

Weathering without realizing inorganic CO₂ removal revealed through base cation monitoring.

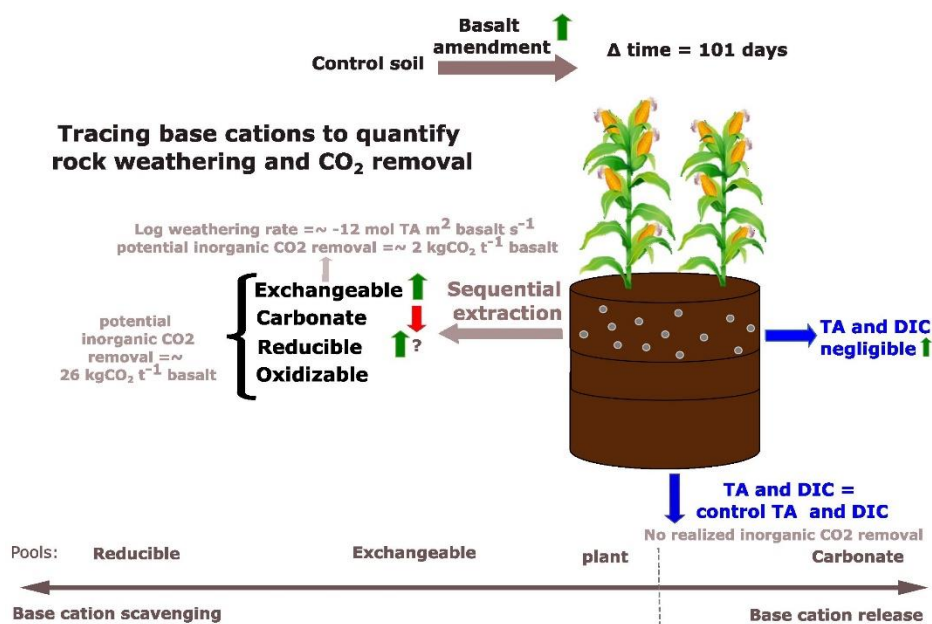
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Graphical abstract



Abstract

Enhanced Weathering using basalt rock dust is a scalable carbon dioxide removal (CDR) technique, but quantifying rock weathering and CDR rates poses a critical challenge. Here, we investigated realized inorganic CO₂ removal (defined as the sum of the change in dissolved inorganic C leaching and in solid inorganic C) and weathering rates by treating mesocosms planted with maize with basalt (0, 10, 30, 50, 75, 100, 150 and 200 t ha⁻¹) and monitoring them for 101 days. We observed no significant realized inorganic CO₂ removal, as leaching of dissolved inorganic carbon did not increase, and soil carbonate content declined over time.

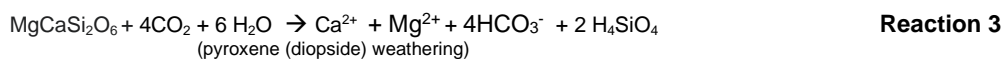
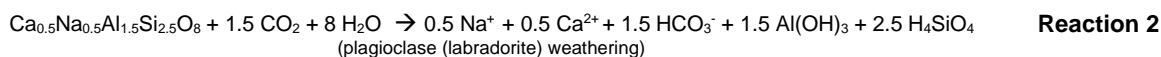
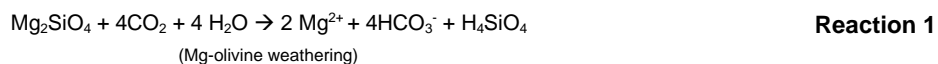
To gain insights into the weathering processes, we traced the fate of base cations in the soil and plants. This analysis showed that most base cations were retained in the topsoil reducible soil pool, typically associated with iron (hydr)oxides, while increases in the exchangeable pool were about a factor 10 smaller. Soil base cation scavenging exceeded plant scavenging by approximately two orders of magnitude. From the base cations in all pools (soil, soil water and plants), we quantified log weathering rates of $-11 \text{ mol total alkalinity m}^{-2} \text{ basalt s}^{-1}$. The

34 potential inorganic CO₂ removal, defined as the maximum inorganic CO₂ removal achievable if all weathered base
 35 cations, adsorbed by soil pools in this experiment, would leach out of the soil and be fully balanced by carbonate
 36 anions, was estimated at 26 kg CO₂ t⁻¹ basalt.

37 In conclusion, despite clear weathering of basalt rock, we found no inorganic CO₂ removal within the timescale of
 38 this experiment. The observed increase of aluminum in association with the reducible soil fraction indicate the
 39 formation of secondary minerals. These, along with enhanced base cation exchange, may contribute to long-term
 40 soil fertility and promote the stabilization of soil organic matter.

41 1. Introduction

42 To meet the "well below 2°C warming" target established by the United Nations' Paris Agreement, Carbon Dioxide
 43 Removal (CDR) must complement conventional climate change mitigation efforts (Minx et al., 2018). One CDR
 44 technology under consideration is enhanced weathering (EW). EW relies on accelerating natural weathering
 45 reactions of silicate minerals with water (H₂O) and carbon dioxide (CO₂) (as in **Reactions 1 to 3**), which increases
 46 the concentration of base cations and dissolved inorganic C (DIC) in water, delivering inorganic CO₂ removal. In
 47 this study, we focus on DIC export from soils to the ocean, as this pathway is considered the most durable form of
 48 carbon sequestration (Phil Renforth & Henderson, 2017), rather than aiming to quantify a full greenhouse gas
 49 budget. DIC (the sum of aqueous [CO₂], [HCO₃⁻] and [CO₃²⁻]) can either be measured directly or estimated
 50 indirectly from total alkalinity (TA) or electrical conductivity, which are less expensive to monitor and can be
 51 empirically linked with DIC through calibration curves (Amann & Hartmann, 2022) (see also **Fig. S10**). This
 52 calibration is feasible because, according to the explicit conservative expression for TA in water, TA = [HCO₃⁻] +
 53 [CO₃²⁻] + [OH⁻] – [H⁺] (Wolf-Gladrow et al., 2007). TA can also be approximated from the sum of base cation
 54 charges, minus the sum of conservative anion charges (e.g. chloride, sulphate, phosphate, nitrate) (Barker, 2013;
 55 Wolf-Gladrow et al., 2007). DIC can also precipitate as soil inorganic carbon (SIC) in the form of solid carbonates,
 56 thereby losing half of the initially captured CO₂ (**Reaction 4**) (Haque et al., 2019). Inorganic CO₂ removal can be
 57 defined as the sum of changes in DIC leaching from a soil system after rock amendment plus the change in solid
 58 inorganic C (SIC) within the soil system after rock amendment. A robust and reliable accounting of Inorganic CO₂
 59 removal must thus include both monitoring of DIC leaching and SIC changes.



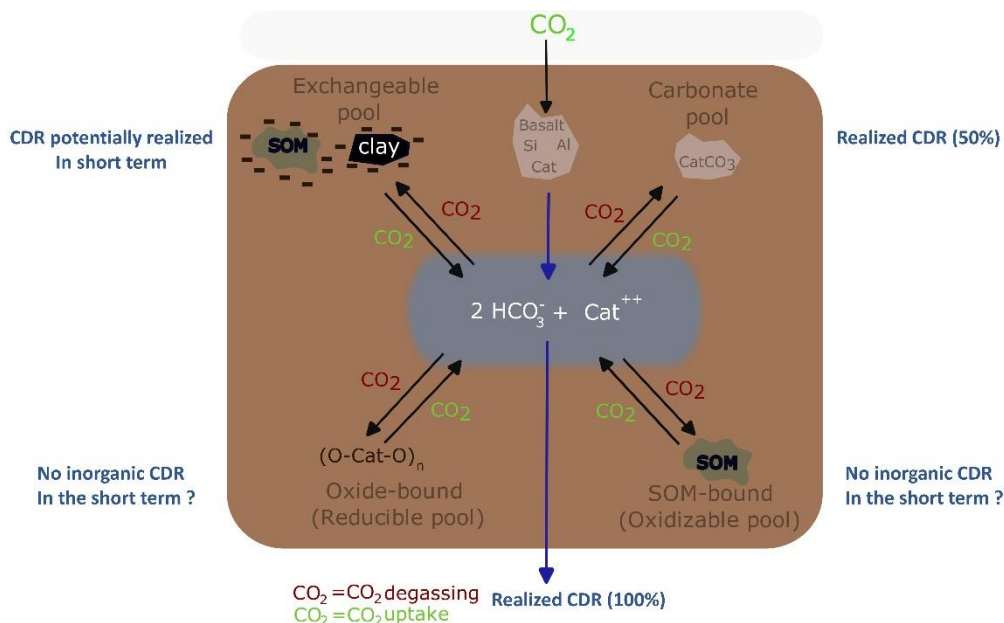
60 EW is an attractive CDR technology for several reasons. First, EW may provide long-lived to permanent CO₂
 61 sequestration: if fixed, DIC is transported via rivers or groundwater to oceans where it may not be released back
 62 into the atmosphere for millennia, the timescale needed for oceanic carbonate precipitation, which would release
 63 50% of the DIC input back into the atmosphere (**Reaction 4**) (Renforth & Henderson, 2017). Secondly, rock dust
 64 amendment has the potential to improve soil fertility and counters soil acidification (Swoboda et al., 2021; Van
 65 Straaten, 2006). Thirdly, unlike some other CDR technologies (such as bio-energy with carbon capture and storage
 66 or afforestation), EW avoids competition for land with food production (Fuss et al., 2018; Janssens et al., 2022;
 67 Smith et al., 2016). Although several rock types are considered for EW, basalt is typically used in EW field trials
 68 and has several advantages. Basalt has relatively high base cation content, particularly of calcium (Ca) and
 69 magnesium (Mg), which translates into a high potential for CO₂ removal (Renforth et al., 2019). Additionally, basalt

70 is comprised of mafic silicate minerals such as plagioclases, pyroxene, and olivine, known for their relatively high
 71 weathering rates (W_r). Furthermore, basalt formations are abundant, widely distributed and close to major
 72 economies, making the adoption of EW using basalt scalable. Importantly, basalt is safer for agricultural application
 73 compared to ultramafic rocks like dunite due to its lower content of heavy metals such as nickel (Ni) and chromium
 74 (Cr) (Beerling et al., 2020).

