



Organic matter-mediated leaching of alkalinity in limed acid soils is affected by dissolved organic carbon adsorption and soil structure

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Abstract. Subsurface soil acidity severely limits crop growth and is challenging to adjust by surface liming. There have been several proposals for subsurface liming using the combination of lime and an organic amendment, as organic anions may migrate deeper in acid subsoil than carbonates. This study aimed to identify mechanisms of subsurface liming, postulating that it is hindered by dissolved organic carbon (DOC) adsorption but enhanced in structured compared to sieved soils due to preferential flow in macropores. Column leaching experiments were set up using three sieved acid soils with contrasting properties, of which one was additionally sampled as undisturbed soil cores. The upper layer of each soil was treated with lime, compost, or a combination of both, in addition to an untreated control and columns were leached with artificial rainwater. Deeper subsurface liming in the lime+compost treatment than in the lime treatment was detected in only one of the three soils. The effect of compost on the migration of alkalinity was explained by differences in DOC sorption among soils, the lowest sorption leading to deepest subsurface liming. Imaging of in situ pH using a planar optode showed evidence of preferential alkalinity flow in the structured soil, however destructive sampling of bulk soil layers did not confirm this. We conclude that combining lime with an organic amendment can effectively ameliorate subsoil acidity but this requires weakly DOC adsorbing subsoils. The role of soil structure on this process needs to be corroborated with plant responses to identify benefits of liming the macropores.

20 **1 Introduction**

About half of the world's potential arable land consists of acid soils ($\text{pH} \leq 5.5$), making soil acidity one of the most important agricultural constraints worldwide (George et al., 2012; Kochian et al., 2004; von Uexküll & Mutert, 1995). Toxic levels of aluminum (Al) and manganese (Mn), and deficiencies in phosphorus (P), calcium (Ca) and magnesium (Mg) severely affect crop yield in acid soils. Restricted root growth, particularly caused by Al, leads to even lower nutrient uptake and increased water stress (Marschner, 1991; Tang et al., 2013). Topsoil acidity is typically amended by applying lime or dolomite (CaCO_3 or $\text{CaMg}(\text{CO}_3)_2$). However, surface application of lime is often inefficient in alleviating subsoil acidity (below 0.1 m) due to the slow downward movement of lime in soil (Conyers & Scott, 1989; Sumner et al., 1986; Tang et al., 2013). The factors explaining the low mobility of lime are its low solubility, the fast consumption of OH^- or HCO_3^- produced during the liming reaction, and the lack of an accompanying anion for the downward transport of Ca^{2+} (Liu & Hue, 2001).



30 For example, Azam & Gazey (2020) found that the subsurface soil pH increase remained limited to 0.049 pH units per year
to a maximum depth of 0.20 m. This was only achieved after repeated surface applications of unrealistically high doses of
lime (up to a total of 8.5 tons ha⁻¹) over 10-24 years. Similarly, Li et al. (2019) showed in an 18-year field trial that pH
increase in the soil profile remained confined to the top 0.3 m depth when pH was continuously maintained above 5.5 in the
top 0.1 m. Consequently, the adverse effects of soil acidity often persist in the root zone, which becomes particularly
35 important when moisture is depleted in the topsoil at the end of the growing season, and plants need to rely on water and
nutrients from the more acid subsoil (Tang et al., 2003, 2013). Therefore, many studies have searched for ways to alleviate
subsoil acidity.

Limited evidence suggests that combining lime with an organic amendment (OA) might enhance alkalinity movement down
the soil profile (Butterly et al., 2021; Lauricella et al., 2021; Liu & Hue, 2001; Miyazawa et al., 2002; Wright et al., 1985).
40 Even on its own, an OA can increase soil pH, depending on the type of residue, its rate of application and the buffer capacity
of the soil (Haynes & Mokolobate, 2001). The main mechanisms for this acid-neutralizing effect are (i) the proton uptake of
the organic anions of humic substances that act as weak bases, (ii) decarboxylation of organic acids during residue
decomposition, and (iii) ammonification of residue nitrogen (N) (Haynes & Mokolobate, 2001; Wong & Swift, 2003; Yan et
al., 1996). Conjugated bases of organic acids derived from the added residues (further referred to as organic anions) can
45 leach down the soil profile and increase soil pH in deeper layers due to continued decarboxylation and ammonification
(Butterly et al., 2021; Tang et al., 2013). However, their effect on soil pH is variable and prone to re-acidification, as the
initial pH increase is often followed by a decrease when nitrification occurs (Yan et al., 1996). When combining lime with
an OA, the temporary increase in pH in deeper soil layers, caused by the OA, reduces the pH gradient between the limed
layer and that below. This can be sufficient for lime-derived alkalinity to leach out from the amended zone and bring about a
50 more long-lasting increase in pH in the subsoil (Butterly et al., 2021). Additionally, the organic anions can function as the
accompanying anion for the transport of lime-derived Ca²⁺. This calcium can replace exchangeable H⁺ and Al³⁺ in the
subsoil, thereby further increasing soil pH and decreasing Al toxicity (Haynes & Judge, 2008; Hue & Licudine, 1999; Smith
et al., 1995; van der Watt et al., 1991). For example, Lauricella et al. (2021) found in a column leaching experiment that soil
pH in columns amended with the combination of lime and vegetable garden compost increased by 0.14 units in the first two
55 centimeters below the amended zone and with 0.08 units in the three cm below that, compared to the lime-only control.
However, very little is known about the factors influencing organic matter-mediated alkalinity leaching in soil. The success
of this process likely depends on the soil's affinity for retaining the organic matter. Binding of DOC occurs on free binding
sites of Fe and Al oxyhydroxides (Kindler et al., 2011). The available binding sites can be inferred from the amorphous
Fe+Al content in the soil, corrected for anions that already occupy these sites. These anions are mainly phosphate and
60 organic anions (RO⁻) (Verbeeck et al., 2017). In spite of this, the specific influence of the (P+RO⁻)/(Fe+Al) ratio on organic
matter-mediated lime leaching has never been tested. This gap is particularly relevant as subsoil acidity is an important issue
in weathered soils with low (P+RO⁻)/(Fe+Al) ratios (Kögel-Knabner & Amelung, 2014).



