have experienced acid deposition (Likens and Bailey, 2014), changes in nitrogen cycling (Goodale and Aber, 2001;McLauchlan et al., 2007) and increases in dissolved organic carbon (DOC) fluxes (Cawley et al., 2014) that may affect CO₂ removal efficiency by ERW processes. In particular, CO₂ consumption as measured by bicarbonate production may be diminished if sulphate (SO₄²⁻), nitrate (NO₃⁻), or naturally-occurring organic acid anions (Fakhraei and Driscoll, 2015) (H₂A⁻) in DOC intervene to inhibit the following mineral weathering reactions. For example:

49
$$CaSiO_3 + H_2O + H_2SO_4 \rightarrow Ca^{2+} + H_4SiO_4 + SO_4^{2-},$$
 (2)

$$50 \quad \text{CaSiO}_3 + \text{H}_2\text{O} + 2\text{HNO}_3 \rightarrow \text{Ca}^{2+} + \text{H}_4\text{SiO}_4 + 2\text{NO}_3^{-}, \tag{3}$$

51
$$CaSiO_3 + H_2O + 2H_3A \rightarrow Ca^{2+} + H_4SiO_4 + 2H_2A^-,$$
 (4)

These environmental effects on stream-water chemistry are well documented at the HBEF (Cawley et al., 2014;Likens and Bailey, 2014;Rosi-Marshall et al., 2016;McLauchlan et al., 2007), and may be exacerbated under future climate change (Sebestyen et al., 2009;Campbell et al., 2009).

Here we exploit the experimental design and long-term monitoring of streamwater chemistry, trees, and soils, for two small forested HBEF watersheds to evaluate the effects of the wollastonite treatment in 1999 on catchment CO_2 consumption via inorganic and organic pathways. Further, we examine how biogeochemical perturbations in S, N, and organic carbon cycling affect catchment inorganic CO_2 consumption. We consider the forest response, the carbon cost for ERW deployment (mining, grinding, transportation and application), and the net greenhouse gas balance for the treatment. Finally, we provide an initial assessment of the net CDR potential of silicate treatments deployed over larger areas of acidified forest in the northeastern United States. 62 2 Methods

63 2.1 Site and treatment

64 2.1.1 Site description





Figure 1: Location of the sampling sites and experimental watersheds. Our streamwater samples were collected just upstream of the weirs in the treated and reference watersheds (gold disks) and our trace gas samples were collected at different elevations in treated and untreated forests (red disks).

69

70

- 71 The HBEF has a temperate climate with ~1400 mm mean annual precipitation of which up to one third falls as snow (Campbell
- et al., 2007). The mean temperatures in January and July are -9 °C and 18 °C respectively, and the period from mid-May to mid-September comprises the growing season (Campbell et al., 2007). There are six small southeast-facing watersheds in the

74 HBEF (Fig. 1) with 20%–30% slopes (Groffman et al., 2006), including one which received the silicate treatment (watershed 75 W1, 11.8 ha, 488–747m asl) and a biogeochemical reference (watershed W6, 13.2 ha, 545–791m asl). Carbonate and evaporite 76 minerals are in very low abundance (<1% calcite in the crystalline rocks and glacial deposits) in these silicate-mineral 77 dominated watersheds (Johnson et al., 1981). Well-drained Typic Haplorthod soils with pH<4.5 and mean depth 0.6m formed 78 from relatively impermeable glacial till, which restricts water flow and protects the underlying schist bedrock from weathering. 79 Overland runoff and flow through bedrock are both thought to be negligible (Likens, 2013). Hydrologically, the HBEF 80 watersheds are typical of small catchments in northern New England (Sopper and Lull, 1965). Flow rates for W1 and W6 81 along with streamwater pH are shown in Fig. S1. Prior to treatment, streamwater calcium concentrations were under 30 µmol L^{-1} while bicarbonate concentrations were under 5 μ mol L^{-1} , below the ranges for typical world rivers (Moon et al., 2014) (60– 82 2293 μmol Ca²⁺ L⁻¹, 179–4926 μmol HCO₃⁻ L⁻¹). 83

Fagus grandifolia, Betula allegheniensis and *Acer saccharum* are the dominant trees in this Northern Hardwood forest,
while *Betula papyrifera, Abies balsamea* and *Picea rubens* are common at the highest elevations where soils tend to be shallow
and wetter (Cho et al., 2012). *A. saccharum* and *P. rubens* are both calcium-sensitive, but soil calcium-bearing minerals are
less available to *A. saccharum* (Blum et al., 2002) and total bioavailable calcium content decreases with elevation (Cho et al.,
2012). This silicate-addition experiment was designed to replace bioavailable calcium which had been stripped from the soils
by decades of acid deposition.

90 2.1.2 Treatment description

On 19 and 21 October 1999, W1 was treated with 344 g/m² of pelletized wollastonite (CaSiO₃) by a GPS-equipped helicopter 91 92 with a motorized spreader to ensure even deployment across the catchment, including the 1804 m² streambed (Peters et al., 93 2004). Following treatment, the lignin-sulfonate binder forming the pellets dissolved within several days (Peters et al., 2004), 94 and the ground wollastonite itself dissolved rapidly in the upper Oie soil horizon, increasing Oie base saturation from 40% to 95 78% and raising soil pH from 3.88 to 4.39 within one year (Johnson et al., 2014). Although the budget of wollastonite-derived 96 calcium (Wo-Ca) has never been closed due to lack of data from vegetation and from deeper soil layers (Shao et al., 2016), it 97 is thought that uptake by vegetation and retention by soil exchange sites delayed transport of Wo-Ca to lower soil horizons 98 and streamwater for three years (Johnson et al., 2014).

99 2.2 Modelling approach

100 2.2.1 Forward modelling of streamwater chemistry including dissolved inorganic carbon

101 We used a forward modelling approach to calculate dissolved streamwater bicarbonate concentrations ([HCO₃⁻]_{stream}) in the

treated and reference watersheds (**Fig. 1**) over ~25 years, including 15 years post-treatment, with the United States Geological

103 Survey (USGS) aqueous geochemistry software PHREEQC version 3.3.12-12704 (Parkhurst and Appelo, 1999) and monthly

104 long-term (1992–2014) streamwater (Driscoll, 2016b, a) and rain/snow precipitation (Likens, 2016b, a) chemistry 105 measurements.

106 Using MATLAB (version R2016a) scripts, we wrote PHREEQC input files and determined the inorganic carbon species 107 for each streamwater sample with PHREEQC. Along with a standard database which decouples ammonium and nitrate 108 (Amm.dat, provided with the PHREEQC software), we included the ionization constants for the organic acid triprotic analogue 109 and the constants for Al complexation described for Hubbard Brook streams (Fakhraei and Driscoll, 2015) in our PHREEQC simulations. These are: $pK_{a1}=2.02$, $pK_{a2}=6.63$, $pK_{a3}=7.30$, $pK_{A11}=4.07$, $pK_{A12}=7.37$, $pK_{A13}=6.65$, and site density m=0.064 mol 110 111 sites mol C⁻¹. Our organic acid concentrations are the product of the corresponding site density of reactions and the measured 112 dissolved organic carbon concentration (Fakhraei and Driscoll, 2015); these were PHREEQC inputs along with total 113 monomeric Al and major ion concentrations from the longitudinal datasets.

Spectator ions (Cl^- and NH_4^+) were adjusted to achieve charge balance given the measured pH for the treated and reference watersheds. Cl^- was only adjusted when charge balance was not achieved using NH_4^+ alone. This was deemed to be the case when PHREEQC failed to converge or when the percent error exceeded 5%. We used original rather than adjusted rainwater Cl to calculate the contribution of rainwater to streamwater chemistry (described below). These adjusted ions were then held constant for our modelled scenarios, while pH was allowed to vary.

Exploratory PHREEQC tests (charge-balancing on DIC) either with or without organic acids suggest that the acids depress total DIC, HCO_3^- and also the saturation state of gaseous CO₂. Similar variability in the saturation is also observed when DIC values from partially degassed samples from the streams are used as input. We chose minimum and maximum values of 1100 and 1700 ppm, or ~3 and 4.6 × 368, the mean value of Mauna Loa pCO₂ (Tans and Keeling, 2017) for 1985–2012. These values correspond to $log_{10}(pCO_2(g)) = -2.87\pm0.09$ SD derived from a prior analysis of this variability for the same time range (Fakhraei and Driscoll, 2015).

