Soil geochemistry as a driver of soil organic matter composition: insights from a soil chronosequence

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Abstract. A central question in carbon research is how stabilization mechanisms in soil change over time with soil development and how this is reflected in qualitative changes in soil organic matter (SOM). To address this matter, we assessed the influence of soil geochemistry on bulk SOM composition along a soil chronosequence in California, USA, spanning 3 million years. This was done by combining data on soil mineralogy and texture from previous studies with additional measurements on total carbon (C), stable isotope values (δ¹³C and δ¹⁵N), and spectral information derived from diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS). To assess qualitative shifts in bulk SOM, we analysed the peak areas of simple plant-derived (S-POM), complex plant-derived (C-POM), and predominantly microbial-derived organic matter (OM; MOM) and their changes in abundance across soils with several millennia to millions of years of weathering and soil development. We observed that SOM became increasingly stabilized and microbial-derived (lower C:N ratio, increasing δ¹³C and δ¹⁵N) as soil weathering progressed. Peak areas of S-POM (i.e. aliphatic root exudates) did not change over time, while peak areas of C-POM (lignin) and MOM (components of microbial cell walls (amides, quinones, and ketones)) increased over time and depth and were closely related to clay content and pedogenic iron oxides. Hence, our study suggests that with progressing soil development, SOM composition co-varied with changes in the mineral matrix. Our study indicates that structurally more complex OM compounds (C-POM, MOM) play an increasingly important role in soil carbon stabilization mechanisms as the mineral soil matrix becomes increasingly weathered.

1 Introduction

Soils harbour the largest and most active terrestrial carbon (C) pool on earth (Jobbágy and Jackson, 2000; Lal, 2008). Over the past decades, many of the detailed mechanistic processes determining the fate of soil organic matter (SOM) in soils have been well studied (Schmidt et al., 2011; Lehmann and Kleber, 2015; Kleber et al., 2021). Yet, it remains unclear how these individual processes evolve over time as stabilization mechanisms depend on the soil matrix properties which in turn are subject to constant changes through soil formation (Bradford et al., 2016; Bailey et al., 2019; Kögel-Knabner and Amelung, 2021). Additionally, differences regarding pedogenic (i.e. soil type, mineralogy, soil microbiome) and environmental (i.e. climate, vegetation cover) properties not only determine the sizes of SOM stocks but also govern the abundance of diverse biochemical compounds (e.g. lignin, polysaccharides, lipids) found in the terrestrial C pool (Paul, 2016; Hall et al., 2020). Hence, SOM composition (the prevalence of certain biochemical compounds) may serve as a proxy for ecosystem properties and soil functioning, as it affects C cycling by the amount of energy provided to soil microorganisms (Nunan et al., 2015; Barré et al., 2019). Furthermore, the peak areas related to biochemical SOM compounds have been successfully used to parameterize fast- and
slow-cycling SOM pools (Todd-Brown et al., 2013; Bailey et al., 2018; Laub et al., 2020; Baldock et al., 2021).

Despite the central role that SOM composition plays in soil C cycling, we lack a thorough understanding of how it is influenced by the input of organic matter (OM) and by changes in soil C stabilization mechanisms following soil formation (von Lützow et al., 2008; Doetterl et al., 2018). At initial soil formation stages, the mainly plant-derived input material determines SOM composition (Khédim et al., 2021). With increasing soil age, soil properties such as soil mineralogy, texture, and soil aggregation become increasingly important for the stabilization of accumulating SOM (Chorover et al., 2004; Mikutta et al., 2006, 2009; Wei et al., 2016). For example, a comparative study of soil aggregation in relation to soil weathering found micro- and macroaggregates to become more stable with increasing weathering (Wei et al., 2016). In addition, there is an emerging consensus that certain SOM compounds are preferentially bound to mineral surfaces in organo-mineral associations (OMAs) and are thus highly important C stabilization mechanisms (Kleber et al., 2021). Many studies reported a high affinity of chemically simple carbohydrates to minerals (i.e. low molecular weight compounds from root exudates or of microbial origin) (Kaiser et al., 1997; Calderón et al., 2011; Kleber et al., 2011; Cotrufo et al., 2015; Lavalle L et al., 2019). Conversely, chemically more complex compounds (i.e. lignin or phenols from woody plant parts), formerly considered to be recalcitrant OM, were found to be less associated with minerals (Spießvögel et al., 2008; Kleber et al., 2011; Lavalle L et al., 2019). However, recent studies reported a preferential binding with iron oxides in the particular case of lignin (Kramer et al., 2012; Hall et al., 2016).