This study was set up to identify the mechanisms of subsurface liming imposed by applying the combination of lime and organic amendments at the soil surface. Our first hypothesis is that alkalinity leaching is negatively influenced by DOC adsorption in the subsoil (i.e. under the treated soil), so in soils with large DOC solid-liquid distribution coefficients (K_D values). Additionally, most research on subsurface liming has focused on leaching in sieved soils, despite evidence that preferential flow through macropores in intact soils can greatly increase chemical leaching (e.g., pesticides, fertilizers, trace metals) compared to matrix flow in sieved soils (de Jonge et al., 2004; Jacobsen et al., 1997; Jarvis, 2007; Lægdsmand et al., 1999; Paradelo et al., 2013; White, 1985). Our second hypothesis is that preferential flow in macropores enhances leaching of alkalinity, meaning that non-dissolved lime particles and lime-bound organic anions could enhance subsoil pH more in intact soils than in sieved ones, especially in short-term lab experiments. Two consecutive column leaching experiments were set up to test these hypotheses. In both experiments, soils were packed in columns, with the topsoil layer treated for each soil with CaCO_3 , an organic amendment, or a combination of both, in addition to an untreated control, after which the columns were leached with artificial rainwater. Two acid soils with contrasting K_D values of the DOC (a Podzol and a Ferralsol) were used in the first experiment. In the second experiment, a third acid soil (a Retisol) was used, with half of the columns sampled as intact soil cores and half packed with the same soil after sieving. These three soils were selected as representative examples of soils exhibiting low, average and high DOC sorption, with the intention of creating a gradient in the success of organic matter-mediated leaching.

2 Materials & methods

2.1 Soil sampling and preparation

Three acid soils were sampled at different locations (Table 1). The first one was sampled from a Podzol in clearcut area after 40 years of Norway spruce (*Picea abies* L.) on former agricultural land in Riel, the Netherlands. The second one was sampled from a Ferralsol in uncultivated land in Da Loan, Vietnam. The third one was sampled from a Retisol in a forest in Bertem, Belgium. In Riel and Da Loan, bulk soil was sampled from the top 20 cm. In Bertem, sampling of bulk soil and an additional sampling of six undisturbed soil columns was performed below the organic layer (forest floor). The undisturbed columns were 14 cm in soil height and were carefully transported to the lab to avoid soil structure disturbance. All bulk soil was air-dried and sieved to 2 mm.

Soil pH was determined in 0.01 M CaCl_2 (1:5 solid-liquid ratio). Soil buffer capacity was defined as the slope of the curve (Fig. S1) of the amount of base added to the soil ($\text{mmol OH}^- \text{kg}^{-1}$ soil) as a function of soil pH (0.001 M CaCl_2 , 1:5) when the soil was limed to different degrees with $\text{Ca}(\text{OH})_2$ until a pH of 5.5. The cation exchange capacity (CEC) was determined using the cobalt hexamine method (Protocol ISO 23470 (2007)). Total soil organic carbon (%SOC) was determined on oven-dried samples at 105° C with an elemental analyser (Carlo Erba EA1108) in tin capsules. The concentrations of amorphous Al and Fe oxyhydroxides (and the P associated with them) were determined with oxalate extraction on soil samples dried at 45° C according to Schwertmann (1991), followed by measurement with ICP-OES (ICP-OES Thermo Scientific iCAP 7000



95 series). All these soil analyses included internal soil reference materials and analytical replicates to ensure accuracy and precision. All results are reported on a dry weight (105 °C) basis. Additional soil properties with a description of corresponding analysis methods are given in Table S1.

A combined index was calculated from the soil analyses to rank the soils in terms of the strength of net DOC sorption. The adsorption of DOC in soils is likely the result of sorption of binding sites of dissolved humic substances to free binding sites
100 on Fe & Al oxyhydroxides. These binding sites are commonly determined as half of the sum of molar oxalate extractable Fe and Al (Fe_{ox} , Al_{ox} , $mmol\ kg^{-1}$) corrected for oxalate extractable P on these sites (P_{ox} , $mmol\ kg^{-1}$) (Renneson et al., 2015). Part of the soil organic carbon is also occupying these site and anion sorption studies on soil has suggested that the competing reactive organic anion (RO^- , $mmol\ kg^{-1}$) is $1.3\ mmol\ RO^-\ g^{-1}$ total soil organic carbon present in the soil (Verbeeck et al., 2017). Hence, the saturation index of the DOC binding sites can be calculated from the oxalate extracts of the soil and from
105 the SOC content as:

$$Saturation\ index\ DOC\ binding\ sites = \frac{P_{ox} + RO^-}{0.5(Fe_{ox} + Al_{ox})} \quad (1)$$