125 2.2.2 Streamwater temperature

Air temperatures for the Hubbard Brook watersheds (Campbell, 2016) were converted to streamwater temperatures (Mohseni and Stefan, 1999). Rainwater temperatures were set equal to streamwater temperatures. These temperatures were used in our PHREEQC modelling, with equilibrium constants for the DIC species as functions of temperature. Only samples measured closest to the weirs and with a valid pH were processed with PHREEQC.

130 2.2.3 Catchment CO₂ consumption

We calculate total annual watershed CO₂ consumption (Eq. 1) as the product of streamwater flow and streamwater bicarbonate concentration [**HCO₃-**] at time *t* corrected for the HCO₃⁻ contribution of rainwater ($\alpha_{rain,HCO3}(t)$, see below):

133
$$CO_{2HCO3}(t) = (1 - \alpha_{rain,HCO3}(t)) [HCO3^{-}](t) \times flow(t),$$

(5)

where $[HCO_3-](t)$ is given in mol kgw⁻¹ and *flow*(t) is the "runoff" in mm year⁻¹. Calculated $[HCO_3-]$ and annual CO₂ consumption for the treated and reference watersheds (Eq. 5) comprise our baseline simulations and represent a primary test of hypothesized increased carbon capture resulting from weathering of the applied silicate.

To isolate a treatment effect for bicarbonate, we used strontium isotopes as a tracer of wollastonite (Wo) weathering within a previously-published mixing function (Nezat et al., 2010;Peters et al., 2004) (Methods, **Fig. S3**). This mixing function provides the fraction *X* of calcium originating from wollastonite. The contribution of all mineral sources other than wollastonite to CO₂ consumption (Eq. 5) is simulated by running simulations with Ca²⁺ concentrations reduced by (1-X):

141

142 Non-Wo-CO_{2,HCO3} (
$$t$$
) = [HCO₃-](t ,(1- X) [Ca²⁺]) × (1- $\alpha_{rain,HCO3}(t)$) × flow(t), (6)

143

where $\alpha_{rain,HCOS}(t)$ is the fractional contribution from rain/snow precipitation (Methods). Eq. 6 is required only to calculate the treatment effect from bicarbonate concentrations, which is the difference between Eq. (5) and Eq. (6):

146

147 *Wo-CO*2,нсоз (*t*)

148 = ([HCO₃-](
$$t$$
, [Ca²⁺]) - [HCO₃-](t , (1- X) [Ca²⁺])) × (1- $a_{rain,HCO3}(t)$) × flow(t)
149 = CO_{2,HCO3}(t) - Non-Wo-CO_{2,HCO3}(t), (7)

150

Bicarbonate-derived CO₂ consumption (Eq. 5) is the most conservative approach to estimating net carbon fluxes related to ERW. For natural freshwaters in equilibrium with the atmosphere, this entails a titration for total alkalinity with a possible correction for the concentration of organic acid anions (Köhler et al., 2000). However, another widely used (Jacobson and Blum, 2003) measure of CO₂ consumption is derived assuming charge-balance of base cations (Ca²⁺, Mg²⁺, K⁺ and Na⁺) by bicarbonate formation (Eq. 1)

156
$$CO_{2,ions}(t) = (2[Ca^{2+}](1-\alpha_{rain,Ca}(t)) + 2[Mg^{2+}](1-\alpha_{rain,Mg}(t)) + [K^+](1-\alpha_{rain,K}(t)) +$$

157 $[Na^+](1-\alpha_{rain,Na}(t)) - 2[SO_4^{2-}](t)) \times flow(t),$ (8)

where streamwater cation equivalents are corrected for contributions from rain/snow precipitation (Methods) and sulphuric acid weathering (Chetelat et al., 2008) (Eq. 2). Eq. 8 is an optimistic measures of CO_2 consumption because it ignores both the weathering agent and streamwater inorganic carbon, assuming charge-balance of cations by carbonate and bicarbonate ions in the oceans. We will present *CO*_{2,ions} results for comparison with *CO*_{2,HCO3}, but Eq. 8 also allows us to derive a more optimistic estimate of the treatment effect than *Wo-CO*_{2,HCO3} (Eq. 7). For an ERW treatment, transient changes in the export of ions not derived from the applied minerals may occur, but we consider that the cations released from the applied minerals comprise the most unambiguous treatment effect in our study. The charge associated with wollastonite-derived Ca^{2+} (Wo-Ca) determines the CO₂ consumption associated with the HBEF wollastonite treatment, and from Eq. 8 our optimistic treatment effect based on calcium rather than bicarbonate reduces to:

167
$$Wo-CO_{2,Ca}(t) = 2 \times X \times [Ca^{2+}](t) \times flow(t),$$
(9)

168

where X is the fraction of Ca²⁺ from weathered wollastonite. Eqs. 7 and 9, together with our flux calculations accounting for sparsity of concentration data compared to daily flow data (Methods), should help avoid major uncertainties in catchmentscale CO₂ consumption calculations: the provenance of the cations and variations in concentration and discharge (Moon et al., 2014).

173 **2.2.4 Fraction of calcium derived from wollastonite**

174 We applied an existing two-component mixing model (Peters et al., 2004):

$$175 \quad \mathbf{X} \quad (\mathbf{t}) = \left[\frac{\left(\left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{post}} - \left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{pre}} \right) \left(\frac{\mathbf{Sr}}{\mathbf{Ca}}\right)_{\mathbf{pre}}}{\left(\left(\left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{post}} - \left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{pre}} \right) \left(\frac{\mathbf{Sr}}{\mathbf{6a}}\right)_{\mathbf{pre}} + \left(\left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{pre}} - \left(\frac{\mathbf{87}_{Sr}}{\mathbf{86}_{Sr}}\right)_{\mathbf{pre}} \right) \left(\frac{\mathbf{Sr}}{\mathbf{6a}}\right)_{\mathbf{pre}} \right) \left(\frac{\mathbf{Sr}}{\mathbf{6a}}\right)_{\mathbf{pre}} \right], \tag{10}$$

where pre-app and post-app refer to pre-application and post-application streamwater concentrations and Wo refers to wollastonite. The Sr data (Blum, 2019) have been extended through 2015 (**Fig. S1a**). See SI Appendix for further discussion of the use of strontium and its isotopes as tracers of Ca^{2+} provenance.

179 2.2.5 Contributions of rain/snow precipitation to streamwater chemistry

We estimated the contribution of rain/snow (Likens, 2016b, a) relative to all other sources using a previously published mixing model (Négrel et al., 1993). We assume all Cl⁻ in the water is from rain/snow, noting that this common treatment of Cl as an unreactive tracer is not always justified (Lovett et al., 2005). We calculate the contribution of precipitation to the streamwater (α_{rain}) using Na and Cl, which are less affected by nutrient cycling and adsorption than other major ions (Négrel et al., 1993):

184
$$\boldsymbol{\alpha}_{\operatorname{rain,Na}}(t) = \frac{\left[\frac{\mathrm{Cl}}{\mathrm{Na}}\right](\operatorname{stream},t)}{\left[\frac{\mathrm{Cl}}{\mathrm{Na}}\right](\operatorname{rain},t)},$$
(11)

185

- - -

To account for attenuation of the rain/snow precipitation leaching through the soil, Cl/Na and HCO₃/Na at any given time (t) are means from the previous three months. We estimate the contribution of rain/snow to other ions such as HCO₃⁻ in the streamwater as follows:

189
$$\alpha_{\text{rain,HCO3}}(t) = \alpha_{\text{rain,Na}}(t) \times \frac{\left[\frac{\text{HCO3}}{\text{Na}}\right](\text{stream,t})}{\left[\frac{\text{HCO3}}{\text{Na}}\right](\text{rain,t})}$$

190

191 2.2.6 Flux calculations

192 To ensure that fluxes from our two watersheds were comparable and to correct for the sparsity of solute measurements 193 compared to flow measurements, we created rolling annual flow-adjusted fluxes using Method 5 of Littlewood et al. (1998) at 194 five evenly-spaced points each year:

195
$$\mathbf{Flux} = \mathbf{scale} \times \left[\frac{\sum_{i=1}^{M} C_i Q_i}{\sum_{i=1}^{M} Q_i} \right] \times \left[\frac{\sum_{k=1}^{N} Q_k}{N} \right], \tag{13}$$

196

where Q_i is the measured instantaneous stream flow, C_i is the concentration for sample *i*, *M* is the number of streamwater chemistry samples in the year (usually 12), Q_k is the k^{th} flow measurement, and *N* is the number of flow measurements. In our case, daily flow measurements (Campbell, 2015) and ~monthly streamwater samples (Driscoll, 2016b, a) were available. Therefore, the mean concentration for the preceding twelve months is multiplied by the mean flow for the same period, suitably scaled to get the total annual flux. Without sub-daily timestamps for the longitudinal streamwater chemistry data, we used daily total flows rather than instantaneous flows. Tests suggested that there was little difference between using mean daily instantaneous flows and the mean daily total flows.