75 Despite the great potential of terrestrial EW and substantial attention by industry in recent years, monitoring rock
 76 weathering and CDR is challenging. Quantification of Inorganic CO_2 removal by EW has often focussed on tracking
 77 DIC or alkalinity leaching in porewaters (Holzer et al., 2023; McDermott et al., 2024). However, it is also important
 78 to consider DIC in exported soil water (leachates)(Larkin et al., 2022) as changes in DIC during soil water transport
 79 are well-established. Numerous studies demonstrated that soil water movement and pH strongly govern DIC
 80 dynamics, both in soil research (Öquist et al., 2009; Schindlbacher et al., 2019) and in EW research (Dietzen et al.,
 81 2018; Niron et al., 2024; Reynaert et al., 2023; Vienne et al., 2024).

82 Focusing solely on changes in DIC and SIC may however overlook other critical soil processes that impact CDR.
 83 Besides the carbonate soil pool, other solid soil pools can also extract base cations from solution (**Figure 1**). These
 84 pools (temporarily) trap base cations, preventing DIC leaching and could stabilize soil organic matter (SOM) (Buss
 85 et al., 2024). Here, we trace the fate of cations in four different soil pools, to gain better estimates of W_r s and CDR.

86



88 **Figure 1:** Schematized weathering of aluminosilicate rock and four soil pools that scavenge base cations (= alkalinity): exchangeable pool, carbonate pool, reducible pool and oxidizable pool. Because of charge balance,
 89 uptake of base cations by these pools releases H^+ that can reconvert bicarbonate (HCO_3^-) (that was a priori
 90 generated from CO_2 through weathering) into CO_2 . Cat^{++} = one divalent base cation or two monovalent base
 91 cations. In each corner, in blue, the significance for CDR is indicated for each of the soil pools.
 92

93 Tracing base cations in soils can be done based on the established methodology of Tessier et al. (1979) in which
94 cations are partitioned into four operationally defined soil pools; The exchangeable pool, the carbonate pool, the
95 reducible pool and the oxidizable pool. First, in the exchangeable soil pool, cations interact with negatively charged
96 clay or SOM surfaces. Secondly, the carbonate pool contains carbonates such as calcium carbonate (CaCO_3) and
97 the C in this pool is reported as SIC. The detection of changes in SIC in basalt amended soils in short-duration
98 experiments is typically challenging (Kelland et al., 2020; Vienne et al., 2022). Focusing on carbonate base cations
99 may avoid typical issues with C heterogeneity and detection of relatively small SIC changes.

100 Thirdly, Tessier et al. (1979) operationally defined a reducible soil pool where base cations are associated with iron
101 (Fe) and manganese (Mn) (hydr)oxides, and an oxidizable pool where cations are bound to SOM or sulfides.
102 Dzombak & Morel (1990) modelled adsorption of Mg to hydrous ferric oxides ($\text{FeO}(\text{OH})$), in which a surface
103 hydroxyl group loses a proton and is replaced by a magnesium ion ($\text{FeOOH} + \text{Mg}^{2+} \rightleftharpoons \text{FeOMg} + \text{H}^+$) and thereby
104 decreases solute TA. In the fourth considered soil pool, the oxidizable pool, organic functional groups such as
105 carboxylic and phenolic groups can form strong bounds with cations after deprotonation (Kalinichev et al., 2011).
106 Cations in the oxidizable pool are expected to chemically stabilize organic matter due to cation bridging and
107 inhibition of decomposing enzymes (Rowley et al., 2018).

108 In conclusion, base cations in these soil pools could decrease solute TA after proton release and hence degas DIC
109 while potentially stabilizing SOM. Last, besides soil pools, also plants can scavenge base cations from solution.
110 Base cations that go to the plant pool can be recycled to the aqueous phase, either through decomposition of plants
111 in the field or through the food chain and sewage system, further complicating base cation mass balancing.

112 The undesirable side-effect of base cation scavenging (by plant/soil pools) is the release of protons to maintain
113 charge balance. This release of protons converts negatively charged DIC (HCO_3^- and carbonate anions (CO_3^{2-})) to
114 H_2CO_3 , which is in equilibrium with gaseous CO_2 ($\text{CO}_3^{2-} + \text{H}^+ \rightarrow \text{HCO}_3^-$ and $\text{HCO}_3^- + \text{H}^+ \rightarrow \text{H}_2\text{CO}_3 \rightleftharpoons \text{H}_2\text{O} + \text{CO}_2$
115 (g)). Hence, inorganic CO_2 removal can be reversed during temporary storage of base cations and realized again
116 when base cations are released back from soil and plant pools into the aqueous phase.

117 From base cations in plant and soil pools, we can thus calculate a 'potential inorganic CO_2 removal', a terminology
118 proposed by Steinwidder et al., (2025). This is a maximum quantity of Inorganic CO_2 removal that can be achieved
119 when all cations released through silicate weathering are charge-balanced by bicarbonate/carbonates and leached
120 from soils. Base cation retention in different soil pools results in a temporal decoupling between weathering and
121 inorganic CO_2 removal. The timeframe in which the potential inorganic CO_2 removal can be achieved is a major
122 uncertainty in EW (Kanzaki et al., 2025). For weakly bound exchangeable cations, potential inorganic CO_2 removal
123 may be achieved in the relative short term of decades. Within this timeframe, because of stronger binding strengths,
124 reducible and oxidizable base cations are more unlikely to be released and thus deliver inorganic CO_2 removal.

125 Last, inorganic CO₂ removal is only achieved if the weathering agent that induced the weathering was H₂CO₃ (as
 126 in Reaction 1-3). If the weathering agent is another acid (e.g. nitric acid (HNO₃) from fertilizers), no inorganic CO₂
 127 removal occurs (McDermott et al., 2024; Taylor et al., 2021).

128 In a mesocosm experiment with basalt rock powder addition, we aimed to accurately quantify the W_r and potential
 129 inorganic CO₂ removal through quantification of base cations in the four abovementioned soil pools, soil water and
 130 maize plants. Tracing the fate of alkalinity after its generation by the weathering of primary minerals is key to
 131 accurately quantify basalt W_rs. Here, we make a mass balance after 101 days of experiment, investigate the fate
 132 of base cations through exploration of sequential extractions as a monitoring strategy for weathering and
 133 implications for C sequestration.

134 2. Materials and Methods

135 2.1 experimental set-up

136 A mesocosm experiment with 30 mesocosms was constructed at the experimental site at the Drie Eiken Campus
 137 of the University of Antwerp (Belgium). This experiment was part of a larger mesocosm experiment that aimed to
 138 investigate heavy metal fate and plant biomass in silicate amended maize plants (Rijnders et al., 2025). The
 139 mesocosms (0.6 m height, radius=0.25m) received natural rainfall and received additional water through manual
 140 irrigation (**Fig. S2**). In May 2021, the lower 40 cm of each mesocosm was filled with a slightly acidic sandy loam
 141 soil (**Table 1**).

142 **Table 1:** properties of control soil.

Control soil properties*	
pH (in a soil: water suspension (1:2.5))	5.66 ± 0.01
Texture (Sand, clay, silt %)	Sandy loam (61, 4, 35 %)
Soil organic C (SOC) (%)**	0.53 ± 0.01
SIC (%)	0.0031 ± 0.0002
Cation exchange capacity (CEC) (meq/100g)	3.03 ± 0.11
Base saturation (%)	50 ± 5
Bulk density (BD) (kg/L)	1.58±0.02

143 *Reported values represent the average ± standard error (SE) of control soil sampled at all depths after the experimental period
 144 of 101 days. ** Determined through loss on ignition (4h heating at 360°C and assuming a SOC/SOM ratio of 0.58 (Van Bemmelen,
 145 1890)).

146 The upper 20 cm was filled with the same soil, either unamended in the control treatment (5 mesocosms), or
 147 amended with basalt (**Figure 2**). Five mesocosms received 50 ton basalt ha⁻¹, while six others received different
 148 amounts of basalt, ranging between 10 and 200 ton ha⁻¹ (**Table 2**). The basalt was mixed homogenously in the
 149 control soil using a concrete mixer. Basalt was obtained from DURABAS (<https://www.rpbl.de>). Particle size
 150 distribution (PSD) was analyzed using a mastersizer 2000 with a Hydro 2000G sample dispersion unit after
 151 removing larger particles with a 2 mm sieve. The P80 was 310.78 μm (see **Fig. S5**). The specific surface area
 152 (SSA) was determined with a Quantachrome Autosorb iQ using the Braunauer-Emmet-Teller (BET) method. The

153 measurement used nitrogen gas as adsorbate with multi-point (5 points) and isotherm (77K) settings. Samples with
 154 the same treatment were pooled in equal quantities into one sample to reduce the cost and time for analysis. All
 155 samples were degassed at 300 °C with 200 minutes of soak time. High measurement quality was ensured by
 156 frequent reference measurements (Bundesanstalt für Materialforschung und -prüfung, Germany) in addition to three
 157 technical repetitions for each measurement. The BET-SSA of the basalt rock was $9.226 \pm 0.08 \text{ m}^2 \text{ g}^{-1}$. X-ray
 158 diffraction (XRD) and x-ray fluorescence (XRF) analyses are provided in the supplement.

159 **Table 2:** Overview of basalt application rates. The 0 and 50 t basalt ha⁻¹ application rates were replicated in five
 160 mesocosms, while other application rates were only tested in one mesocosm. We added these replicates within
 161 individual application rates to learn about the variability between mesocosms receiving the same treatment.

Ton silicate ha ⁻¹ (replications)	0 (5x)	10	30	50 (5x)	75	100	150	200
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162
 163 Basalt was mixed into the top soil on 17/5/2021. To allow leachate collection, mesocosms had a 2 cm diameter
 164 hole at the bottom. On the inside, the bottom of the pot was covered with a root exclusion mesh to prevent soil
 165 export. Glass collectors (2.3L volume) were placed under the mesocosm to collect the leachates. Leachate volumes
 166 were determined throughout the experiment and were collected for chemical analyses on seven occasions. On
 167 3/6/2021, two sweet maize seedlings (variety Tom Thumb) were planted in each the mesocosms and all pots
 168 received fertilization with nitrogen (N), phosphorus (P) and potassium (K) (96 – 10 – 79) kg ha⁻¹ by adding Ca nitrate
 169 (Ca(NO₃)₂), triple super phosphate (TSP, 45% P₂O₅) and potassium sulfate (K₂SO₄). The experimental duration
 170 was 101 days; plants were harvested on 26/8/2021.