Adsorption isotherms of DOC were constructed for each soil at three different soil pH levels: native pH, $pH \approx 5$, and $pH \approx 8$. The OA used as a DOC source in the adsorption test was a green compost originating from ILVO in Ghent, Belgium. The compost had a pH of 8.8 (0.01 M $CaCl_2$, 1:5 solid-liquid ratio). The DOC concentration was determined by extracting the
110 compost with 0.001 M $CaCl_2$ at a solid-liquid ratio of 1:10. Samples were shaken for two hours, centrifuged for 15 minutes at 1400 RCF and filtered through 1.2 μm Chromafil filters. The DOC concentration in the extract was $781\ mg\ DOC\ L^{-1}$ and was measured using the combustion catalytic oxidation method (Shimadzu TOC-L CPH). The degree of aromaticity of the samples was 30%, determined using the Specific UV Absorbance (SUVA), as detailed in Amery et al. (2010). For the DOC adsorption tests, soils were first mixed with $Ca(OH)_2$ at the correct doses. For each soil and $Ca(OH)_2$ dose, aliquots of 3 g
115 were mixed with 30 mL of a 0.001 M $CaCl_2$ solution with increasing compost derived DOC (extracted from the compost in advance as described above and diluted with 0.001 M $CaCl_2$ to varying initial DOC concentrations (0-781 $mg\ DOC\ L^{-1}$)). Soil suspensions were shaken for 16 hours, centrifuged at 1400 RCF for 10 min, and filtered through a 1.2 μm Chromafil filter. The DOC concentration in the filtrate was determined with a TOC analyzer (Shimadzu TOC-L CPH). Adsorption isotherms were constructed by plotting the adsorbed DOC concentration ($mg\ kg^{-1}\ soil$) as a function of the equilibrium DOC
120 concentration in solution ($mg\ L^{-1}$) (Fig. S2). The net adsorbed DOC concentration was calculated from the difference in DOC concentration in solutions between those before and after the reaction with the soil. In the Ferralsol, the DOC concentrations were always lower after than before the reaction, i.e., there was a net DOC adsorption. Conversely, in the Podzol and Retisol, DOC concentrations were always higher after reaction with the soils than before, i.e. net DOC desorption from soil occurred, even at high added DOC concentrations. Therefore, the curves of negative sorption (i.e.,
125 desorption) versus equilibrium DOC were extrapolated to zero DOC concentration in solution ($mg\ L^{-1}$), yielding a soil sorbed DOC ($mg\ DOC\ kg^{-1}$, a negative value) with equal fate as added DOC equal to zero to determine the native adsorbed DOC concentration. The y-axis was then reconstructed by adding this value to the net sorbed DOC concentration, resulting



in positive values for all total adsorbed DOC concentration. The adsorption isotherms were fitted as Freundlich curves. The linear parts of these curves were summarized with the initial slope, i.e. the solid-liquid distribution coefficient K_D ($L\ kg^{-1}$) and the linear parts were used by considering all the points between 0-25 mg DOC L^{-1} (Ferralsol) and 0-150 mg DOC L^{-1} (Podzol and Retisol). A calculation of the initial DOC concentration in the soil solution upon adding the OA to the soil in the column experiment described below showed initial DOC concentrations of 0.6 mg L^{-1} (Ferralsol), 33.9 mg L^{-1} (Podzol) and 9.7 mg L^{-1} (Retisol), i.e. these soil+compost mixtures were within that linear part of the curves.

2.2 Set-up of the column experiments

Two consecutive leaching experiments were performed in a column setup previously described in detail in Bergen et al., 2023. In short, plexiglass cylinders of 6 cm diameter were filled with soil until a height of 16 cm. The upper 2 cm of each soil column was treated either with $CaCO_3$ at a dose of 5 g kg^{-1} , an OA at a dose of 10 g dry matter kg^{-1} , or a combination of both, in addition to an untreated control. The OA was the green compost from ILVO, Ghent, described in the previous section. In Experiment 1, one replicate was included for the control and OA treatments, while two replicates were included for the lime and lime+OA treatments, resulting in 12 columns. In Experiment 2, the same design was made and half of the columns ($n=6$) were sieved while the other half were intact ($n=6$), with a 2 cm treated layer of the sieved soil added on top. The soil densities after filling of the columns were 1.35 g cm^{-3} (Podzol), 1.15 g cm^{-3} (Ferralsol), 1.05 g cm^{-3} (Retisol, sieved) and 1.29 g cm^{-3} (Retisol, intact). The soil columns were wetted to field capacity and placed on ceramic plates pre-wetted with ultrapure Milli-Q water. The columns and plates were then placed in PVC housing with rubber rings for sealing. The bottom of each housing was attached to an Erlenmeyer flask to collect the percolate. The Erlenmeyer flasks were connected to a vacuum pump, which maintained the pressure at 900 mbar to achieve unsaturated flow conditions. Each column was irrigated with artificial rainwater composed of 1 mM $CaCl_2$, 0.003 mM KOH, 0.02 mM NaOH and 0.02 mM H_2SO_4 (pH of 5.26) from a separate container through a peristaltic pump (Watson-Marlow 205 U). The average simulated rainfall intensity was 2 mm day^{-1} . In Experiment 1, columns were leached for three weeks with a total leaching of 0.61 pore volumes for the Podzol and 0.53 pore volumes for the Ferralsol. In Experiment 2, columns were leached for eleven weeks with a total leaching of 1.8 pore volumes for the sieved Retisol and 2.13 pore volumes for the intact Retisol.