Variable	Units	Sections	Eq.	Fig.	Table	Description
[HCO3-]	mol kgw ⁻¹	2.2.1	5	2a, S5a,d,g S6a		Bicarbonate concentration in streamwater
<i>A</i> rain,HCO3	fraction	2.2.3, 2.2.5	5,12	S 3		Fraction of bicarbonate from rainwater
flow	mm year-1	2.2.3	5–9	S2d		Streamwater flow
<i>СО</i> 2,нсоз	mol C year ⁻¹	2.2.3	5	2b, S5b,e,h	3,S1	Total watershed CO ₂ consumption as calculated from bicarbonate
<i>Non-Wo-</i> <i>CO</i> 2,нсоз	mol C year-1	2.2.3	6			Watershed CO ₂ consumption (calculated from bicarbonate) not due to wollastonite weathering
<i>Wo-CO</i> 2,нсоз	mol C time ⁻¹	2.2.3	7	2c,5, S5c,f,i S6b,c	3,4,S1	Watershed CO ₂ consumption (calculated from bicarbonate) due to wollastonite weathering. Our conservative/pessimistic $\Delta CONS$ estimate in our GHG balance is the 15-year sum.
X	fraction	2.2.3	6,7,9,10	S1a		Fraction of total calcium originating from wollastonite
t	time	2.2.3	5–7			Time
CO _{2,ions}	mol C time ⁻¹	2.2.3	8		3	Total watershed CO ₂ consumption as calculated from major ions

204 Table 1. Summary of variables presented in Methods (Section 2, Eqs 5 through 14)

<i>A</i> rain,Ca	fraction	2.2.3	8	S4		Fraction of calcium from rainwater	
<i>A</i> rain,Mg	fraction	2.2.3	8			Fraction of magnesium from rainwater	
<i>A</i> rain,K	fraction	2.2.3	8			Fraction of potassium from rainwater	
<i>A</i> rain,Na	fraction	2.2.3, 2.2.5	8,11	S4		Fraction of sodium from rainwater	
[Ca ²⁺]	mol kgw ⁻¹	2.2.3	6–9	2d		Calcium concentration in streamwater	
[Mg ²⁺]	mol kgw ⁻¹	2.2.3	8			Magnesium concentration in streamwater	
[K+]	mol kgw ⁻¹	2.2.3	8			Potassium concentration in streamwater	
[Na+]	mol kgw ⁻¹	2.2.3	8			Sodium concentration in streamwater	
[SO4 ²⁻]	mol kgw ⁻¹	2.2.3	8	S5a		Sulphate concentration in streamwater	
Wo-CO _{2,Ca}	mol C year-1	2.2.3	9	2f, S6b,c	3, S1	Watershed CO ₂ consumption (calculated from calcium) due to wollastonite weathering. Our optimistic estimate for $\Delta CONS$ in our GHG balance is the 15-year sum.	
⁸⁷ Sr/ ⁸⁶ Sr	fraction	2.2.4	10	S1b,c		Strontium isotope ratio	
Sr/Ca	fraction	2.2.4	10	S1b,c		Strontium to calcium ratio	
Ci	mol kgw ⁻¹	2.2.6	13			Concentration of solute for sample <i>i</i> (collected ~monthly for chemical analysis)	
Q_i	mm time ⁻¹	2.2.6	13			Streamflow for sample <i>i</i>	
Qk	mm day-1	2.2.6	13	S2b		Streamflow for day <i>k</i>	
N	number	2.2.6	13			Number of daily flow measurements	
ΔGHG	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Net treatment effect on watershed greenhouse gas balance	
∆wood	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on woody biomass over ten years, positive if wood production increases relative to reference watershed.	
ΔСН4	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on soil CH_4 sink since 2002, positive if the soil CH_4 sink increases relative to reference watershed.	
ΔSRESP	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on soil CO ₂ emissions since 2002, positive if emissions decrease relative to reference watershed.	
ΔCONS	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on CO ₂ consumption over 15 years, range from <i>Wo-CO</i> _{2,HCO3} and <i>Wo-CO</i> _{2,Ca}	
ΔΝ2Ο	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on soil N ₂ O emissions since 2002, positive if emissions decrease relative to reference watershed.	
ΔΝΟ3Ν2Ο	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on downstream N_2O emissions (due to nitrate export) over 15 years, positive if emissions decrease relative to reference watershed.	
ΔDOC	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Treatment effect on dissolved organic carbon export over 15 years, positive if export decreases relative to reference watershed. This	

						represents carbon loss from the watershed and likely CO ₂ emissions downstream.	
LOGPEN	t CO ₂ ha ⁻¹	2.3.1	14	5	4	Logistical emissions penalty associated w mining, milling, pelletization, transport a application of the wollastonite treatme expected to be negative.	
S	$m^2 kg^{-1}$	2.3.3	15			Specific surface area of material being milled	
<i>Ө</i> р	kJ kg ⁻¹	2.3.3	15			Specific potential energy of material being milled	

205

206 2.3 Greenhouse gas balance

207 2.3.1 Greenhouse gas budget for a treatment

The success of any treatment for climate change mitigation is determined by the net greenhouse gas (CO_2 equivalent) fluxes prior to and following treatment, at the treatment site and downstream. Desireable outcomes for a treatment include increased ecosystem carbon storage in biomass and soils, increased CO_2 consumption, and decreases in ecosystem, downstream and logistical greenhouse gas emissions.

At the HBEF, we have measured the CO_2 consumption due to the wollastonite treatment in two different ways and these determine our range of values to be incorporated in our GHG budget. Several other treatment effects can be estimated relative to the reference watershed, but some aspects of the total GHG balance are missing. For example, we have measurements of soil respiration (root+heterotrophic) and dissolved organic carbon (DOC) export in streamwater, but we lack measurements of canopy respiration from leaves and stems, and export of particulate organic carbon in streamwater. Our partial greenhouse gas budget for the HBEF wollastonite treatment will therefore be given by

218

$219 \quad \Delta GHG = \Delta wood + \Delta SRESP + \Delta CH4 + \Delta N20 + \Delta CONS + \Delta NO3N20 + \Delta DOC + LOGPEN, \tag{14}$

220

where our partial GHG treatment effect (ΔGHG) is the sum of greenhouse gas sink and source responses. Measured sinks for the wollastonite experiment include biomass in wood ($\Delta wood$), CO₂ consumption ($\Delta CONS$), and a soil sink for methane ($\Delta CH4$). Sources include N₂O emissions both from soil ($\Delta N2O$) and exported nitrate ($\Delta NO3N2O$), and CO₂ emissions from soil respiration ($\Delta SRESP$), exported dissolved organic carbon (ΔDOC), and logistical operations (LOGPEN).

Sink effects are defined as positive if the sink increases and are given by the difference (treated-reference) between the two watersheds, whereas source effects are defined as positive for reductions in greenhouse gas emissions (reference-treated). With these definitions, penalties are negative and reduce ΔGHG in Eq. 14. Logistical emissions and CO₂ consumption due to weathering of applied wollastonite are zero for the reference watershed, so we expect *LOGPEN* to be negative and $\Delta CONS$ to be positive. 230 Wood is a longer-term carbon sink than leaves or twigs so we have chosen to let this represent our biomass increment.

231 Eq. (14) neglects ecosystem disturbances including fire, and possible carbonate mineral precipitation in soils. There is no

232 evidence for the latter at the HBEF.

We used a range of emissions factors for N₂O to estimate the penalty associated with nitrate export ($\Delta NO3N2O$); low: 0.0017 kgN₂O-N kg⁻¹ DIN (Hu et al., 2016) and high: 0.0075 kgN₂O-N kg⁻¹ DIN (De Klein et al., 2006), where DIN is dissolved inorganic nitrogen dominated by nitrate. This N₂O was then converted to CO_{2e} (CO₂ equivalents in terms of cumulative radiative forcing) given the 100-year time horizon global warming potential (Pachauri et al., 2014) (GWP₁₀₀) for N₂O: 265 gCO_{2e} g⁻¹ N₂O. Likewise, $\Delta CH4$ was converted to CO_{2e} (CO₂ equivalents in terms of cumulative forcing) given GWP₁₀₀ for CH₄: 28 gCO_{2e} g⁻¹ CH₄.