171 Soil water content and temperature were recorded using Cambell Scientific sensors (CS616) that are 30 cm in
 172 length. Watering (using rain water collected from a tank) was executed manually and total water manual inputs
 173 were tracked. In addition, daily precipitation amounts (in mm) were obtained for Wilrijk (Belgium) using the open
 174 source tool (visualcrossing.com). In the supplement, an overview of environmental conditions (rainfall, total water
 175 inputs, temperature and soil moisture) is given.

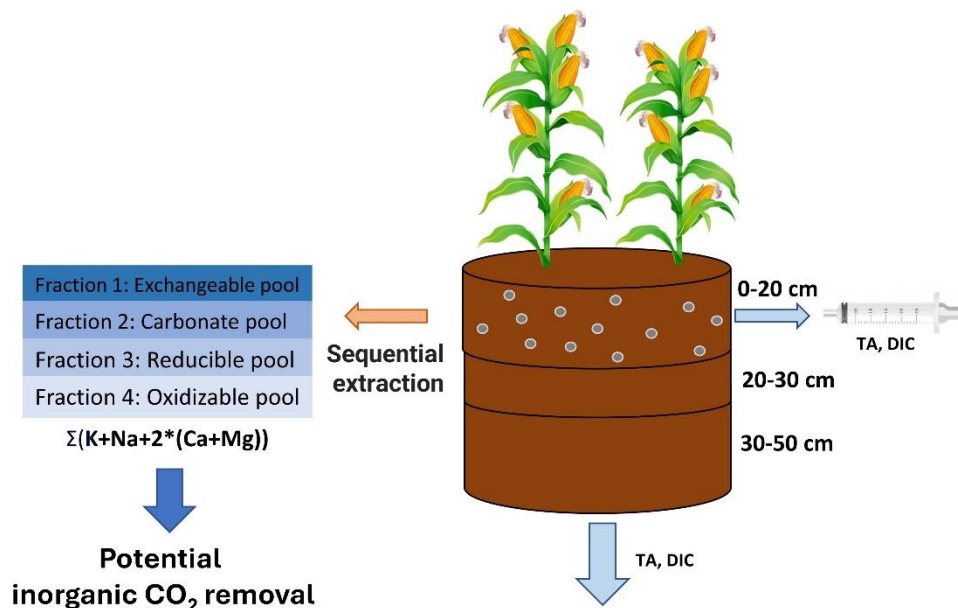


Figure 2: Overview of experimental set-up and measurements.

2.2 Leachate and pore water analysis

Weekly pore water sampling was performed with rhizons (Rhizon Flex, Rhizosphere Research Products B.V., Wageningen, NL) installed at 5 cm depth in each mesocosm. Leachate and porewater samples were filtered through a 0.45 μm PET filter. The major cations (Ca, Mg and K) were measured through ICP-OES (iCAP 6300 duo, Thermo Scientific). Before analysis, ICP samples were conserved using 1.5 mL (HNO_3 69%) per 30 mL sample. TA was determined using a SAN++ continuous flow analyzer (Skalar - NLD). The pH was measured using a HI3220 pH/ORP meter. Dissolved organic carbon (DOC) and DIC were measured using a FormacsHT with LAS sampler (Skalar - NLD). DIC and DOC were measured on eight and 12 occasions in leachates and pore water respectively.

Two quality control (QC) standards were analyzed for individual elements (Ca, K, Mg, sodium (Na), silicium (Si) and Fe). The mean precision of the QC standards was 0.84%, 1.12%, 0.54%, 2.79%, 1.67% and 1.30% for the respective elements. The mean accuracy for the two QC standards was 1.87%, 2.30%, 0.17%, 1.88%, 1.39% and 2.65% for Ca, K, Mg, Na, Si and Fe respectively. For TA soil water samples, mean accuracy and precision for two different QC standards were 1.51 and 1.72% respectively. The DIC measurements with FormacsHT had an accuracy and precision 1.09 and 0.23% respectively. Accuracy and precision were determined based on 12 measurements of a QC for TA (standards: 150 and 350 $\text{mg CaCO}_3 \text{ L}^{-1}$) and DIC (range between 10 and 100 mg L^{-1}) and based on eight measurements of two different QC concentrations for each individual element.

2.3 Soil collection and pretreatment

Top soil pH was measured on five dates. To determine top soil pH, 4 g of air dried topsoil sample was dissolved in 10 mL deionized water and shaken before pH measurement using a HI3220 pH/ORP meter (Hanna Instruments, Temse, Belgium). After harvesting, soils were sampled using cylindrical soil cores (100 cm^3 , 5 cm length x 20 cm^2).

198 Samples were taken across the depth of the mesocosm and three sampling depths were considered (0-20cm, 20-
 199 30cm, 30-50cm). The cores were dried at 70°C for 48 hours to determine water content (gH₂O/g soil) and bulk
 200 density. An additional soil sample was taken at each depth, dried at 70 °C for 48 hours and used for chemical
 201 analyses.

202 2.4 Sequential base cation extractions

203 As conceptualized by Tessier et al. (1979), base cations can reside in four different soil pools: the exchangeable
 204 pool (where O-atoms on hydroxyl or carboxyl groups of clays or SOM associate with cations), the carbonate pool
 205 (cations bound in pedogenic carbonates), the reducible pool (cations bound to Al/Mn/Fe hydr(oxide)) and the
 206 oxidizable pool (cations bound to SOM). SOM bound to cations, extracted with weak salt solutions in the
 207 exchangeable pool typically has a low turnover time (Poeplau et al., 2018) and is therefore thought to be more
 208 susceptible to microbial decomposition than oxidizable SOM.

209 We adapted the original Tessier scheme by replacing 1M magnesium chloride with 1M ammonium acetate
 210 (NH₄(CH₃COO)) for extraction of the exchangeable pool, in order to be able to measure all base cations in the
 211 exchangeable pool. Likewise, Na-acetate was replaced with a mixture of acetic acid and water to be able to measure
 212 Na in the carbonate pool. We quantified SIC changes from the base cations in these acetic acid extracts (as in
 213 (Larkin et al., 2022)) (see also Equation S4). Additionally, three other SIC measurement techniques were explored
 214 to compare and the sensitivity of detecting SIC changes after amending with a range of basalt (see section S3.7).

215 **Table 3:** overview of sequential extraction method
 216 (extraction time, temperature, conditions, volume of extractants and chemical composition of extractants).

217 Extraction scheme	218 Extraction scheme Adapted Tessier et al. (1979)*
219 Pool 1: Exchangeable pool	10 mL 1M NH ₄ (CH ₃ COO) 1h, 20°C, shaker → centrifuge → sample
220 Pool 2: Carbonate pool	5 mL 1M acetic acid (2h, 20°C, shaker) + 4 mL H ₂ O + 1 mL 3M NH ₄ Acetate → sample
221 Pool 3: Reducible pool	20mL 0.04M hydroxylamine (NH ₂ OH.HCl) in 25% (v/v) acetic acid (pH 2) 6h, 96°C, heat bath
222 Pool 4: Oxidizable pool	3mL 0.02M HNO ₃ +5mL 30% peroxide (H ₂ O ₂) (to pH 2 with HNO ₃): 2h, 85°C, heat bath +3mL 30%H ₂ O ₂ (to pH 2 with HNO ₃) 3h, 85°C, heat bath +5mL 3.2M NH ₄ (CH ₃ COO) (in 20 vol% HNO ₃) +4 mL H ₂ O 0.5h, 20°C, shaker

227 Prior to extractions, approximately 1g of soil was air dried. We also conducted the extractions for the pure basalt
 228 that was initially added to the mesocosms to be able to correct for the cations that were initially already present as
 229 exchangeable, carbonate, reducible and oxidizable pool cations. After each extraction, samples were centrifuged
 230 for 2 minutes at 2000 rpm, supernatants were collected for analysis. The remaining soil pellet after centrifugation
 231 was washed with 10 mL of demineralized water before the following step. Relevant elements (K, Na, Mg, Ca, Al,
 232 Fe and Si) were measured in each pool using ICP-OES (iCAP 6300 duo, Thermo Scientific) for each pool. Si was

233 only assessed in the reducible pool and in the oxidizable pool to investigate whether Si forms amorphous oxides or
234 allophane-like compounds or binds with organic matter. Al carbonates were not quantified here as naturally these
235 carbonates are not commonly formed (Takaya et al., 2019).

236 **2.5 Plant responses**

237 On 26/8/2021(101 days after basalt amendment in soils), the aboveground biomass was harvested and dried for
238 48h at 70 °C to determine dry weight. Plant material was ground with an ultra-centrifugal mill (Model ZM 200, Retsch
239 GmbH, Haan, Germany). Base cations (Ca, Mg and K) were measured through ICP-OES (iCAP 6300 duo, Thermo
240 Scientific) in aboveground biomass to calculate plant base cation stocks. Base cations were measured separately
241 in all aboveground biomass parts: stems, leaves, flowers and maize ears.