2.3 Column dismantling and soil analyses

At the end of each experiment, the columns were dismantled. In Experiment 1, the soil columns were sliced at 1 cm intervals until a depth of 16 cm. Soil pH at each depth was measured in 0.01 M $CaCl_2$ at a 1:5 solid-liquid ratio after 2 hours of shaking. The DOC concentration at each depth was determined by performing a 1:1 solid-liquid extraction with 0.001 M $CaCl_2$. The DOC concentration of the extracts was measured via the combustion catalytic oxidation method (Shimadzu TOC-L CPH) after samples were shaken for 30 mins, centrifuged for 15 mins at 1400 RCF and filtered over 0.45 μm Chromafil filters. In Experiment 2, the soil column was removed from the plexiglass cylinder and carefully sliced lengthwise into two equal parts with a galvanized iron wire. Half of the column was used to determine the pH and DOC concentration as



160 a function of the depth as described for Experiment 1. The other half was used to image the *in situ* soil pH in two dimensions using a planar optode (PO).

2.4 Planar optode imaging

Imaging of the *in situ* soil pH along the depth profile was accomplished using a PO system (“VisiSens TD”, PreSens GmbH, Regensburg, Germany) in Experiment 2. A planar optode is an optical device that uses sensor foils containing an analyte-
165 sensitive dye immobilized in an analyte-permeable matrix brought into contact with the sample. When excited by a light source, the dye emits a fluorescence signal that changes dynamically with varying analyte concentrations. A digital camera captures the signal, and the software translates it into a color image of the analyte distribution (Kreuzeder et al., 2018; Li et al., 2019; Santner et al., 2015; Tschiersch et al., 2011). In this study, pH-sensitive foils (7 x 2.5 cm) were fixed to a glass plate and applied to the cut-open half of the soil column. The foils were left to equilibrate with the soil solution for 24 h
170 before imaging. The PO was calibrated with 12 citrate buffers at an ionic strength (IS) of 25 mM and a pH ranging from 3.06 to 5.36. This IS was chosen to mimic the IS of the soil solution in the samples.

2.5 Statistical analysis

Statistical differences in pH values between soil slices (=depth) within the same column were determined by ANOVA followed by a Dunnett’s test (control = 5-6 cm depth) at 0.05 level of significance. Statistical differences in pH values
175 between different treatments at corresponding depths within each soil were determined by analysis of variance (ANOVA) followed by a Tukey’s test at 0.05 level of significance. Treatment effects on the pH profiles were also analyzed with a functional approach, i.e. non-linear regression. First, all pH values of soil treatments were corrected to that of the untreated control. This yielded $\Delta\text{pH}_{(\text{depth})}$ data, i.e. the difference in pH between a treated soil and the untreated control at a given depth. The $\Delta\text{pH}_{(\text{depth})}$ exhibited a sigmoidal decreasing trend towards depth under the treated layer and this was fitted with the
180 following 4-parameter model:

$$\Delta\text{pH}_{(\text{depth})} = \Delta\text{pH}_{\text{max}} - \frac{\Delta\text{pH}_{\text{max}} - \Delta\text{pH}_{\text{back}}}{1 + e^{-\text{slope} \cdot (\text{depth} - b)}} \quad (2)$$

where $\Delta\text{pH}_{\text{max}}$ (-) is the maximum value of ΔpH , $\Delta\text{pH}_{\text{back}}$ (-) is the background ΔpH , slope (cm^{-1}) represents the steepness of the curve and b (cm) is the inflection point of the curve.

Equation 3 shows that $\Delta\text{pH}_{\text{max}}$ is the asymptote at the soil surface (depth = 0 cm):

$$185 \quad \Delta\text{pH}_{(\text{depth}=0)} = \Delta\text{pH}_{\text{max}} - \frac{\Delta\text{pH}_{\text{max}} - \Delta\text{pH}_{\text{back}}}{1 + e^{\text{slope} \cdot b}} \approx \Delta\text{pH}_{\text{max}} \quad (3)$$

when $e^{\text{slope} \cdot b} \gg 1$.

The ΔpH -depth profiles of all columns were fitted with the nonlinear fitting option in the JMP software (JMP pro 17, SAS Institute Inc.). For the OA+lime and lime treatments, data of all four columns were fitted in one set with an assumed difference in parameter values for each of the four parameters. The statistical difference in parameter values were tested to



190 identify treatment effects on the extent and depth of penetration of the alkalinity. The depth profiles of the Δ DOC concentrations were fitted by replacing Δ pH with Δ DOC in Equation 2 (see supplementary information).

3. Results

3.1 Soil properties

195 Selected soil properties are given in Table 1. All three soils were acid (pH 3.4 - 4.2). The distribution coefficients of DOC at a common pH = 5 ranked Ferralsol > Retisol > Podzol and varied over an order of magnitude. This ranking also follows the ranking of the Saturation index for DOC binding sites (high value indicates low sorption). Other soil characteristics can be found in Table S1.

Table 1: Selected characteristics of the soils used in the three-column leaching experiments including the DOC solid-liquid distribution coefficient K_D .

Soil property	units	Dutch soil	Vietnamese soil	Belgian soil
Origin		Riel	Da Loan	Bertem
Land use		Clearcut	Uncultivated	Forest
WRB soil type		Podzol	Ferralsol	Retisol
pH		3.7	4.2	3.5
pH buffer capacity	mmol OH ⁻ kg soil ⁻¹ pH unit ⁻¹	24.4	24.3	24.1
CEC	cmol _c kg ⁻¹	1.4	3.6	3.2
SOC	%	0.89	0.53	1.12
K_D DOC (pH ≈ 5)	L kg ⁻¹	2.7	175.4	10.4
K_D DOC (pH ≈ 8)	L kg ⁻¹	2.4	92.3	4.3
RO ⁻	mmol kg ⁻¹	11.6	6.9	14.6
P _{ox}	mmol kg ⁻¹	10	0.6	3.9
Fe _{ox} + Al _{ox}	mmol kg ⁻¹	46.8	58.9	81.8
Saturation index	-	0.92	0.26	0.45
DOC binding sites [§]				

200 [§]Eqn. 1

3.2 Soil pH

The pH of the treated topsoil (0-2 cm) was significantly increased by lime and lime+OA addition compared to the control in all soils (Fig. 1 and Tables S2 & S3). The OA only significantly increased topsoil pH compared to the control in the Podzol.