239

240 2.3.2 Carbon sequestration in wood

We calculate our treatment effect on wood production as the difference between the treated and reference watershed mean wood production (Battles et al., 2014) over two five-year periods. We considered these differences (treated-reference) to be an estimate of the treatment effect on potentially long-term (decades to centuries) biomass carbon sequestration. Assuming 46.5% of the woody biomass is carbon (Martin et al., 2018), our calculated cumulative additional C sequestration in the treated watershed over ten years was 20.7 mol C m⁻² (9.1 t CO₂ ha⁻¹). Our optimistic and pessimistic values are derived from the 95% confidence intervals for the five-year mean values (Battles et al., 2014).

247 2.3.3 Greenhouse gas emissions from soils

Measurements (Groffman, 2016) were taken at four elevations in the treated watershed and at points just west of the reference watershed starting in 2002 (**Fig. 1**). Gas samples were collected from chambers placed on three permanent PVC rings at each of these eight sites (Groffman, 2016). The data were not normally distributed so were analyzed with Kruskal-Wallis tests at the 0.05 significance level; however, tests with one-way ANOVA produced the same overall results. All analyses were done in Matlab R2016a.

Cumulative curves for each of the 24 chambers were generated by matching the dates of the measurements, excluding points which were missing data for any chamber and allowing up to a week's discrepancy between catchments. Nearly all discrepancies were within one day. Assuming diurnal variation was minor compared to seasonal variation, each datum (g C m^{-2} hour⁻¹) was multiplied by 24 hours and by 30 days to get gC m^{-2} month⁻¹. There was no extrapolation to fill gaps in the dataset; results are internally consistent but not comparable to other datasets. We were particularly interested in the elevationspecific responses, as the different elevations have distinct tree species compositions and below-ground responses to the wollastonite treatment (Fahey et al., 2016). 260 The HBEF experimental watersheds are divided into 25×25m plots on slope-corrected grids. Vegetation has been surveyed four times since the late 1990s and assigned a zone designation in each plot (Driscoll et al., 2015;Driscoll Jr et al., 261 262 2015;Battles et al., 2015b, a) (Fig. S12). To estimate the respiration savings over the whole watershed, we added the areas of individual plots which were assigned to our four vegetation types (Low, Mid and High hardwoods, and Spruce-Fir). Because 263 264 there were seven vegetation types in the datasets, we compared all types with pairwise Kruskal-Wallis tests at the 0.05 265 significance level using the basal area data for the six dominant tree species. Kruskal-Wallis tests were appropriate because 266 the data, and therefore the differences from the means (residuals), were not normally distributed. These tests suggested that 267 the "extra" vegetation types ("Birch/Fern Glade", and "Poor Hardwoods" at High and Mid elevations) could be combined with Spruce-Fir, High and Mid Hardwoods respectively. Watershed fractions for our combined forest types were 0.155 for 268 269 SpruceFir, 0.16 for High Hardwoods, 0.415 for Mid Hardwoods, and 0.27 for Low Hardwoods. When creating our composite 270 treatment effects for the entire watershed, we considered a treatment effect to be present only where our statistical analyses 271 suggested significantly different fluxes.

272 2.3.4 Logistical carbon emissions costs

We used the 1999 upstate New York CO_2 emission factor for electricity generation from oil (United States Environmental Protection Agency, 1999) (0.9 Mg CO_2 MWh⁻¹), and rearranged Equation 28 of Stamboliadis (Stamboliadis et al., 2009):

275
$$\boldsymbol{e}_{\boldsymbol{p}} = \frac{\left[e^{\frac{(\ln S_{\alpha})}{\mu}}\right]}{3600 \times 1000},$$
 (14)

where the specific surface area s (1600 m² kg⁻¹ for our treatment) is related to the specific potential energy e_p of the material (kJ kg⁻¹), with theoretical parameters (Stamboliadis et al., 2009) α =139 m² kJ⁻¹ and μ =0.469 (dimensionless). We convert this potential energy to MWh t⁻¹ Qz (3600 seconds per hour and 1000 kWh MWh⁻¹). The equation was derived for quartz (Qz) which has hardness 7. Because wollastonite hardness is in the range 5–5.5, this equation may overestimate the energy needed to grind the wollastonite.

The main energy source in Allerton will have been coal, and the 1999 Illinois emissions factor (United States Environmental Protection Agency, 1999) is $1.1 \text{ Mg CO}_2 \text{ MWh}^{-1}$. The monetary cost is USD0.041 kWh⁻¹ for pelletization of limestone fines and USD0.85 t⁻¹ product, so we estimate 20.73 kWh t⁻¹ product.

Road transport distances were estimated using Google Maps (1397 km Gouverneur to Allerton, 1757 km Allerton to Woodstock, 408 km Gouverneur to Woodstock). We used standard emissions ranges (Sims et al., 2014) for Heavy Duty Vehicles (HDVs) (70–190 gCO₂ km⁻¹ t rock⁻¹) and for short-haul cargo aircraft (1200–2900 gCO₂ km⁻¹ t⁻¹). Calculation details are given in Table 2. The Matlab script used for these calculations is available on request. Note: t refers to megagrams, not US short tons.

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- 290

291 Table 2. Logistical penalty calculations for the Hubbard Brook wollastonite treatment

Penalty element	Value and calculation with units
Mass of wollastonite (CaSiO ₃) shipped to Allerton (t ^a Wo)	109665 lbs or 49.7432073 t Wo
Mass of pellets shipped from Allerton (t pellets)	112992 lbs or 51.2523091 t pellets
Ratio of pellet mass to Wollastonite mass	1.0368 = 51.25 t pellets / 49.74 t Wo
HDV transport distance (km)	3154 km = 1397 km (Gouverneur to Allerton) + 1757 km
	(Allerton to Woodstock)
Transport distance for "local pelletization" calculation (km)	408 km (Gouverneur to Woodstock)
Optimistic transport emissions (g CO ₂ g ⁻¹ Wo applied)	0.229 g CO₂ g⁻¹ Wo applied = 70 gCO ₂ km ⁻¹ shipped t ⁻¹
	shipped × ((1397 km ×49.74 t Wo shipped)+(1757 km ×
	51.25 t pellets shipped)) / 48.86×10^6 g Wo applied
Pessimistic transport emissions (g CO_2 g ⁻¹ Wo applied)	0.620 g CO₂ g⁻¹ Wo applied = 190 gCO ₂ km ⁻¹ shipped t ⁻¹
	shipped × ((1397 km ×49.74 t Wo shipped)+(1757 km ×
	51.25 t pellets shipped)) / 48.86×10^6 g Wo applied
Mass of pellets deployed by helicopter (t pellets applied)	110992 lbs or 50.3451243 t pellets applied
Mass of wollastonite deployed by helicopter (t Wo applied)	48.86 t Wo applied = 50.345 t pellets applied / 1.03684
Total area treated (ha)	14.2 ha = 11.8 ha watershed plus 2.4 ha "destructive area"
	along the western edge
Nominal mean round trip flight distance (km, Woodstock	5 km
to watershed and back)	
Number of flights (1 short ton hopper capacity) ^b	55.5 = 50.345 t pellets / 0.907 t per trip
Molar mass of wollastonite CaSiO ₃ (g Wo mol ⁻¹ Wo)	116.17 g Wo mol⁻¹ Wo = 40.08 g Ca mol ⁻¹ Ca + 28.09 g Si
	$mol^{-1}Si + 3 \times 16 \text{ g O mol}^{-1}O$
Molar mass of CO_2 (g CO_2 mol ⁻¹ CO_2)	44.01 g CO₂ mol⁻¹ CO₂ = 2×16 g O mol ⁻¹ O + 12.01 g C
	mol ⁻¹ C
Optimistic spreading emissions (mol CO ₂ ha ⁻¹)	483.36 mol CO₂ ha⁻¹ = 1200 gCO ₂ km ⁻¹ t ⁻¹ × 5 km ×
	50.345 t pellets / 44.01 g CO ₂ mol ⁻¹ CO ₂ / 14.2 ha
Optimistic spreading emissions (g CO ₂ g ⁻¹ Wo)	0.006 g CO₂ g⁻¹ Wo = 1200 gCO ₂ km ⁻¹ t ⁻¹ \times 5 km \times 50.345
	t pellets /48.86/10 ⁶ g Wo
Pessimistic spreading emissions (mol CO ₂ ha ⁻¹)	1168.1 mol CO₂ ha⁻¹ = 2900 gCO ₂ km ⁻¹ t ⁻¹ \times 5 km \times
	50.345 t pellets / 44.01 g CO ₂ mol ⁻¹ CO ₂ / 14.2 ha
Pessimistic spreading emissions (g CO ₂ g ⁻¹ Wo)	0.015 g CO₂ g⁻¹ Wo applied = 2900 gCO ₂ km ⁻¹ t ⁻¹ \times 5 km
	\times 50.345 t pellets / 48.86 \times 10 ⁶ g Wo applied

292

293 ^aMegagrams or metric tons, not short tons

²⁹⁴ ^bNumber of flights does not explicitly enter into penalty calculations because the emissions for shorthaul aircraft are multiplied

by the 5km round trip distance and the entire mass transported, rather than the mass transported during one round trip (one

short ton).