242 **2.6 Calculation of W_r and potential inorganic CO₂ removal**

243 We use the delta (Δ) symbol to denote the difference relative to unamended control soil. Accordingly, we quantify
244 ΔTA (the change in total alkalinity in the basalt-amended soil relative to the control) based on the difference in base
245 cation concentrations between amended and unamended soils. As basalt only contains cations and no conservative
246 anions, we assume that ΔTA can be quantified from the change in base cation charges (**Equation 1**).

$$247 \quad \Delta TA \approx 2 * (\Delta Ca + \Delta Mg) + \Delta Na + \Delta K - \Delta \text{conservative anions} \quad (1)$$

with $\Delta \text{conservative anions} = 0$

248 The W_r corresponds to the rate of rock dissolution. The W_r can be expressed per element or as moles of alkalinity
249 equivalents (i.e. the sum of base cation charges (**Equation 1**) per amount of rock surface area per unit of time (in
250 mol m⁻² rock s⁻¹). In addition, we calculate a 'potential inorganic CO₂ removal'. We use the same definition for
251 potential inorganic CO₂ removal as in Steinwider et al. (2025). A 'potential inorganic CO₂ removal' can be defined
252 as the maximum amount of inorganic CO₂ that could be removed if all experimentally determined, weathered, soil-
253 retained base cations were to leach from the soil and be completely balanced by carbonate anions. Potential
254 inorganic CO₂ removal was previously 'CDR potential' by Niron et al. (2024)). The concept of CDR potential was
255 first introduced by Phil Renforth (2019) to describe the maximum inorganic CO₂ removal achievable if all base
256 cations within a rock were to completely weather. More recently, Beerling et al., (2024) quantified base cation losses
257 from topsoils using an immobile/mobile tracer approach (see Section 4.2), from which they also derived a measure
258 of CDR potential. To maintain conceptual clarity, we avoid using the term CDR potential for purposes other than its
259 original definition by Renforth (2019). When the term is employed, its meaning should always be explicitly stated.

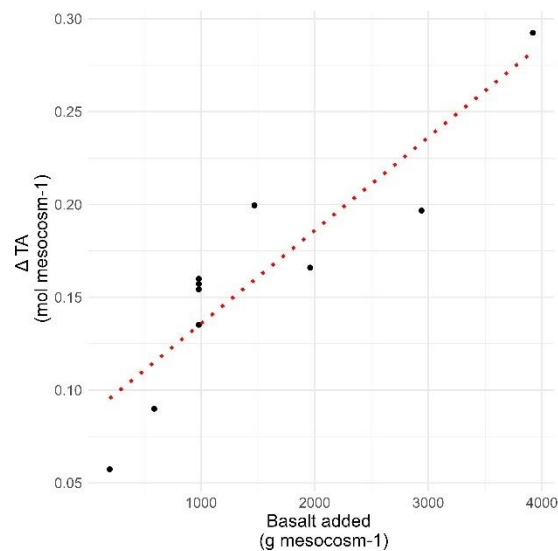
260 To calculate the W_r (from all base cation increases relative to controls in plants, extracted soil fractions and soil
261 water leachates), we sum changes in TA in the following pools: exported soil water (leachates), plants and soil
262 pools. We can express changes in the cation pool of each reservoir as the equivalent W_r required to supply the

263 cations ($W_{leachate}$, W_{plant} and W_{soil}) (**Equation 2**). Conventionally W_{rs} are expressed using a logarithmic scale as
 264 absolute values can vary strongly.

$$265 \quad \text{Log } W_r \left[\frac{\Delta \text{mol TA}}{\text{m}^2 \text{rock.s}} \right] = \text{Log} \left(\frac{\Delta \text{mol TA}_{soil} + \Delta \text{mol TA}_{plant} + \Delta \text{mol TA}_{leachate}}{\text{m}^2 \text{rock.s}} \right) \quad (2)$$

266
 267 We used a gradient of rock applications, where we calculated the slope of the molar change in base cation charges
 268 (expressed as an equivalent “alkalinity” change if these base cations were dissolved in water) with higher rock
 269 amendment (TA slope) (**Figure 3**). We compared logarithmic and linear regression and selected the linear
 270 regression approach, as both approaches had comparable R^2 and Akaike Information Criterion (AIC) values values
 271 and linear slopes ease further data processing (**Fig. S23 and Table S5**). We opted for the linear regression
 272 approach to simplify subsequent calculations. To make our gradient approach more robust, we also calculated the
 273 log W_r for individual application rates in **Fig. S13**.

274



275
 276 **Figure 3:** Illustration of the calculation of TA slope: The alkalinity scavenging by a given pool was plotted in function
 277 of the applied basalt, after which the slope was used to quantify a W_r . This figure is an example regression with
 278 data for the top soil exchangeable pool. All regressions can be found in **Fig. S22**.

279 Then we converted units of the alkalinity slope for increasing rock application (TA slope) per unit of rock mass to
 280 moles of alkalinity per rock surface area and per time (**Equation 3**). Equation 3 was used to calculate $\Delta \text{mol TA m}^{-2}$
 281 rock s^{-1} of base cations in leachates, plants and of every measured soil pool at every soil depth.

$$282 \quad \frac{\Delta \text{mol TA}}{\text{m}^2 \text{rock.s}} = \frac{\text{Scavenged alkalinity (= TA slope)} \left[\frac{\Delta \text{mol TA}}{\text{g rock}} \right]}{\text{Experimental duration [s]} * \text{SSA}_{\text{silicate}} \left[\frac{\text{m}^2 \text{surface area}}{\text{g rock}} \right]} \quad (3)$$

283 $\Delta \text{mol TA m}^{-2} \text{rock s}^{-1}$ was thus quantified per pool, based on the change in base cations in the basalt treatment
 284 compared to the control treatment. For plants, we calculate TA slope through regression of harvested base cations

285 with basalt application. Harvested base cations were calculated as the product of harvested aboveground biomass
 286 and their base cation content. Charge contributions of Na were not included; Na was not quantified at the time of
 287 plant biomass elemental analysis, which may lead to an underestimation of the alkalinity equivalent increase in the
 288 plant pool. However, given that base cation charges in the plant pool were about two orders of magnitude smaller
 289 than in the soil pool, we expect the effect of this omission to be limited. In addition, maize plants aim to actively
 290 increase their K/Na ratio which avoids salt stress, the K content of maize shoots is typically about 2 orders of
 291 magnitude larger than Na (Gao et al., 2016; Suarez & Grieve, 1988). For leachates, TA slope was calculated as
 292 the product of mean cumulative leachate volume and mean leachate TA concentration for each application rate and
 293 regressing them with the applied basalt as dependent variable.

294 Finally, the Wr attributable to the change in cation content of the soil pools (Wr_{soil}) was calculated by summing the
 295 $Wr_{soil_layer_k_pool_j}$ for each pool and depth (**Equation 4**). Here, we sum changes in all pools of the topsoil (0-20 cm)
 296 and lower depths (20-30 cm and 30-50 cm) to obtain an aggregate value for Wr_{soil} . With
 297 $Wr_{soil,layer\ k, fraction\ j}$ calculated as in **Equation 4** (with k = the number of depths and j = the number of considered
 298 soil pools). TA slope at every depth and soil pool was calculated as in **Equation 5**.

$$299 \quad Wr_{soil} = \sum_{k=1}^3 \sum_{j=1}^4 Wr_{soil,layer\ k, pool\ j} \quad (4)$$

$$300 \quad \begin{aligned} & scavenged\ TA\ (TA\ slope) \left[\frac{\Delta mol}{g\ rock} \right] = \\ & \frac{\frac{\mu mol\ TA}{g\ dry\ Soil} (Amended\ Soil) - \frac{\mu mol\ TA}{g\ dry\ Soil} (control\ Soil)}{Application\ rate\ (Amended\ soil) [g\ rock\ m^{-2}\ ground\ area] * 1000} * Bulk\ Density \left[\frac{kg\ dry\ Soil}{m^3\ soil} \right] * thickness\ soil\ layer [m] \end{aligned} \quad (5)$$

301 Equivalents of soil retained base cation charge equivalents per gram of dry soil mixture can be calculated for each
 302 mesocosm by summing the charges from each base cation (**Equation 6**).

$$303 \quad \frac{\mu mol\ TA}{g\ dry\ Soil} = \sum_{j=1}^4 \left(\left(\frac{\mu g\ Ca_{pool\ j}}{g\ dry\ soil} + \frac{\mu g\ Mg_{pool\ j}}{g\ dry\ soil} \right) * \frac{2 mol\ TA}{mol\ cat^{++}} + \left(\frac{\mu g\ Na_{pool\ j}}{g\ dry\ soil} + \frac{\mu g\ K_{pool\ j}}{g\ dry\ soil} \right) * \frac{1 mol\ TA}{mol\ cat^+} \right) \quad (6)$$

304 These individual base cations (e.g. Ca in pool j) are calculated from the difference of cations weathered during the
 305 weathering operation minus the cations initially present in that fraction in the applied feedstock (Power et al., 2025)
 306 (**Equation 7**). For example, some cations can already exchange on the surface or edges of the applied minerals,
 307 so that these cannot be counted as weathered, they will however contribute to CDR once leached.

308 To calculate in-situ Wr , it is thus necessary to correct for the cations that had already been weathered from primary
 309 minerals at the time of silicate amendment. This correction is currently not being done in EW literature yet. As basalt
 310 is only added to the top soil and not deeper, this correction is only done for the 0-20 cm soil layer here.

$$\left(\frac{\mu\text{g element}_{i\text{pool}j}}{\text{g dry soil}} \right)_{\text{Post weathering, soil mixture}} = \left(\frac{\mu\text{g element}_{i\text{pool}j}}{\text{g dry soil}} \right)_{\text{added with feedstock initially}} \quad (7)$$

The mass of a specific element (i) in each of the four (j) soil pools (in $\mu\text{g element/g soil}$) is calculated using **Equation 8**.

$$\left(\frac{\mu\text{g element}_{i\text{pool}j}}{\text{g dry soil}} \right)_{\text{Post weathering, soil mixture}} = \frac{\text{concentration element}_i \text{ in pool}_j \left[\frac{\text{mg}}{\text{L}} \right] * \text{Volume extract}_j [\text{mL}]}{\text{mass of solid extracted} [\text{g}]} \quad (8)$$

The initial addition of element i to pool j is calculated as in **Equation 9**.

$$\left(\frac{\mu\text{g element}_{i\text{pool}j}}{\text{g dry soil}} \right)_{\text{added with feedstock initially}} = \frac{\mu\text{g element}_i \text{ pool}_j}{\text{g silicate}} * \frac{\text{Application rate} \left[\frac{\text{g silicate}}{\text{m}^2} \right]}{\text{Bulk density} \left[\text{g dry soil} \frac{\text{m}^3}{\text{m}^3} \right] * \text{depth of soil amendment} [\text{m}]} \quad (9)$$

According to the charge balance (**Reaction 1-3**) during mineral dissolution, 1 mol HCO_3^- mol⁻¹ TA is generated (and thus 1 mol of CO_2 is sequestered). We define a factor η , that is equal to the ratio of HCO_3^- per mol of generated TA. According to charge balance, $\eta=1$. A more conservative approach is to assume that all this generated alkalinity will be exported to the ocean, after which chemical equilibrium degasses a portion of the alkalinity ($\eta = 0.7$ mol CO_2 mol⁻¹ TA, assumed for oceans) (Renforth et al., 2012; Renforth et al., 2019; Renforth & Henderson, 2017). According to Renforth et al. (2019), the ocean alkalinization efficiency η ranged between 0.7 and 0.85. This η parameter is relatively uncertain given that model studies indicate that η can range between 0.65 and 0.8 mol CO_2 mol TA⁻¹ (see section S6 in the supplement of (Kowalczyk et al., 2024)). Alternatively, we can assume that all base cations will form solid carbonates in soils or rivers. In this case $\eta=0.5$ mol CO_2 mol⁻¹ TA (**Reaction 4**). In **Table 4**, we calculated potential inorganic CO_2 removals assuming conservative values of $\eta=0.5$ (carbonate precipitation scenario) and $\eta=1$ (the highest possible η without any downstream DIC losses).