205 The lime+OA combination did not increase topsoil pH more than lime alone in any soil. Alkalinity movement down the soil profile was observed in the first layer below the treated layer (2-3 cm). Red asterisks in Fig. 1 mark ΔpH values that are significantly higher than the ΔpH value of the 5-6 cm layer of the same soil column, indicating an increase in pH compared to the original pH in the unamended part of the soil column. In the Podzol, soil pH in the 2-3 cm layer increased, on average, 0.6 units in the limed treatment and 1.5 units in the lime+OA treatment, compared to the original pH in the respective columns. The inflection point of the sigmoidal curve (i.e. parameter b in Eq. 2) was significantly larger (=deeper) in the

210 lime+OA treatment than in the limed treatment in this soil, indicating larger alkalinity leaching in the former (details not shown). In the Ferralsol, no such increase in pH compared to the original pH was observed below the treated layer. In the sieved and intact Retisol, soil pH in the 2-3 cm layer of the lime+OA treatment was significantly higher than in deeper layers, whereas that increase was not statistically significant in the lime only treatment. The lime+OA treatment increased pH by 1.3 units in the sieved soil and 0.9 units in the intact soil. The functional analyses of the ΔpH did not show a

215 significantly larger penetration of alkalinity ($p>0.05$) of in the lime+OA compared to the lime only.