297

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299 3 Results





Figure 2: Inorganic CO₂ capture at the Hubbard Brook Experimental Forest. (a) Observed calcium and (b) calcium export in the reference (grey) and treated (blue) watersheds along with the contribution from sources other than wollastonite (red) and the time of treatment (vertical dotted line). (c) Calculated CO₂ consumption due to the treatment (*Wo-CO*_{2,Ca}, Eq. 9). (d) Modelled streamwater bicarbonate, (e) CO₂ consumption (*CO*_{2,HCO3}, Eq. 5), and (f) CO₂ consumption due to the treatment (*Wo-CO*_{2,HCO3}, Eq. 7), colours as for calcium. Simulations (d–f) account for the presence of organic acids (+OA). All calcium export (b) and CO₂ consumption curves (c,e,f) were calculated with flow-normalised concentrations and corrected for sparsity of samples (Methods).

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We first consider the time-series of streamwater changes in Ca^{2+} concentrations in the treated ([Ca]_{Treated}) and reference 309 310 ([Ca]_{Ref}) watersheds. Immediately after treatment, [Ca]_{Treated} increased from $<30 \mu$ mol L⁻¹ to $\sim60 \mu$ mol L⁻¹, and then slowly 311 declined over the next decade, remaining persistently above [Ca]_{Ref} for 15 years (Fig. 2a). The initial post-treatment peak represents dissolution of wollastonite within the stream (Peters et al., 2004) and release of calcium from hyporheic exchange 312 during the first few years (Shao et al., 2016;Nezat et al., 2010). Retention of Ca^{2+} ions liberated by wollastonite dissolution 313 314 (Wo-Ca) in the watershed soils (Nezat et al., 2010) and sequestration into tree biomass (Balogh-Brunstad et al., 2008;Nezat et 315 al., 2010) delayed appearance in streamwater for three years (Shao et al., 2016;Nezat et al., 2010). Subsequently, [Ca]_{Treated} remained approximately double [Ca]_{Ref}, with a ~30% contribution from non-wollastonite Ca²⁺ until 2012. Towards the end of 316 the time-series, increased seasonal NO₃⁻ export in the treated watershed between 2012 and 2014 (Rosi-Marshall et al., 2016) 317

318 led to Wo-Ca displacing non-Wo-Ca from the soil exchanger.

We derived the annual export of Ca^{2+} from the treated and reference watersheds as the product of mean annual flowadjusted Ca^{2+} streamwater concentrations and annual flow (**Fig. 2b**) (Methods). After accounting for variations in flow, increased streamwater Ca^{2+} concentrations in the treated watershed are translated into a 2-fold increase in total Ca^{2+} export relative to the reference watershed that was maintained for 15 years until 2015 through this analysis period. Overall, the wollastonite treatment resulted in a sharp spike in calculated CO_2 consumption (*Wo-CO*_{2,Ca}) that decreased but remained elevated as a result of the treatment (**Fig. 2c**).

Temporal patterns in modelled streamwater bicarbonate concentration in both treated and reference watersheds (**Fig.** 2d), and the corresponding total annual CO₂ consumption (*CO*_{2,HCO3}) (**Fig. 2e**) and CO₂ consumption resulting from treatment (*Wo-CO*_{2,HCO3}) (**Fig. 2f**), largely mirror changes in streamwater Ca²⁺ concentrations but are modified by the supply and loss of anions. Calculated flow-adjusted CO₂ consumption (**Fig. 2e**) peaked 2–3 years post-treatment with a broader peak in CO₂ consumption evident in 2007–2012 corresponding to declining legacy effects of acid rain until transient NO₃⁻ peaks appeared 2012–2015. *Wo-CO*_{2,HCO3} shows a pattern that mirrors *Wo-CO*_{2,Ca} but is generally 5 times lower (**Fig. 2c,f**).

331 3.2 Sulphuric, nitric and organic acids reduce CDR

332 We next undertook sensitivity analyses to investigate the effects of acid deposition, increased NO₃⁻ and organic acid export 333 from the treated watershed on bicarbonate concentrations and resulting CO_2 consumption (Fig. S5). In a 'Low SO_4 ' scenario 334 (Fig. S5a-c), we sought to understand the effects of acid deposition by replacing the mean monthly time-series of streamwater and rainwater SO_4^{2-} for the treated watershed with a new time-series (purple curve, **Fig. S5a**) created by repeating the post-335 336 2010 datasets, which reflect diminished acid deposition following emission controls from the US Clean Air Act (Likens and 337 Bailey, 2014). Removing acid rain effects in this manner dramatically increased the calculated bicarbonate concentrations and 338 total annual CO_2 consumption (**CO_{2,HCO3}**), increasing the initial spikes resulting from the wollastonite treatment in both by at least four-fold (purple curves, Fig. S5 b.c). An additional legacy of acidification in North American forests (Harrison et al., 339 340 1989) is SO_4^{2-} retention on soil clay mineral Fe and Al oxides (Fuller et al., 1987), which were subsequently released by 341 increased soil pH following wollastonite weathering (Shao et al., 2016;Fakhraei et al., 2016). To assess the effect of this legacy SO_4^{2-} , we ran simulations for the treated watershed substituting the lower streamwater SO_4^{2-} concentrations from the 342 343 reference watershed (T REF, green curves, Fig. S5b,c). Results suggest that legacy SO_4^{2-} accounts for over half of the total 344 acid deposition effect on increased $[HCO_3^-]$ and CO_2 consumption in the simulations.

In the 'Ref NO₃' scenario (**Fig. S5 d-f**), seasonal spikes in streamwater export of NO₃⁻ recorded from the treated watershed between 2012 and 2015 were removed by substituting the reference watershed streamwater NO₃⁻ concentration measurements lacking these spikes. This manipulation markedly increased modelled bicarbonate (**Fig. S5e**) and mean annual CO₂ consumption (**Fig. S5f**). To quantify the effects of organic acids on bicarbonate production in the treated watershed, we ran "+OA" and "-OA" simulations, i.e., with and without accounting for organic acids, respectively (**Fig. S5 g-i**). Results showed that removing OA from our simulations also increased modelled streamwater bicarbonate concentration (**Fig. S5h**), and resulting CO₂ consumption (**Fig. S5i**), in the treated watershed.

352 **3.3 Effects of increasing wollastonite treatment**

Because the HBEF application rate (3.44 t ha⁻¹) is smaller than the 10–50 t ha⁻¹ suggested for ERW strategies (Strefler et al.,

2018; Beerling et al., 2018), we simulated the possible effects of a ten-fold increase in the streamwater Ca^{2+} concentrations on

bicarbonate production (Fig. S6a) and CO_2 consumption (Fig. S6b). In this initial assessment, we assume streamwater

356 responses are directly proportional to wollastonite application rate, i.e., 34.4 t ha⁻¹, and that all other variables remained

unchanged. Results show that after 15 years, cumulative *Wo-CO_{2,HCO3}* is 73% of *Wo-CO_{2,Ca}* (Fig. S6c), as opposed to less than

358 20% for the actual rate of 3.44 t ha⁻¹ (**Table 3**). These results suggest that at higher application rates of wollastonite, the details

359 of the CO_2 consumption calculations become less important.