While calculating W_r , cations added with the rock feedstock that were already weathered were subtracted as in **Equation 7**, yet these cations are not subtracted to calculate the potential inorganic CO_2 removal as base cations in weathered fractions of the rock feedstock can also leach to soil water whereby HCO_3^- is generated. Last, base cation changes in the plant pool were excluded from the potential inorganic CO_2 removal pool here, as a conservative approach we assume that base cations in plants will not reach the ocean. The latter assumption had a negligible impact on the potential inorganic CO_2 removal estimate (**Table 4**).

$$\text{potential inorganic CO}_2 \text{ removal} \left[\frac{\text{kg CO}_2}{\text{t rock}} \right] = \text{Scavenged TA} \left[\frac{\text{mol TA}}{\text{g rock}} \right] * \frac{\eta \text{ mol CO}_2}{\text{mol TA}} * \frac{44\text{g CO}_2}{\text{mol CO}_2} * 1000 \quad (10)$$

with $\left(\frac{\mu\text{g element}_{i\text{pool}j}}{\text{g dry soil}} \right)_{\text{added with feedstock initially}} = 0$

335

336 2.7 Calculation of the carbonate saturation indices (SIc) using Phreeqc

337 To assess whether carbonate precipitation was theoretically possible during this experiment, we calculated SIc for
 338 dolomite and calcite. For Mg and Ca the SIc as the logarithm of the ion activity product and the solubility product
 339 constant if dolomite and calcite ($\text{SIc} = \log \text{IAP/K}$). Minerals have the potential to precipitate when a log SIc >0 is

340 reached although substantial oversaturation of calcite ($\log \text{SIc} > 1$) without calcite formation is possible in rivers due
341 to ion inhibition, e.g. by phosphate (Morse et al., 2007; Zhang et al., 2022). Likewise, they are in equilibrium at a
342 $\log \text{SIc} = 0$ and dissolve if $\log \text{SIc} < 0$. The R phreeqc package was used and the phreeqc.dat database was used.
343 As an input, the experimental pore water (10 cm depth) composition of Mg and Ca was entered, as well as measured
344 pH and TA. Daily SIc values were calculated by feeding unique combinations of Mg, Ca, pH and into the PHREEQC
345 solution function for each day.

346 **2.8 Data analysis**

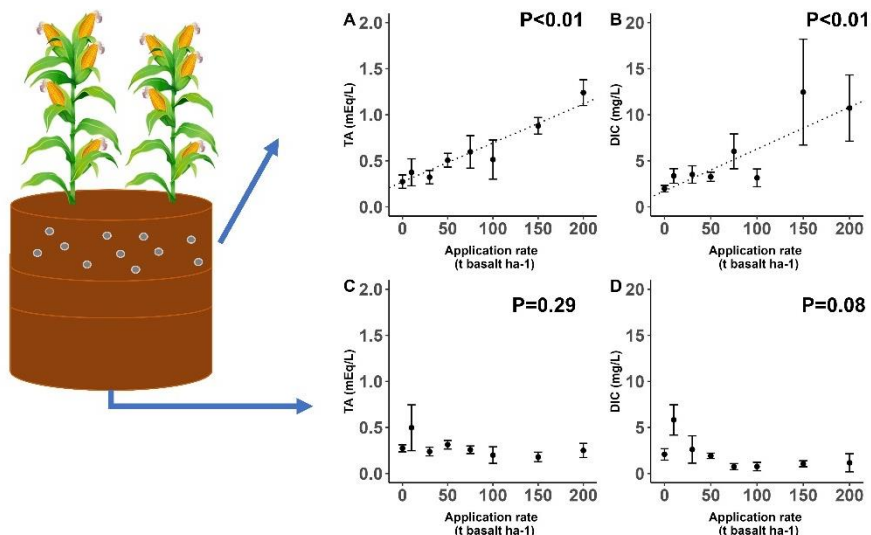
348 For SIC and elemental stocks in plant biomass, soil pools and soil water export, a linear regression with basalt
349 application rate as a dependent variable was performed to test for a basalt effect. For measurements that were
350 repeated in time (pore water and leachate DIC and DOC compositions), a linear mixed model was used with basalt
351 and time as fixed factors and mesocosm as a random factor using the lme4 R package (version 1.1-33). For
352 measurements repeated in time, we assessed basalt x time interaction effects and discarded these if not significant.
353 All analyses were executed in R version 4.3.2. As an additional sensitivity analysis for the determination of W_r using
354 the slope of application rates approach described in the main text, we quantified W_r also for individual application
355 rates in **Fig. S13**.

356 To propagate uncertainty between basalt and controls in Figure 5, averages and standard errors for every replicated
357 application rate (0 or 50 t/ha) were determined. The average from the 50 t ha⁻¹ was subtracted with the average
358 from the control soil and $se = \sqrt{(se_control^2 + se_basalt^2)}$. For non-replicated application rates (10,30, 75,100,
359 150 and 200 t/ha, $se=0$) the measurement was subtracted from the control soil average and errors were also
360 propagated with $se = \sqrt{(se_control^2 + se_basalt^2)}$.

361 **3 Results**

362 Basalt amendment significantly increased DIC and TA in the top soil water (**Figure 4**). TA in soil water correlated
363 positively with DIC ($R^2 = 0.68$, $p < 0.01$, **Fig. S10**). TA was thus generated in the basalt amended soil layer, yet we
364 did not observe DIC or TA increases with higher basalt application rates in water exported from the soil column at
365 60 cm depth (**Figure 4**). Temporal dynamics show that DIC in top soil pore water gradually increased in time with
366 higher basalt amendment, while DOC decreased in time with more basalt (**Fig. S7 and Table S4**).

368



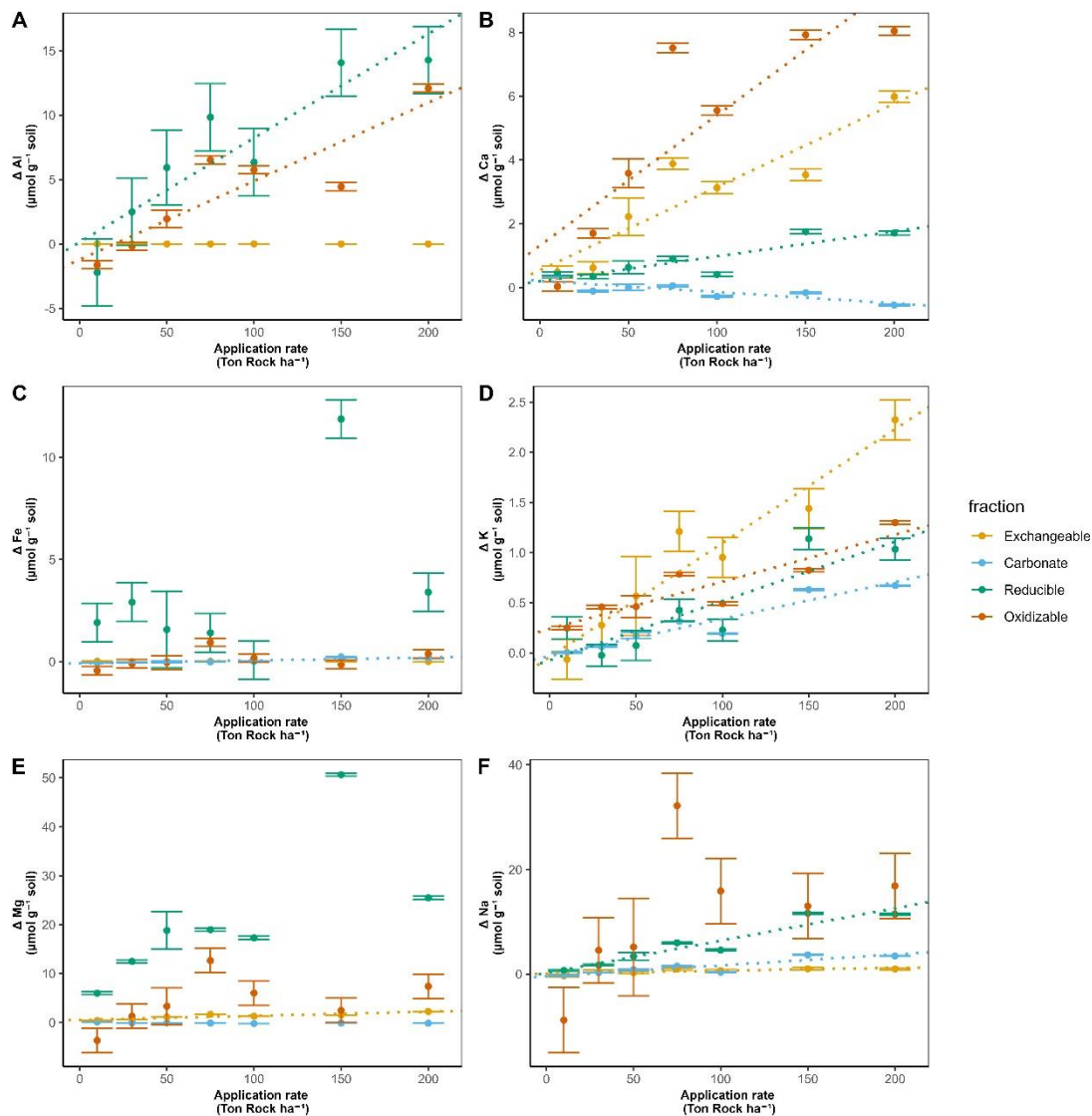
369
 370 **Figure 4:** Top soil (0-20 cm) pore water (A) TA and (B) DIC concentrations. Export water (50 cm depth) (C) TA and
 371 (D) DIC concentrations. Values represent average concentrations +- standard error across all sampling occasions
 372 over the 101 day duration experiment (n= 125, 44, 95 and 60 for pore water DIC, TA and leachate DIC and TA
 373 concentrations respectively). Significant trends are indicated with a dotted regression line. Raw data for TA and DIC
 374 in function of time is visualized in **Fig. S7(B,D)** and **Fig. S9(B,D)**.
 375