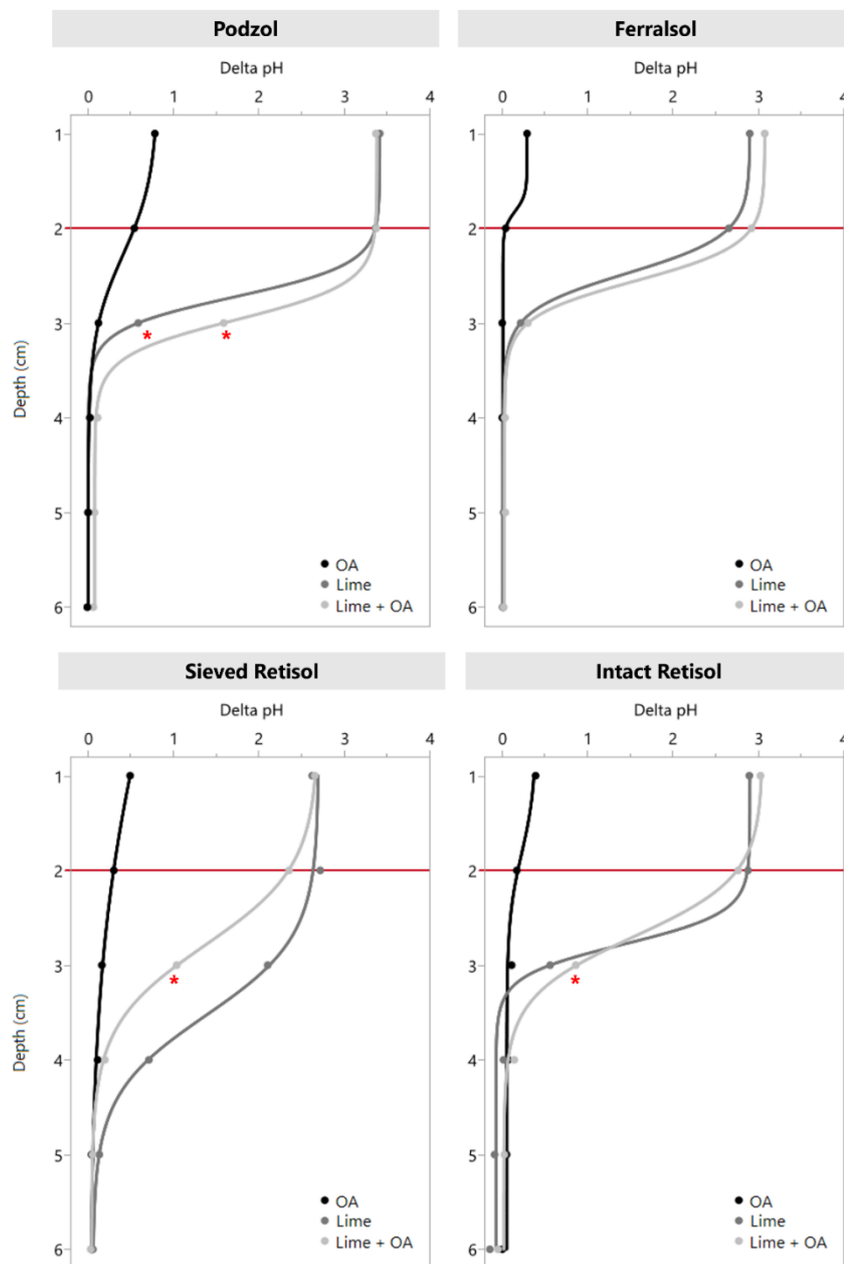


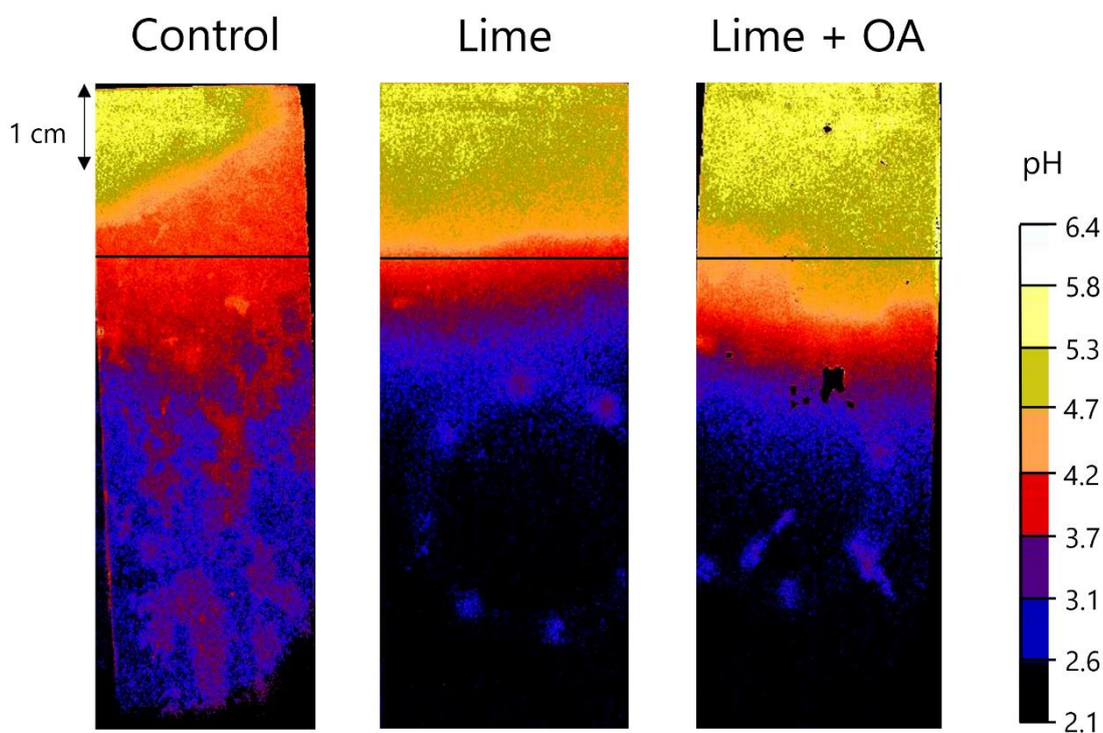
Figure 1: Data points and sigmoidal fits (Eq. 2) of the depth profiles of ΔpH (difference between pH in specific soil layer and the pH of the control treatment in the corresponding soil layer) values in soil slices after dismantling of the columns. The red horizontal lines represent the border of the treated layer. Red asterisks represent significantly higher ΔpH values (Dunnett, $p < 0.05$) than the ΔpH value of the 5-6 cm soil layer of the same column.

220

The *in situ* pH in the columns measured by the planar optode is shown in Fig. 2. Only three columns were selected for illustrative purposes: the control treatment, a limed treatment, and a lime+OA treatment of the intact Retisol. The movement



of the alkalinity front down the soil profile is visible, with deeper leaching of the alkalinity in the lime+OA treatment than in the lime-only treatment. The absolute values of the soil pH are solely indicative since incomplete contact between the sample and sensor foil may occur in unsaturated samples. The upper 2 cm layer in the intact columns consists of sieved soil with a slightly higher pH than that of the intact soil below the treated layer due to drying and rewetting, explaining the yellow-red color of the upper 2 cm layer in the control treatment.



230 **Figure 2: Planar optode image of in situ soil pH for control, lime and lime+OA treatments in the intact Retisol. The top of the image coincides with the soil surface. The horizontal black lines indicate the border of the treated layer. Note that the treated layer is a 2 cm layer of sieved soil (higher pH, Table 1) imposed on the intact column, hence the increased soil pH in the upper 2 cm of the control treatment.**

3.3 DOC concentrations

235 Clear trends in DOC concentrations are shown in Table 2 and Fig. S3. Generally, DOC concentrations were raised by the soil amendments in the following order: lime+OA > lime >> OA. The factor increase in DOC among the treatments was similar across all soils. However, absolute DOC concentrations in the Ferralsol remained markedly lower than in the other soils.



240

Table 2: DOC concentrations in soil extracts from the treated layer (0-2 cm) and the layer below (2-3 cm) after dismantling of the columns.

	DOC concentration (mg C L ⁻¹)			
	Control	OA	Lime	Lime + OA
Podzol				
0-2 cm	12.2	37.8	43.9	56.0
2-3 cm	13.5	17.5	22.9	30.3
Ferralsol				
0-2 cm	1.1	2.0	3.3	5.2
2-3 cm	0.9	0.8	1.2	1.7
Sieved Retisol				
0-2 cm	13.6	14.3	60.2	67.1
2-3 cm	15.4	16.6	57.1	65.5
Intact Retisol				
0-2 cm	13.6	17.2	53.0	60.0
2-3 cm	15.4	22.5	84.5	72.9