360

Table 3. Cumulative fluxes from treatment date calculated with streamwater partial pressure of CO_2 (gas) = $3.63 \times atmospheric$

362 CO₂ partial pressure measured at Mauna Loa (Tans and Keeling, 2017) (see Methods). DIC = dissolved inorganic carbon.

363 Scenarios are defined in the main text.

Cumulative fluxes 1 year post-treatment date (19 October 2000)								
Watershed	Scenario	Org.	CO _{2,ions}	Wo-CO _{2,Ca}	DIC	HCO.	<i>СО</i> 2,нсоз	<i>Wo-CO</i> 2,нсоз
		acids	(Eq. 8)	(Eq. 9)	DIC	11003	(Eq. 5)	(Eq. 7)
				I	mo	C m ⁻²		
REF (6)	baseline	+OA	-0.003	0	0.084	0.002	0.002	0
Treated (1)	baseline	+OA	0.047	0.052	0.086	0.011	0.011	0.011
Treated (1)	baseline	–OA	0.047	0.052	0.094	0.019	0.019	0.018
Treated (1)	Low SO4	+OA	0.083	0.052	0.117	0.043	0.042	0.039
Treated (1)	REF NO3	+OA	0.047	0.052	0.105	0.030	0.030	0.029
Treated (1)	WoX10	+OA	0.513	0.534	0.533	0.457	0.457	0.457
Cumulative	fluxes 15 yea	rs post-	treatment (2	0 November 20	14)			
Watershed	Scenario	Org.	CO _{2,ions} Wo-CO _{2,Ca} DIC CO _{2,HCO3} Wo-CO _{2,HCO3}					
		acids	(Eq. 8)	(Eq. 9)	DIC	ncos	(Eq. 5)	(Eq. 7)
			mol C m ⁻²					
REF (6)	baseline	+OA	-0.274	0	1.307	0.052	0.036	0
Treated (1)	baseline	+OA	-0.044	0.294	1.299	0.083	0.064	0.057
Treated (1)	baseline	–OA	-0.044	0.294	1.414	0.198	0.179	0.145
Treated (1)	Low SO4	+OA	0.269	0.294	1.523	0.307	0.270	0.179
Treated (1)	REF NO3	+OA	-0.044	0.294	1.410	0.194	0.175	0.127
Treated (1)	WoX10	+OA	2.600	3.275	3.626	2.406	2.387	2.380

Cumulative fluxes 1 year post-treatment date (19 October 2000)

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369 **3.4 Amplification of organic carbon sequestration by wollastonite treatment**

370 In reversing long-term Ca^{2+} depletion of soils, the silicate rock treatment significantly increased forest growth and wood 371 production between 2–12 years post-treatment relative to the reference watershed (Battles et al., 2014). This forest response 372 increased total carbon sequestration by 20.7 mol C m⁻² or 9.1 t CO₂ ha⁻¹ during those ten years as a result of the treatment 373 (Methods).

374 Changes in greenhouse gas (GHG) emissions from soils represent a further route to affecting the climate mitigation 375 potential of the wollastonite treatment. Despite a rapid increase of one pH unit in the upper organic soil horizon (Oie), soil 376 respiration CO₂ fluxes showed no significant difference between watersheds during the first three years after treatment 377 (Groffman et al., 2006). However, our analysis of newly available longer-term datasets indicates that the treatment 378 significantly reduced soil respiration in the high elevation hardwood zone (~660–845m a.s.l.) ($\gamma^2(1,270)=17.2$, P < 0.001), 379 possibly due to reduced fine-root biomass (Fahey et al., 2016) rather than changes in microbial activity (Groffman et al., 2006). 380 No significant effects on soil respiration were detected in any of the other HBEF vegetation zones (Fig. 3). The wollastonite treatment increased the soil sink strength for CH₄ ($\chi^2(1,266)=30.8$, P < 0.001) in the low-elevation hardwood zone (482–565m 381 382 a.s.l.), while it decreased in the high elevation zone ($\chi^2(1,268)=22.3$, P < 0.001) (SI Appendix, Fig. S8). There were no 383 significant treatment effects on soil N₂O fluxes in any vegetation zone (SI Appendix).



384

Figure 3: Long-term soil respiration responses to wollastonite treatment at Hubbard Brook Experimental Forest. Cumulative soil CO₂ respiration responses of treated and untreated (a) high elevation hardwoods, (b) high elevation conifers, (c) low elevation hardwoods or (d) mid-elevation hardwoods. Plots show cumulative means ± 1 SE for three chamber measurements at each site and time. Reference data were collected from untreated forests immediately adjacent to the western edge of our reference catchment. *P*-values from Kruskal-Wallis tests comparing treated and reference raw data (SI Appendix) are shown.

390

391 3.5 Logistical CO₂ emissions and net CDR

392 We next considered carbon emissions (penalties) for logistical operations involved in mining, grinding, transporting and

393 applying the wollastonite (Fig. 4, Table 2). In the HBEF experiment, wollastonite was mined and milled on site near

394 Gouverneur, New York. We used CO₂ emissions factors for electricity generation in upstate New York (United States

395 Environmental Protection Agency, 1999) to estimate the maximum CO_2 penalty for mining and grinding to the mean particle

396 size 16 µm diameter (Methods). However, local hydropower (Energy Information Administration, 1997) and regional nuclear

397 power suggest these costs could have been zero. This would represent a substantial carbon saving for the overall ERW process

398 relative to prior expectation of ERW studies in which grinding CO₂ emissions account for up to 30% reduction in ERW-CDR

399 efficiency (Renforth, 2012; Moosdorf et al., 2014).



400

401 Figure 4: Carbon penalties for the wollastonite treatment. Carbon penalties for logistic elements of the treatment are compared with 402 literature estimates for large-scale rollout of enhanced rock weathering for the HBEF treatment (3.44 t ha⁻¹), with and without long-distance 403 transport for pelletization.

404

405 In the HBEF experiment, the milled wollastonite was transported by highway to Allerton, Illinois, for pelletization 406 and then returned to the staging area near Woodstock, New Hampshire (round trip >3150 km). Transportation CO₂ emissions were 0.22-0.61 t CO₂ t Wo⁻¹. Given coal power in central Illinois, we estimate pelletization emitted up to 0.02 t CO₂ t Wo⁻¹ 407 (Methods). Application at Hubbard Brook occurred via 55 \sim 5-km helicopter flights, which gives a further CO₂ cost of 0.01– 408 409 $0.15 \text{ t CO}_2 \text{ t Wo}^{-1}$. In total, these logistical operations emitted $0.23-0.69 \text{ t CO}_2 \text{ t Wo}^{-1}$, or $0.8-2.4 \text{ t CO}_2 \text{ ha}^{-1}$ for the 11.8 ha of the HBEF treated watershed (Table 4). However, local pelletization could have reduced heavy duty vehicle (HDV) transport 410 411 distance to ~400 km and lowered total CO₂ emitted during logistical operations to 0.04–0.15 t CO₂ t Wo⁻¹. At other forested sites, where wind-drift of material is not critical, pelletization may not be necessary. 412

413

- 414 Table 4. Measured elements of the treatment effect on the greenhouse gas budget for the Hubbard Brook Experimental Forest
- 415 wollastonite experiment.

Eqn 14	Greenhouse gas sinks ^a and emissions ^a (t CO _{2e} ha ⁻²)	Pessimistic	Optimistic						
Ecosystem res	Ecosystem responses ^b								
∆wood	Wood production sink increased over ten years ^c	8.946	9.542						
ΔSRESP	Soil respiratory CO ₂ emissions have reduced ^a since 2002	2.213	2.646						
∆CH4	Soil methane sink has increased since 2002	0.015	0.029						
<i>∆N20</i>	Soil N ₂ O emissions since 2002 (no significant difference)	0	0						
	Net ecosystem response at the treatment site through 2014	11.174	12.218						
Downstream s	sequestration and emissions responses								
ΔCONS	CO_2 consumption sink through 2014 (<i>Wo-CO</i> _{2,HCO3} and <i>Wo-CO</i> _{2,Ca})	0.025	0.129						
<i>∆NO3N2O</i>	Downstream N ₂ O emissions ^d from treatment date through 2014	-0.071	-0.016						
ΔDOC	DOC export emissions ^{d,e} from treatment date through 2014	-0.203	0						
	Net downstream balance through 2014	-0.228	-0.129						
Logistics:									
	Mining/Grinding given hydro or nuclear/petroleum power	-0.162	0						
	Helicopter (~55 5-km flights)	-0.051	-0.021						
	HDV transport (New York to Illinois to New Hampshire)	-2.135	-0.787						
	Pelletization (in Illinois, coal power)	-0.068	0						
LOGPEN	Total logistical emissions	-2.416	-0.808						
∆GHG	Partial treatment effect on greenhouse gas balance	8.509	11.523						

⁴¹⁶ ^aDefined as the difference between watersheds: treated–reference for sinks and reference–treated for emissions

⁴¹⁷ ^bSome possible treatment responses such as canopy respiration and particulate organic carbon export are unknown.