376 Overall, base cations were mostly retained in the top soil, where Ca significantly increased in the exchangeable,
 377 reducible and oxidizable pools with higher basalt addition. Only in the carbonate soil pool, Ca (and also Mg)
 378 significantly decreased with more basalt (**Figure 5 and 6**). With higher rock amendment, Mg accumulated in the
 379 top soil exchangeable pool ($p < 0.01$). The Mg accumulation in the reducible pool was higher compared to the
 380 exchangeable, but the slope was borderline significant for the reducible pool ($p = 0.07$) due to higher variability in Mg
 381 concentrations with increasing basalt amendment.

382 Changes in Na followed similar patterns as Mg, as also significantly more Na exchanged in top soil ($p = 0.02$) and a
 383 larger signal of reducible Na was found ($p < 0.01$). In contrast with divalent cations, monovalent cations increased in
 384 the carbonate fraction if basalt increased (**Figure 5 and 6**). With more basalt, Al is being found in association with
 385 the oxidizable and reducible fraction. Si increased only significantly in the oxidizable pool ($p = 0.04$) (**Fig. S15**).
 386 Increases in oxidizable Si, Ca, Al with higher basalt addition suggest the formation of mineral-associated organic
 387 matter.

388 In the soil layer just below the soil-basalt mixture (20-30 cm), the cations did not increase significantly in any of the
 389 measured soil pools and oxidizable Na, Fe and Mg decreased significantly (**Fig. S11, Figure 6**). We did not observe
 390 significant changes in any element with higher basalt amendment in the 30-50 cm soil layer (**Fig. S12**).

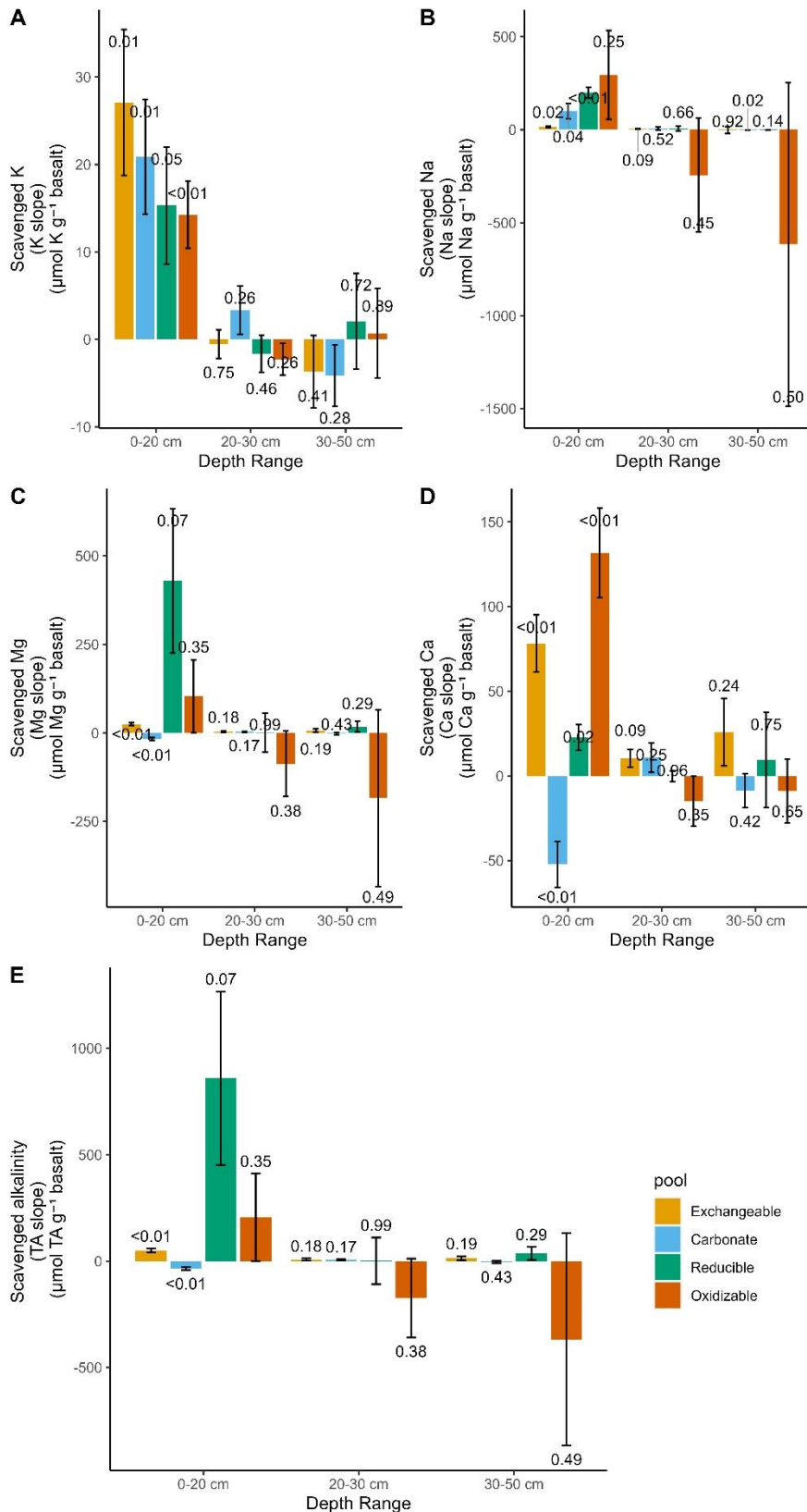
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 395 **Figure 5:** Change in top soil (0-20 cm) elements relative to the control soil (corrected as in **Equation 7**), 101 days
 396 after basalt amendment, as a function of basalt application rate for (A) Al (B) Ca (C) Fe (D) K (E) Mg and (F) Na for
 397 four different soil pools. Dots and error bars represent averages and standard errors. For basalt application rates
 398 other than 50 t ha⁻¹, error bars correspond to those of the control soils, as these basalt treatments were not
 399 replicated and the data are shown as control-normalized results. Significant effects (p<0.05) of basalt application
 400 rate on cation concentrations are indicated by dotted linear regression lines. Measurements were repeated on at
 401 least four samples per fraction for the control soils (N ≥4 for each fraction) and N=4 for 50 t ha⁻¹ treatment (fewer
 402 than 5 replicates were available due to technical issues). Note that y-axes scales differ among subplots to better
 403 visualize small changes for some elements. Unnormalized (raw) data are presented in **Fig. S19-21**.
 404

405 From all significant element changes in soil pools, we calculate that 8.4%, 52.1% and 9.4% Of basalt Na, K and Ca
 406 were weathered while we do not observe an increase in Mg if we only consider significant (p<0.05) slopes. If we
 407 consider all (also p>0.05) regression slopes, the estimates become 48.0%, 10.6%, 9.35% for K, Ca and Mg, while
 408 Na did not increase in this approach (mass balance per element, see **Fig. S23**).

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Figure 6: Equivalent alkalinity uptake 101 days after basalt amendment in different soil pools and depths. P-values of linear regressions are shown above and below bar plots of positive and negative changes respectively. Error bars represent the standard error that was derived from linear regression. Underlying regressions for slopes of TA scavenged by each pool and depth can be found in **Fig. S22**. Base cation changes in top soil pore water are not included in this figure as we only include charge equivalent adsorbed by soil pools here, yet base cations in top soil pore water were negligible (see **Fig. S24**).

419 Base cations were not only scavenged by soils, but also by plants. Although two orders of magnitude smaller than
 420 in soil pools, TA scavenging by plants was higher than soil water exported TA and increased significantly with larger
 421 basalt amendment ($p < 0.01$) (**Figure 7**). The increase in base cation charges in plants was attributed to K (81%),
 422 Ca (11%) and Mg (8%).
 423

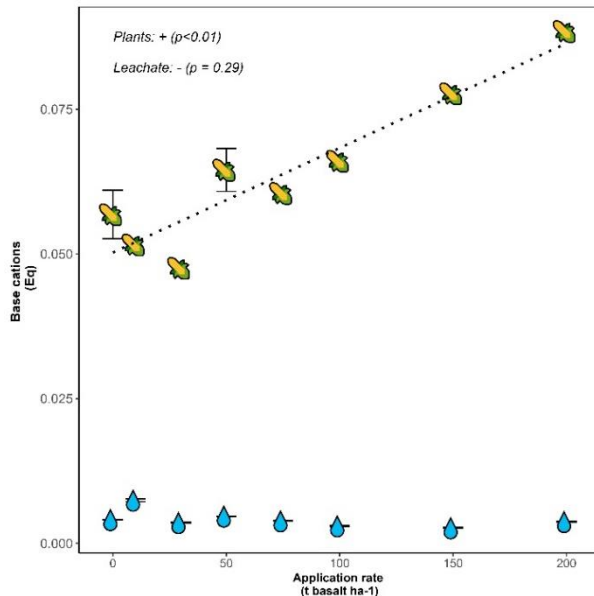


Figure 7: Moles of base cation charge equivalent (Eq) per mesocosm after 101 days retained in maize plants (stems, leaves and maize ears), indicated with maize fruit symbols, and flushed with leaching water, indicated with droplets. Error bars represent averages and standard errors ($n=5$ for both control and 50 t ha^{-1} basalt replicates for plant measurements). For leachate TA, a total of 60 measurements were done in total at different dates. Na was not analyzed in plants and is thus not included in the harvested base cations. Errors on leachate TA were small and appear as horizontal lines within the droplets. Raw data for leachate TA measurements can be found in **Fig S9D**.