4 Discussion

4.1 Effect of DOC adsorption on alkalinity leaching

The results of this study indicate that combining lime with an OA leads to enhanced alkalinity leaching to the subsoil compared to a lime-only treatment (Fig. 1 and Tables S2 & S3). In the Podzol, combining lime with an OA increased the pH in the 2-3 cm layer more than twice as much as applying lime alone. Moreover, in the Retisol, statistically significant pH increases below the treated layer were only observed when lime was combined with an OA. However, no alkalinity leaching was observed in the Ferralsol. This soil type is characterized by a high Fe and Al (hydr)oxide content due to intense soil weathering (Kögel-Knabner & Amelung, 2014). These Fe and Al oxides are considered to be the most important adsorbents for DOC in soils. Therefore, the oxalate-extractable Fe and Al content in soil is a good indicator of the DOC adsorption capacity, when corrected for the already adsorbed species (mainly P_{ox} and RO⁻) (Kaiser et al., 1996; Kindler et al., 2011; Moore et al., 1992). On top of that, weathered soils typically contain clay minerals like kaolinite, which are also important adsorbents for DOC in soils (Jardine et al., 1989; Kalbitz et al., 2000). Indeed, the K_D value of the DOC adsorption isotherms (Table 1 and Fig. S2) was considerably larger in the Ferralsol than in the two other soils. Table 2 shows that the DOC concentrations measured in this soil ranged from <1 mg C L⁻¹ to about 5 mg C L⁻¹, while typical DOC concentrations in soil solution range from 1 mg C L⁻¹ to 50 mg C L⁻¹ (Herbert & Bertsch, 1995). The low concentration of organic anions in the solution probably prevented the facilitated transport of lime-derived alkalinity, confirming our first hypothesis. Our results imply that DOC-mediated alkalinity leaching is unlikely to occur in weathered soils, typically located in the humid tropics (Werts, 2023). This is a troublesome outcome, as most of the potentially arable acid soils are located in the humid tropics,



260 and resource-limited farmers in these areas could particularly benefit from cost-effective solutions to remediate (sub)soil acidity (von Uexküll & Mutert, 1995).

The pH effects in the subsoil are logically rather small due to the limited duration of column experiment, i.e. on 0.5-2.1 pore volumes. Two short calculation examples are added to estimate long-term impacts of the organic amendments for subsurface liming. For any adsorbing compound, in this case DOC, a retardation factor R represents the time required to travel through
265 the soil relative to that of water, and is defined as $R = 1 + \rho * K_D * \theta$ with ρ the density of the absorbent (kg L^{-1}), K_D the solid-liquid distribution coefficient (L kg^{-1}) and θ the volumetric moisture content (-). Since water travels about 2.3 m year^{-1} (assuming a net drainage excess in tropical soils of 0.7 m and a volumetric moisture content of 0.3), it follows that DOC leaches $2.3 \text{ m}/R$ annually. With the given DOC K_D value at pH 5 (Table 1), a soil bulk density of 1.3 kg L^{-1} and a volumetric moisture content of 0.3 , the DOC leaching depth is only 3.1 mm year^{-1} for the Ferralsol. The adsorption of DOC can be
270 lowered by increasing soil pH through liming, due to the decreasing positive charge on the Fe and Al (hydr)oxides in the soil at increasing pH (Tipping, 1981). Indeed, when recalculating for pH 8, a DOC leaching depth of 5.8 mm year^{-1} is found. Although the yearly leaching depth almost doubled by increasing the pH from 5 to 8, it would still take about 17 years before the DOC has leached to the subsoil ($> 0.1 \text{ m}$) in case of surface application. In contrast, a DOC leaching of 7.9 cm year^{-1} can be calculated for the Podzol at pH 5, when assuming a drainage excess of only 0.3 m year^{-1} in temperate regions.

275 The second calculation is based on the total alkalinity generated by the DOC leaching to deeper layers on the longer term. First, the amount of negatively charged ligands originating from the added DOC in the topsoil layer that leaches to the subsoil in the Ferralsol is calculated using Visual MINTEQ (*Visual MINTEQ*, 2021). The IS was fixed at 0.001 M and Ca^{2+} was added at a total concentration of 0.001 M . The DOC was included using the Nica-Donnan model, at 5 mg C L^{-1} , in line with the measured DOC concentration in the topsoil layer of the lime+OA treatment (Table 2). The model was run at pH 7
280 and at pH 4.2, corresponding to the pH of the topsoil and subsoil layer in the lime+OA treatment (Table S2). The concentration of Ca^{2+} bound to DOC was $2.72 * 10^{-5} \text{ M}$ in the topsoil and $1.97 * 10^{-5} \text{ M}$ in the subsoil. The difference between these two values is the concentration alkalinity released upon this pH change, or $7.5 * 10^{-6} \text{ M Ca}^{2+}$. The DOC that was bound to this Ca can bind two protons for each Ca^{2+} -ion released, meaning that the amount of negatively charged ligands able to bind protons in the subsoil is $1.5 * 10^{-5} \text{ M}$. A subsoil layer of 10 cm in depth, 1 m^2 in surface area and with a bulk density of
285 1.3 kg L^{-1} was considered. With a net drainage excess of 700 L year^{-1} through this unit area, the negative charge leaching to the subsoil is $10 \text{ mmol year}^{-1}$. At a total amount of 130 kg soil in the subsoil layer under consideration, this corresponds to $0.08 \text{ mmol year}^{-1} \text{ kg}^{-1} \text{ soil}$ that can contribute to neutralize the protons in the subsoil. The pH increase in the subsoil layer for the Ferralsol with a buffer capacity of $24.3 \text{ mmol OH}^- \text{ kg}^{-1} \text{ soil pH unit}^{-1}$ (Table 1) is then equal to $0.003 \text{ pH units year}^{-1}$, i.e. vanishingly small. The same calculation for the Podzol with a DOC concentration of 50 mg C L^{-1} (Table 2), pH values of
290 topsoil and subsoil layers equal to 7 and 3.6 (Table S2), net drainage excess of 0.3 m year^{-1} and a buffer capacity of $24.4 \text{ mmol OH}^- \text{ kg}^{-1} \text{ soil pH unit}^{-1}$ (Table 1) yields an pH increase of $0.02 \text{ units year}^{-1}$. That value is within the same range as experimentally found by Azam & Gazey (2020), i.e. subsurface soil pH increase remained limited to $0.049 \text{ pH year}^{-1}$. Considering these values, it would take over 300 years to increase the pH in the subsoil with 1 pH unit in the Ferralsol, while