418 ^cAfter Battles et al. (2014). We have not attempted to extrapolate these results.

419 d Δ DOC and Δ NO3N2O are penalties because these lead to CO₂ and N₂O emissions downstream.

420 "The "optimistic" value for DOC assumes complete burial and undesireable low oxygen conditions in downstream waters.

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These carbon emission penalties must be subtracted from watershed carbon removal to calculate net CDR for the wollastonite treatment at HBEF (**Fig. 5**; **Table 4**). Compared in this way, we find increased wood production over ten years (Battles et al., 2014) repays the total logistical CO₂ costs 4–12 times over. The components (**Fig. 5**; **Table 4**) comprise 8.5– 11.5 tCO₂ ha⁻¹ of the total GHG budget associated with the wollastonite treatment (Methods). These figures would increase to 10.4–12.2 tCO₂ ha⁻¹ if the wollastonite had been pelletized anywhere along the route from Gouverneur to New Hampshire. Wollastonite treatment effects on streamwater chemistry play a minor role in the greenhouse gas budget (**Fig. 5**; **Table 4**). For our hypothetical ten-fold higher treatment ($34.4 \text{ t} \text{ ha}^{-1}$), CO₂ consumption calculated by assumed calcium release is ~10 times higher, but carbon emission penalties scale with increased rock mass. Assuming pelletization near the mine to reduce transport costs, the total logistical penalty would be 1.2–5.1 tCO₂ ha⁻¹. In total, net CDR would be 6.8–12.4 tCO₂ ha⁻¹ for the the ten-fold larger treatment if none of the other GHG fluxes changed. We have not attempted to extrapolate other forest biomass and soil GHG fluxes or streamwater DOC and NO₃⁻ responses.

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GHG Source GHG Sink

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Figure 5: Carbon responses for the wollastonite treatment. Elements of the greenhouse gas balance associated with the wollastonite treatment (Table 4). The CO₂ consumption range is given by *Wo-CO_{2,HCO3}* calculated by Eq. (7) and *Wo-CO_{2,Ca}* calculated by Eq. (9), time-integrated from the application date through 2014. Nitrate export in streamwater leading to N₂O greenhouse gas emissions downstream and a small increase in the soil CH₄ sink have been converted to CO₂-equivalents (Methods). Exported DOC is assumed to be respired downstream.

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444 **3.6 Potential for deployment at larger scales**

445 The HBEF forests are representative of a major area of eastern North America receiving acid deposition since the 1950s (Likens and Bailey, 2014) which may be suitable for remediation and carbon capture via ERW treatment with a silicate rock 446 447 or mineral. For example, the Appalachian and Laurentian-Acadian Northern Hardwood Forests (NHWF) covering a combined 448 area of 0.137 Mkm² in the United States (Ferree and Anderson, 2013) have the same dominant hardwood trees as the HBEF 449 experimental watersheds (Fagus grandifolia, Betula allegheniensis and Acer saccharum). Acid deposition exceeded "critical loads" likely to harm ecosystems in almost 9000 ha of New Hampshire's Acer saccharum stands (NHAs) (Schaberg et al., 450 2010). These forests might be expected to respond similarly to a wollastonite treatment. The acid-sensitive trees Acer 451 452 saccharum and Picea rubens are also widely distributed along the high elevation acid sensitive regions of the Appalachian 453 Mountains which have already been impacted by acid deposition (Lawrence et al., 2015). We define this as a 40-km corridor 454 along the Appalachian Mountains comprising 0.14 Mkm² and overlapping with the High Alleghenv Plateau Ecoregion (HAL) where Acer saccharum is declining above ~550 m a.s.l. (Bailey et al., 2004) (0.07 Mkm²). 455

456 We examined the potential CO₂ consumption for a range of wollastonite application rates encompassing those suggested 457 for ERW strategies (Strefler et al., 2018; Beerling et al., 2018) (Fig. 6). In this analysis, we adjusted mean (2003–2012) Wo- $CO_{2,ca}$ for the actual 3.44 t ha⁻¹ treatment (~0.2 mol C m⁻² yr⁻¹) proportionally for 10–50 t ha⁻¹ treatments. We assume logistical 458 459 carbon penalties are minimised and balanced by forest biomass carbon sequestration responses to treatment. This analysis suggests net CDR potential of 0.3–1.7 Mt CO₂ yr⁻¹ along the Appalachian corridor, which is 2–12% of New Hampshire state 460 emissions (13.8 Mt CO₂) in 2016 (Energy Information Administration, 2019). However, world wollastonite reserves (Curry, 461 462 2019) (≥ 0.1 Pg) are insufficient to treat large areas of eastern North America at rates of 10–50 t ha⁻¹, highlighting the 463 requirement for alternative sustainable sources of silicate materials.

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Figure 6: Projected CO₂ consumption following higher-dosage treatments. We considered the possibility of higher-dosage silicate treatments on other northeastern United States higher-altitude forests affected by acid rain, such as *Acer saccharum* forests in New Hampshire (NHAs), the High Allegheny Plateau Ecoregion (HAL), the Appalachian trail corridor (AT), or Northern Hardwood forests (NHWF) dominated by the same tree species as at Hubbard Brook. Because the world's wollastonite reserves (yellow disks) are insufficient to treat these areas, other calcium-rich silicate minerals would be required. CO₂ consumption due to higher dosage (t ha⁻¹) is estimated as: (mean observed CO_{2,Ca} between 2004 and 2012) × area × dosage / 3.44 t ha⁻¹.

473

474 4 Discussion

475 Our analyses of wollastonite application at the HBEF provide a unique long-term (15 year) perspective on the whole watershed

476 carbon cycle responses and net CDR by accounting for the associated CO_2 costs of logistical operations. By 2015, net CDR

477 amounted to 8.5-11.5 t CO₂ ha⁻¹ at a low rate of wollastonite application, with increased carbon sequestration into forest

478 biomass playing the dominant role. We estimate that if the HBEF application rates were increased ten-fold, net CDR would

479 increase by 8%, assuming 400-km transport distances and no change in forest responses. Amplification of organic carbon 480 capture may therefore represent a major CDR benefit of ERW when applied to forested lands affected by acid rain. Forest 481 management practices, disturbance regimes and the ultimate fate of any harvested wood are also important in determining the 482 storage lifetime of the sequestered carbon. Our results highlight the need to carefully monitor the net carbon balance of forested 483 ecosystems in response to a silicate treatment, including wood and canopy respiration (Fahey et al., 2005) (Methods). This 484 challenging goal might best be achieved with fully instrumented eddy covariance plots, although the HBEF topography is not 485 well suited for this approach (Fahey et al., 2005).

486 Inorganic CO₂ consumption calculated based on streamwater bicarbonate fluxes approximately doubled in the treated watershed relative to the reference watershed 15 years post-treatment (0.028 and 0.016 tCO₂ ha⁻¹, respectively) (**Table 3**). The 487 488 presence of $SO_4^{2^-}$, NO_3^{-} and organic acid anions lowered the efficiency of CO_2 consumption by alkalinity generation, with acid 489 deposition having the single largest calculated effect (**Table 3**). The cause of increased NO_3^- export from the treated watershed 490 is not as yet understood (Rosi-Marshall et al., 2016). If it proves a general feature of terrestrial ecosystem responses to silicate 491 mineral treatment, this could affect the efficiency of carbon capture via bicarbonate export. Overall, we suggest that continued 492 recovery of eastern North American and European forests and soils from acid deposition creates conditions beneficial to 493 watershed health, carbonic acid-driven weathering and inorganic carbon export following application of crushed silicate 494 minerals.

In Asia, acid rain is an ongoing problem with an estimated 28% of Chinese land area (~2.7 Mkm²) receiving potentially 495 496 damaging S deposition in 2005 (Zhao et al., 2009), and critical loads were exceeded in ~0.36 Mkm² of the European Economic 497 Area (EEA) in 1999 (Larssen et al., 2003), approximately double the affected area of US Northern Hardwood Forests (Fig. 6). 498 Fig. 6 suggests that a single 30t Wo ha⁻¹ treatment over 0.14 Mkm² (Appalachian Trail corridor) could, in principle, sequester ~1 MtCO₂ y⁻¹ or 15 MtCO₂ over 15 years via wollastonite-derived Ca export in streamwater alone. Adding the Chinese and 499 European acidified areas could potentially sequester 0.34 GtCO₂, approximately 0.2–0.7% of the ~50–150 Gt CDR required 500 501 by 2050 to avoid warming in excess of 1.5° (Rogelj et al., 2018). Inclusion of biomass and soil responses increases CDR 502 contributions from ERW on acidified forests, but these will still be modest. Assuming no further forest responses beyond the 503 15-year HBEF timeframe, we report a GHG balance of ~10 tCO_{2e} ha⁻¹. This translates to 1 GtCO_{2e} Mkm⁻² suggesting 3.2 GtCO_{2e} over 15 years for the Appalachian Trail, the EEA and China combined, or 2–6% of global required CDR as described 504 505 above.