446
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 448
 449 Converting the base cations to moles of equivalent TA and considering only the exchangeable pool as only soil
 450 cation reservoir we derive a log weathering rate of $-12.13 \pm 0.34 \text{ mol TA m}^{-2} \text{ rock s}^{-1}$ (**Table 4**). When we consider
 451 also the decrease in base cation equivalents in the carbonate pool, the mean estimate decreases to -12.23 mol TA
 452 $\text{m}^{-2} \text{ rock s}^{-1}$, translating into mean estimated potential inorganic CO_2 removals of $0.36\text{-}0.5 \text{ kg CO}_2 \text{ t}^{-1} \text{ basalt}$
 453 (assuming $\eta=0.5\text{-}0.7$). If we include all soil pools and non significant regressions the estimates becomes one order
 454 of magnitude higher, yet with substantial uncertainty.

455 **Table 4:** Overview of the Wr and potential inorganic CO_2 removal that can be quantified from changes in base
 456 cations in specific soil pools. Rows where (scavenged) TA increased significantly with increasing basalt amendment
 457 are indicated in bold.

Soil Pool	Depth	Reservoir	Log Wr (Log mol TA/m ² basalt /s)	CDR Potential* (kg CO ₂ /ton basalt) ($\eta = 0.5$)	CDR Potential* (kg CO ₂ /ton basalt) ($\eta = 1$)
/	/	Plant*	-12.93 ± 0.07	/	/
/	/	Leachate*	$Wr < 0$	/	/
Exchangeable	0-20cm	soil	-12.20 ± 0.41	1.10 ± 0.04	2.21 ± 0.07
Carbonate	0-20cm	soil	$Wr < 0$	-0.75 ± 0.03	-1.50 ± 0.06
Reducible	0-20cm	soil	-10.96 ± 0.21	18.91 ± 1.76	37.82 ± 3.51
Oxidizable	0-20cm	soil	-11.58 ± 0.43	4.53 ± 0.89	9.06 ± 1.77
Exchangeable	20-30cm	soil	-13.02 ± 0.29	0.17 ± 0.02	0.34 ± 0.04
Carbonate	20-30cm	soil	-13.17 ± 0.29	0.12 ± 0.02	0.23 ± 0.03
Reducible	20-30cm	soil	-	-	-
Oxidizable	20-30cm	soil	13.17 ± 50.43	0.02 ± 0.47	0.04 ± 0.95
Exchangeable	30-50cm	soil	$Wr < 0$	-3.82 ± 0.80	-7.56 ± 1.60
Carbonate	30-50cm	soil	-12.76 ± 0.30	0.30 ± 0.04	0.61 ± 0.08
Reducible	30-50cm	soil	$Wr < 0$	-0.10 ± 0.02	-0.21 ± 0.05
Reducible	30-50cm	soil	-12.34 ± 1.15	0.79 ± 0.13	1.58 ± 0.27

Oxidizable	30-50cm	soil	Wr<0	-8.10±2.15	-16.19±4.31
Exchangeable	0-20	Exchangeable+plant+leachate	-12.13±0.34	1.30±0.04	2.21±0.07
Exchangeable + Carbonate	0-20	Exchangeable+plant+leachate+ carbonate	-12.23±1.05	0.36±0.05	0.71±0.10
All soil pools	All	All soil pools + plant + leachate	-11.11±2.70**	13.17±3.07	26.33±6.13

458 *For leachates (which represents realized CDR) and also for plants there is no potential inorganic CO₂ removal in
459 this approach. ** Abs (Wr/standard error (Wr)*LN(10)) was used to propagate the error using the log10
460 transformation, resulting in substantial uncertainty for the Wr estimate of all pools.

461 4. Discussion

462 4.1 Weathering rates and CO₂ removal

463 EW is typically considered as a durable CDR pathway that removes CO₂ from the atmosphere by producing DIC
464 that is either transported to the ocean (Strefler et al., 2018) or precipitates as carbonates in the soil (Manning et al.,
465 2013). Here, we observe a clear weathering signal (a TA and DIC increase) in top soil pore water (**Figure 4**). These
466 TA and DIC increases in the pore water of amended top soil are consistent with recent findings (Holzer et al., 2023;
467 McDermott et al., 2024; Vienne et al., 2024). Increased DIC in basalt soils relative to controls may result from
468 enhanced plant root respiration or DIC exudation or from mineral weathering; our dataset does not allow these
469 effects to be separated. DIC did however not leach from our soil columns within this experimental timeframe of 101
470 days.

471 Absence of substantial DIC leaching is in line with other short-term recent studies (Amann et al., 2020; Larkin et al.,
472 2022; Niron et al., 2024; Vienne et al., 2024). For example, the log Wr of approximately -13 mol TA m² s⁻¹ quantified
473 from DIC export after 1 year in a mesocosm trial with 220 ton ha⁻¹ olivine-rich rock (Amann et al., 2020) was about
474 three orders of magnitude lower than what would be expected from lab-scale weathering studies (roughly -10 mol
475 TA m² s⁻¹, (Palandri & Kharaka, 2004)). Vienne et al. (2024), amended soils with 100 ton basalt ha⁻¹ and quantified
476 a CDR from exported TA that was in the same order of magnitude as in the work of Amann et al. (2020). Although
477 the studies of Amann et al. (2020), Vienne et al. (2024) were relatively short (<= 1 year) and used a relatively low
478 water infiltration flux, also a longer (3 year duration) catchment-scale study in Malaysian oil palm plantations with
479 high annual rainfall (>2000 mm year⁻¹) detected no significant increase in TA leaching in the catchments (Larkin et
480 al., 2022). There may thus be a substantial delay for DIC leaching.

481 A DIC leaching delay can have multiple causes (**Figure 1**); A first possibility is pedogenic carbonate formation.
482 We observe that solid carbonates did not increase in our experiment, in contrast, SIC decreased in time. PHREEQC
483 calculations for our experiment suggest that dolomite and calcite were undersaturated, so that carbonate dissolution
484 was possible (**Fig. S17**). Saturation states are expected to be low in our experiment because control soil was
485 undersaturated and dissolved base cations accumulated in other soil pools than the carbonate pool (**Figure 5 and**
486 **6**). A decrease in SIC is in contrast with substantial SIC increases found after wollastonite rock amendment (Haque

491 et al., 2019, 2020). The observed SIC increase in the latter field study may be partly attributed to residual carbonates
492 from prior liming activities instead of new carbonate formation related to silicate weathering (Haque et al., 2020).
493 Thus, not all measured SIC may reflect new carbonate formation in the study of Haque et al. (2020). For short-term
494 basalt studies, using elemental C analysis, also no significant changes in SIC could be detected previously (Kelland
495 et al., 2020; Vienne et al., 2022, 2024). In contrast, in the study of Larkin et al. (2022), a relatively small SIC increase
496 was detected in amended soils, using carbonate pool extractions.

497

498 While TA was not exported or taken up by soil carbonates here and plant base cation losses were minor (**Table 4**)
499 it was retained in top soil where the exchangeable and reducible pools reduced solute TA. We expect cations to be
500 primarily associated with Fe- and Mn-(oxyhydr)oxides in the reducible pool and with organic matter in the oxidizable
501 pool, as supported by literature (Tessier et al., 1979); However, the extraction chemicals of this sequential extraction
502 scheme (hydroxylamine and H₂O₂) are known to have a limited specificity and may have also partially targeted
503 other mineral phases (such as clays) (Ryan et al., 2008), which could explain the elevated Si observed in the topsoil
504 pools (**Fig. S15**). In addition, the observed increase of aluminum in association with the reducible soil fraction
505 indicate the formation of secondary minerals. While we cannot pinpoint the exact Mg-phases formed in our soils,
506 our results do demonstrate substantial base cation retention in the soil and show that there can be more base cation
507 losses to soils than to the exchangeable pool alone.

508

509 Our estimate of log W_r, derived solely from significant increases in TA uptake at higher basalt amendment rates,
510 was approximately -12 mol TA m⁻² s⁻¹. This estimate reflects changes in the exchangeable and carbonate soil
511 pools, plant uptake, and leachate composition. Notably, this value aligns with previous studies that estimated log W_r
512 values between -12 and -11 based on base cation depletion from the exchangeable pool alone (Kelland et al.,
513 2020; Reershemius et al., 2023; te Pas et al., 2023), as summarized in Vienne et al., (2024). Also in a batch leaching
514 experiment with 1mM CaCl₂ (designed to mimic soil solutions) the quantified log W_r of basalt was found to be -11
515 (Van Der Bauwhede et al., 2024).

516

517 Estimates from Buckingham et al. (2022), based only on leachates, gave a much lower log W_r of -15, partly due to
518 low water infiltration rates. With a high infiltration flux (8000 mm/year), Amann et al. (2022) estimated log W_r
519 between -12.5 and -13.5 from basalt leachates. This highlights the importance of including scavenged alkalinity to
520 determine W_r in soils. When we also include non-significant regression slopes we derive a mean log W_r estimate
521 with substantial uncertainty (-11.11±2.70) mol TA m⁻² s⁻¹. From individual application rates, we quantify log W_r
522 ranging between -11 and -10 (**Fig. S13**); These values are comparable to those observed in soil-free, laboratory-
523 scale basalt dissolution experiments conducted at circumneutral pH (Brantley et al., 2008; Gislason & Oelkers,
524 2003). They also approximate the dissolution rates of key mineralogical components in basalt (such as plagioclases

525 (between -12 and -9 for Na and Ca endmembers respectively) and augite (-11.97)) under room temperature and
526 neutral pH conditions (Gudbrandsson et al., 2011; Hermanska et al., 2022; Palandri & Kharaka, 2004a).