it would only take 50 years in the Podzol. In reality, increasing soil pH in the strongly sorbing Ferralsol would be even more
295 difficult due to fixation of DOC on the soil matrix (see first calculation example). These simplistic calculations show that
DOC-mediated alkalinity leaching is unlikely to happen at realistic timescales in soils such as the Vietnamese one with high
concentrations of DOC reactive binding sites.

4.2 Effect of soil structure on alkalinity leaching

The pH maps with the planar optodes revealed that alkalinity migrates deeper with OA in structured soils. However, bulk
300 soil measurements did not confirm increased alkalinity leaching in structured soils compared to that in sieved soils. This is in
contrast to our hypothesis that enhanced alkalinity leaching would take place in the structured Retisol, driven by (i) non-
equilibrium transport of dissolved organic anions complexed with lime-derived Ca^{2+} , and (ii) preferential particle transport of
non-dissolved lime particles and mobile colloids containing organic matter, possibly bound to Ca^{2+} through ligand exchange
on acid functional groups. Previous studies did show enhanced chemical leaching in structured soils, attributed to
305 macropores (pores larger than ~ 0.3 mm) that allow rapid, non-equilibrium flow of water and dissolved substances (Jarvis,
2007; White, 1985). Additionally, strongly sorbing solutes such as pesticides and P have been observed to leach more readily
in structured soils than in sieved soils due to their tendency to sorb onto mobile colloids (de Jonge et al., 2004; Larsson &
Jarvis, 2000; Paradelo et al., 2013). These colloids are efficiently filtered in matrix flow within sieved soils but are readily
transported via macropore pathways in structured soils (Jacobsen et al., 1997; Jarvis, 2007). It is possible that such
310 preferential transport of alkalinity did happen in this study, but bulk measurements of soil pH and DOC concentrations failed
to detect localized effects. Figure 2 reveals regions of elevated soil pH (dark blue) in untreated zones of lime and lime+OA
treatments of the intact soil, suggesting the presence of preferential flow in macropores. Such effects, though minor, could
hold substantial implications in field conditions where plant roots actively exploit macropores for water and nutrient uptake
(Atkinson et al., 2020; Colombi et al., 2017). Yet, the possibility of imaging artifacts due to incomplete sample-sensor
315 contact in Fig. 2 cannot be excluded. Two possible explanations are given for the lack of pronounced preferential flow
observed in that case. First, the applied irrigation rate, averaging 2 mm day^{-1} , may have been insufficient to generate non-
equilibrium flow in soil macropores. Literature indicates that irrigation intensities exceeding approximately 1 mm h^{-1} are
typically required to activate such flows (Beven & Germann, 1982; Jarvis, 2007). Higher rates could not be achieved in this
experiment, as excessive irrigation risked overflowing the columns. This risk increased further by potential clogging of the
320 porous ceramic plates at the base of the columns. Second, the used Retisol may be low-structured, with limited macropore
presence and weak pore connectivity. Water primarily moves through the soil matrix in such soils, exposing solutes to a
larger surface area and increasing interaction with soil particles, compared to preferential flow in structured soils. This leads
to more adsorption and dispersion of solutes, which is aggravated by the high retention time of water in low-structured soils
(Jarvis, 2007; Norgaard et al., 2013).



325 **5 Conclusion**

This study confirms that combining lime with an organic amendment can enhance alkalinity leaching to the subsoil compared to a lime-only treatment, yet the success of this process depends on soil properties. Specifically, highly weathered soils, such as the Vietnamese Ferralsol, show limited DOC-mediated alkalinity transport due to strong DOC adsorption, which likely prevents lime-derived Ca^{2+} from reaching the subsoil. These results suggest that DOC-mediated alkalinity leaching is unlikely to occur in highly weathered tropical soils. This is a challenging outcome for acid-soil management in these regions, where low-cost liming solutions are needed. Contrary to our hypothesis, alkalinity leaching was not more pronounced in structured soils than in sieved soils. While preferential flow through macropores has been shown to promote chemical leaching in structured soils, this effect was not observed in our study, potentially because local effects remained undetected by bulk measurements, or due to limited irrigation rates and/or low macropore connectivity in the Retisol. Our results underscore the need for further research into the complex interactions of soil chemistry, structure, and hydrology that govern alkalinity leaching, especially in field conditions where macropore flow and root uptake may alter the transport dynamics of organic anions and lime.

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Conflict of interest

The authors declare that there is no conflict of interest.

345 **Author CRediT statement**

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Lynn Van Aelst: Formal analysis, Investigation

Toon van Dael: Conceptualization, Methodology, Writing – Review & Editing, Supervision

350 Erik Smolders: Conceptualization, Methodology, Writing – Review & Editing, Supervision, Funding acquisition



Declaration of generative AI and AI-assisted technologies in the writing process

During the preparation of this work, the author(s) used ChatGPT in order to improve their writing style. After using this tool/service, the author(s) reviewed and edited the content as needed and take(s) full responsibility for the content of the publication.

355 Data availability

Data will be made available upon request.

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