It is uncertain whether other acidified forest ecosystems would respond similarly to the HBEF *Acer saccharum* forests in New Hampshire. Many Chinese soils (Duan et al., 2016), as well as old deep soils in areas such as the Virginian Blue Ridge Mountains and the German Harz and Fetchel Mountains (Garmo et al., 2014) have high SO_4^{2-} sorption capacity. These soils may retain substantially more SO_4^{2-} than the HBEF soils, with potential for prolonged SO_4^{2-} flushing following ERW treatment and lower bicarbonate production. Liming studies suggest a range of other effects, some of which may also occur with silicate treatments. Liming increases nitrate export, migration of heavy metals and acidity to deeper soil, and fine root production in topsoils leading to frost damage (Huettl and Zoettl, 1993). 513 Many forests have been limed with carbonate minerals such as calcite and dolomite to mitigate acidification in the past. 514 Dolomite has also helped reverse Mg deficiency in conifers (Huettl and Zoettl, 1993). Liming generally improves water 515 quality, although it also forms mixing zones with high-molecular-weight Al complexes toxic to fish (Teien et al., 2006). With 516 silicate treatments, nontoxic hydroxyaluminosilicates form instead (Teien et al., 2006). Unfortunately, carbonates are 517 contraindicated for CDR on acid soils because they can be a net source of CO₂ in the presence of strong acids (Hamilton et al., 2007). Treatments of European and North American acidified forests with calcite (1-18 t ha⁻¹ CaCO₃) or dolomite (2-8.7 t 518 519 ha $^{-1}$ CaMg(CO₃)) have, in general, resulted in increased DOC export and soil respiration without increasing tree growth, 520 regardless of forest composition (Lundström et al., 2003). As calcite and dolomite are 44% and 48% CO₂ by weight, these treatments will have released 0.44-7.9 and 0.96-4.54 t CO₂ ha⁻¹ respectively when fully dissolved, although dissolution may 521 be slow. Over six years following a 2.9 t dolomite ha⁻¹ treatment (90% 0.2–2.0 mm grains) in a Norwegian coniferous 522 watershed equating to 1.36 t CO₂ ha⁻¹, less than 1% of the dolomite dissolved (Hindar et al., 2003). We estimate that CO₂ 523 524 consumption corrected for CO₂ release and as measured with dolomite-derived Ca and Mg in streamwater (Dol-CO_{2,Ca+Mg}) averaged 0.02 mol CO₂ m⁻² yr⁻¹. CO₂ release from carbonate minerals equals Ca and Mg release on a molar basis, so 0.02 mol 525 Dol-CO₂ m⁻² vr⁻¹ was also either exported in streamwater or lost to the atmosphere. This experiment may have a negative 526 527 greenhouse-gas balance depending on logistical penalties and soil respiration, as there was no significant treatment effect on 528 tree growth or vitality (Hindar et al., 2003). Ca-sensitive Acer saccharum is present at Woods Lake in New York State, yet 529 tree biomass decreased with no significant differences relative to reference catchments during the 20 years following a 6.89 t 530 Mg-calcite ha⁻¹ application (Melvin et al., 2013), equivalent to 3.07 t CO₂ ha⁻¹ given 8% Mg content of the pellets. In contrast 531 to our study and other liming studies, root biomass and soil carbon stocks increased in response to this treatment, although soil 532 respiration was reduced (Melvin et al., 2013). Acer saccharum basal area and crown vigour increased over 23 years in response to 22.4 t dolomitic limestone ha⁻¹ (equivalent to 10.0 t CO_2 ha⁻¹) on the Allegheny Plateau, although basal area and survival of 533 another dominant canopy species, Prunus serotina, was reduced (Long et al., 2011). Clearly, forest responses to mineral 534 535 treatments are species- and site-specific.

536 Although the HBEF experiment used wollastonite, this is not a target mineral for ERW, both because of its limited 537 reserves (Curry, 2019) and high monetary costs (Schlesinger and Amundson, 2018). Recent all-inclusive guide prices of ~700 USD Mg⁻¹ for helicopter deployment of pelletized lime along the Appalachian Mountain corridor are comparable to the price 538 of 694 USD Mg⁻¹ for unpelletized 10-µm wollastonite in 2000 (Virta, 2000). Less expensive materials such as locally-sourced 539 540 waste fines from mines or volcanic ash (Longman et al., 2020) should be considered if their heavy metal content is low, but 541 the choice of treatment material should be considered together with the vegetation and the native minerals. Application of 542 magnesium-rich materials (e.g. olivine), for example, may help reverse Mg deficiency in *Pinus sylvatica* and *Picea abies* as 543 dolomite has done (Huettl and Zoettl, 1993), but some other tree species, such as Acer saccharum, have a higher demand for 544 calcium than for magnesium (Long et al., 2009). The treatment of ecologically sensitive catchments always requires caution 545 as some species, such as *Sphagnum* mosses and lichens, may respond poorly to treatment (Traaen et al., 1997).

546 Finally, we consider integration of ERW treatments with forest management practices. Dominant CDR pathways 547 depend on biogeochemical cycling which in turn depends on the life cycle of the dominant trees. For example, base cation 548 export and therefore CO₂ consumption temporarily increases following clear-felling, then decreases while trees are young and 549 growing due to base cation uptake, and remains low after trees mature due to nutrient recycling (Balogh-Brunstad et al., 2008). 550 These dynamics may be less obvious in forests which are not clear-felled; Acer saccharum forests are often thinned and retain 551 a canopy as the seedlings are adapted to shade. Individual Acer saccharum trees can live for over 300 years, growing relatively slowly for the first 40 years and attaining maximum height during the first 150 years (Godman et al., 1990). One may expect 552 553 to maximise wood production of growing trees with ERW treatments meeting or exceeding the forest demand for previously 554 limited nutrients such as calcium, which would also minimise soil respiration if the trees allocate less carbon to roots. 555 Treatments could be repeated as necessary to meet the nutritional needs of sensitive trees or to maintain high CO₂ consumption. 556 Conversely, rising soil pH may not suit some species. For example, Acer saccharum normally grows in organic-rich soils with pH under 7.3 (Godman et al., 1990) and its growth may be hindered at higher pH following large treatments. Outside the main 557 558 tree growth phase, and in forests without responsive tree species, CO₂ consumption could become the dominant GHG response 559 to ERW treatments depending on the extent to which it is counteracted by DOC export as soil pH rises (Johnson et al., 2014) 560 and decomposition rates and fluxes rise (Lovett et al., 2016). Site-specific research is required to determine the optimum 561 dosage, timing, efficacy and suitability of ERW treatments on acid-impacted forests.

562 Code availability

The aqueous geochemistry software PHREEQC software, along with documentation, is freely available from the USGS website (<u>https://www.usgs.gov/software/phreeqc-version-3</u>). MATLAB[®] may be purchased from the MathWorks website (<u>https://uk.mathworks.com/products/matlab.html</u>). Our MATLAB code and scripts used for this project are provided as a supplementary .zip file, without guarantees that these will run with MATLAB versions other than R2016a or on non-Linux operating systems.

568 Data availability

- 569 Our data are available from the Long Term Ecological Research (LTER) Network Data Portal. This public repository can be
- 570 accessed via the Hubbard Brook Ecosystem Study website: https://hubbardbrook.org/d/hubbard-brook-data-catalog
- 571 See Supplement for a full list of filenames, package IDs, DOIs and access dates.

572 **Author contributions**

573 All authors contributed to project conceptualization and interpretation of model results. L.L.T. undertook model simulations

and data analysis. L.L.T. and D.J.B. drafted the manuscript with edits and revisions from all authors. C.T.D. designed the 575 wollastonite watershed study, provided data and observations for model simulations. J.D.B. provided strontium isotope

576 datasets. P.M.G. provided soil respiration, nitrous oxide and methane flux data.

577 **Competing interests**

578 The authors declare that they have no conflict of interest.

579 Disclaimer

574

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