527

528 Although this and other experiments show relatively consistent weathering rates from exchangeable base cations
529 (comparable to those observed in lab-scale studies) we emphasize that, unlike laboratory conditions where base
530 cations remain far from equilibrium in excess water, soils experience solid-phase cation scavenging, which
531 promotes DIC degassing (**Figure 1**). From the sum of significant TA slopes we calculate a relatively low potential
532 inorganic CO₂ removal, equalling to approximately 0.36-0.71 kg CO₂ ton⁻¹ basalt or 0.018-0.036 tCO₂ ha⁻¹ for a
533 basalt application rate of 50 t ha⁻¹ (**Table 4**). Also the highest possible potential inorganic CO₂ removal realized
534 within this experimental timeframe is modest; Including also non-significant TA increases and assuming $\eta = 1$, the
535 potential inorganic CO₂ removal is quantified to be 26.33±6.13 kg CO₂ ton⁻¹ basalt (**Table 4**). We emphasize that
536 a potential inorganic CO₂ removal is a maximum inorganic CO₂ removal that can be realized with the delivered
537 amount of base cation weathering as strong acids associated with fertilizers (such as nitric acid and sulphuric acid),
538 or organic acids and not carbonic acid may have initially weathered silicate rock which does not lead to a CDR
539 within the soil system (McDermott et al., 2024; Taylor et al., 2021). Moreover, life-cycle emissions associated with
540 mining, grinding and transporting rock are typically of the same order of magnitude as our relatively low potential
541 inorganic CO₂ removal (Lefebvre et al., 2019).

542

543 Furthermore, for climate change mitigation, not only the amount of potential inorganic CO₂ removal is important,
544 but also the timescale at which this CDR is realized (Kanzaki et al., 2025). A mass balance of base cations indicates
545 that exported TA was negligible compared to base cation charges that were retained in the soil over the timeframe
546 of our experiment (101 days) (**Table 4**). As long as TA is retained in soil pools, inorganic CO₂ removal through DIC
547 export is delayed as equivalent amounts of protons have then been released into the soil water to maintain charge
548 balance (**Figure 1**). Realization of this delayed inorganic CO₂ removal depends on liberation of base cations from
549 these soil pools and their transport out of the soil, charge-balanced by HCO₃⁻. This export may take decades or
550 longer, depending on the circumstances (Kanzaki et al., 2025).

551

552 The realization of CDR may be further delayed through the formation of base cation bearing clay minerals. Clay
553 formation has previously been suggested for EW application based on changes in soil water Ge/Si ratios and Si
554 isotopes (Vienne et al., 2024) and also based on Li isotope measurements (Pogge von Strandmann et al., 2022).
555 These measurements indicated basalt induced clay formation, but it remains unclear what type of clays were formed
556 and hence what the effect on inorganic CO₂ removal may be. In the best case for the inorganic CO₂ removal time
557 lag, the formed clays are 1:1 phyllosilicates such as kaolinite that do not sequester base cations. In this case, DIC
558 leaching is only retarded by base cation exchange. Worst case for the inorganic CO₂ removal time lag, the formed

559 secondary minerals bear substantial amounts of base cations such as chlorite or chrysotile. These clays exhibit a
560 log W_r between -12 and -12.5 at neutral pH (Palandri & Kharaka, 2004), so that dissolution within decadal
561 timescales is unlikely (Bullock et al., 2022).

562

563 Although unfavourable for inorganic CO₂ removal, if base cation bearing secondary clay minerals would form, they
564 can increase SOC (Georgiou et al., 2022; Heckman et al., 2022; Steinwider et al., 2025). Georgiou et al. (2022)
565 refer to base-cation bearing clays (e.g. smectitic or illitic clays) as ‘high-activity minerals’ due to their higher SOM
566 stabilization capacity compared to secondary minerals that do not contain base cations (i.e., ‘low-activity minerals’,
567 with a lower CEC such as kaolinite). Both high- and low-activity minerals can adsorb DOC and form mineral-
568 associated organic matter-C (MAOM-C), which is believed to have a relatively high permanence (decades-
569 centuries) in soils (Lavalée et al., 2020). Besides mineral surface however, plant inputs can also limit SOC accrual.
570 In the latter case, SOC stocks can only increase if belowground plant C inputs increase, which could follow from
571 increases in exchangeable bases or pH (Haque et al., 2019; Shamshuddin et al., 2011). Nonetheless, increases in
572 decomposition can also stimulate SOC losses if rock dust increases soil pH (Klemme et al., 2022).

573

574 **4.2 Implications for monitoring inorganic CO₂ removal**

575

576 Different base cation monitoring strategies are possible. A first option is to quantify TA in soil water (Clarkson et
577 al., 2024). A disadvantage is however that soil water samples have to be sampled across the soil depth.
578 Alternatively, TA could be only monitored in top soil, yet then uncertain TA leaching models must be used (Kanzaki
579 et al., 2025). To calibrate TA leaching models, soil measurements in depth profiles could be used.

580

581 A first soil measurement approach is a total cation accounting approach, which quantifies the loss of base cations
582 from top soils (te Pas et al., 2025). However, this approach only focuses on the top soil and fails to account for
583 physical cation transport from top soils due to erosion or vertical feedstock transport via infiltration or bioturbation.
584 Alternatively, in a mobile/immobile tracer element approach (often named ‘TiCat’ by the EW community), cation
585 losses from amended top soils are quantified along with immobile tracers, which can account for cation losses
586 through bioturbation or erosion (Reershemius et al., 2023). Nonetheless the disadvantage of TiCat is that it does
587 not track potential TA scavenging (e.g. by organic matter or clays) at larger depth. Our potential inorganic CO₂
588 removal estimate will thus differ from a potential inorganic CO₂ removal estimate quantified using a TiCat approach.

589

590 Alternatively, entire depth profiles could be analyzed to spatially calibrate TA leaching models. Analysis of the
591 exchangeable soil pool is already a well-established method in EW research (Beerling et al., 2024; Reershemius et
592 al., 2023; Reynaert et al., 2023; Vienne et al., 2024). Adding also the carbonate, reducible and oxidizable soil pools
593 to the analysis could make base cation mass balancing more complete. These protocols could calibrate predictive

594 TA leaching models spatially. In addition there is an opportunity to quantify SOC and MAOM-C changes in the same
595 samples, which have recently gained traction in EW research due to their role in stabilizing SOM (Buss et al., 2024;
596 Sokol et al., 2024; Xu et al., 2024). Integration of these measurements can provide more accurate estimates of the
597 climate impact of EW, but should take into account the difference in permanence of inorganic and organic carbon
598 stocks.

599
600 However, this monitoring approach involves complexities such as feedstock correction, leaching solution strength
601 and soil heterogeneity. Although correcting for pre-weathered elements was crucial in this study, it assumes perfect
602 mixing based on a silicate-to-soil ratio. This correction was particularly significant for carbonate and reducible soil
603 pools, where for some base cations, over half of the cation increase with basalt amendment originated from
604 feedstock addition and not from weathering (**Fig. S18**). An alternative approach could involve creating time series
605 from sequential extraction data and quantifying base cation changes based on the change in time between multiple
606 measurements taken after rock amendment.

607
608 As discussed in the previous section, another key challenge is that the fate of base cations may remain uncertain
609 if strongly bound crystalline organo-minerals (see Lopez-Sangil & Rovira, 2013) form that are unextractable by the
610 Tessier scheme. Such processes may have contributed to the observed decrease in oxidizable elements at larger
611 depth, although this could also be an artefact of the applied extraction procedure. Pogge von Strandmann et al.
612 (2022) proposed substituting the H₂O₂ leaching step of the Tessier scheme with a dilute HCl leach, which is thought
613 to extract clays as well. Alternatively, post-extraction analysis of residual solids using techniques such as XRD or
614 QEMSCAN may be necessary to rigorously assess changes in rock mineralogy (Mason et al., 2022). Although deep
615 soil core sampling and extensive mineralogical analysis are resource-intensive and not feasible for large-scale
616 application, this monitoring strategy could be valuable during the initial adoption of EW in targeted 'measure-all'
617 experiments, as reliable TA leaching models require extensive calibration.

618

619 **5. Conclusions**

620 This study presents a detailed examination of EW and its effectiveness as a climate mitigation technique, revealing
621 both its potentials and limitations. A novel aspect of this work is the in-depth investigation of entire soil profiles for
622 base cations in different soil fractions, paired with soil water TA monitoring. We highlight the value of sequential
623 extractions as a method for monitoring base cations throughout soil profiles for calibrating TA leaching models.

624 Our findings indicate that basalt-based enhanced weathering may not immediately lead to the inorganic CO₂
625 removal previously anticipated in projections and IPCC reports (Babiker et al., 2022; Minx et al., 2018). We observed
626 rock weathering without inorganic CO₂ removal; despite the absence of DIC leaching or carbonate precipitation,
627 exchangeable bases increased with higher basalt amendments, demonstrating that rock weathering occurred.

628 Additionally, we observed a borderline significant but substantial increase in base cations in the reducible topsoil
629 pool with greater basalt application, which may further suppress TA leaching.

630 As base cation exchange increased with higher basalt amendments, we infer that greater application rates can
631 further delay the release of DIC from soil minerals to surface waters. However, in practice, EW is typically applied
632 at application rates below 30 t ha⁻¹. These lower, more practical rates may also enhance effectiveness of inorganic
633 CO₂ removal by reducing lag times for DIC release. It remains unclear if clays were formed here and whether EW
634 can deliver CDR within the urgent decadal timeframe needed to mitigate climate change. Despite its limitations for
635 short-term inorganic CO₂ removal, the generated secondary minerals and increased CEC could enhance plant
636 productivity and SOC retention in soils, contributing to long-term soil health, fertility, and potentially carbon
637 sequestration beyond inorganic pathways.

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647 Author contribution

648 AV: research conceptualization, data gathering, development methodology, data analysis and writing. PF:
649 conceptualized sequential extraction methodology, writing and discussion. JR: research conceptualization, data
650 gathering. TJS: writing and discussion. TR: writing and discussion. RP: data gathering, rock characterization and
651 writing. JH: writing and discussion. HN: development extraction methodology, writing and discussion. MPE:
652 elemental C measurements and proofreading. LS: writing, development methodology and discussion. LB: writing
653 and discussion. SV: supervising research, conceptualization, writing and discussion.

654 Data and code availability

655 Data and code used in this manuscript are freely available at: <https://zenodo.org/records/15129984>

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