OMAs are preferentially found in the fine-grained, heavy soil fraction (<2 µm = clay) which is therefore often regarded as the most important fraction to explain how SOM persists in soils (Six et al., 2000; Kleber et al., 2021). Recent conceptualizations argue that the organo-mineral interaction is subject to dynamic changes through mineral weathering (Cotrufo et al., 2013; Lehmann and Kleber 2015; Kleber et al., 2021). With ongoing soil weathering, primary silicates originating from the parent material are replaced by newly formed secondary clay minerals (White et al., 1996). The sorptive capacity of clay minerals decreases as 2:1 layer type silicates (e.g. smectite) are substituted by 1:1 layer type silicates (e.g. kaolinite) (Sposito et al., 1999). In contrast, iron oxides, particularly nano-sized iron (Fe)- and aluminium-bearing (Al) oxides, gain importance with increasing soil weathering as they conserve high specific surface areas. As soil acidification progresses with soil weathering, the importance of these oxides to stabilize OM compounds becomes more pronounced (Kleber et al., 2021). Hence, the potential of different SOM stabilization mechanisms changes dynamically as a function of mineral weathering (Mikutta et al., 2009). Given that different SOM compounds have differing binding preferences, mineral weathering is thus likely to affect SOM composition. Therefore, a promising approach to gain further insights into the controls of SOM composition is the observation of changes in SOM composition along pedogenic gradients.

The aim of our study was to assess how bulk SOM composition changed along an undisturbed soil chronosequence spanning 3 million years, located under rangeland vegetation in the Mediterranean climatic conditions of California, USA. To this end, we used the C : N ratio and the stable isotope signatures δ15N and δ13C as proxies for the degree of microbial transformation of the bulk SOM and the selected peak areas of diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) measurements as proxies for SOM composition. We then combined this information with previously published data on soil mineralogy and texture from the same samples (Doetterl et al., 2018) in order to identify which drivers are most important in explaining shifts in SOM composition with soil development. Based on previous studies, we hypothesized that absorbance peak areas related to simple plant-derived OM (S-POM) would decrease with increasing soil formation as binding sites for the formation of OMAs are reduced. Consequently, the protection of S-POM compounds from microbial degradation and transformation decreases. The peak areas of complex plant-derived OM (C-POM) were expected to increase with soil age due to preferential association with pedogenic iron oxides (Hall et al., 2016; Zhao et al., 2016; Huang et al., 2019). Finally, MOM was expected to strongly increase with soil age since microbial uptake of OM and the subsequent stabilization in soils leads to an increasing share of microbial-derived SOM (Cotrufo et al., 2013, 2015).

2 Material and methods

2.1 Site description

This study links information on soil mineralogy and texture (Doetterl et al., 2018) with proxies for microbial transformation and SOM composition to derive a better understanding of long-term SOM dynamics along weathering gradients. The study area is located in the California Central Valley (Merced County, CA, USA), where granitic debris with a minor share of mafic minerals from the Sierra Nevada foothills was deposited on alluvial terraces of the Merced River during interglacial periods (Harden, 1987; White et al., 1996, 2005). The alluvial deposits cover a temporal range of 3 million years, with the youngest terrace only 100 years old (Harden, 1987; Doetterl et al., 2018). We assume that the soil age corresponds to the age of the terrace depositions. The present climatic conditions across the study region are semiarid with a mean annual temperature of 16.3 °C and a mean annual precipitation of 315 mm (Reheis et al., 2012). On the flat, plateau-like terraces, extensive, unfertilized rangeland dominates, which is not affected by wildfires. The vegeta-
tion cover is uniform along the chronosequence, consisting of annual Mediterranean grasslands, with common species including *Bromus mollis*, *Festuca* spp. (grasses), *Erodium* spp. (forbs), and legumes (*Trifolium* spp.) (Jones and Woodmansee, 1979).

### 2.2 Soil sampling

Soil samples were collected in December 2013 from 1 m³ soil pits located within a diameter of approximately 40 km in the north of Merced County (Fig. 1). The terraces are named based on different quaternary periods that led to the alluvial deposition (Harden, 1987). The two youngest terraces are named Post-Modesto (PM) 24II (0.1 ka) and Post-Modesto (PM) 22 (3 ka). The intermediate terrace Modesto 2.4 m was deposited between 8 and 30 kyr (819) ago. The two oldest terraces are Riverbank deposits (from 260 to 330 kyr (8295) ago) and Chinahat (3000 kyr old deposits). On each terrace one plot was sampled, and three field replicates were taken per pedogenic horizon (Doetterm et al., 2018). Yet, for the youngest 0.1 ka terrace, soils were only taken to a depth of 30 cm, and in the case of the intermediate Modesto 2.4 m terrace, soil samples were taken from three different soil pits. To convert the pedogenic horizons to homogeneous depth classes (0–10 and 10–30 cm), weighted arithmetic means were calculated based on the contribution of each pedogenic horizon to the respective depth class. All soils were classified as chronic Luvisols except for the 3 ka terrace (PM 22) which was classified as a dystric Fluvisol. The total sample set was comprised of 30 observations.

### 2.3 Soil analysis

Doetterm et al. (2018) quantified elemental composition, pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic organo-mineral associations and oxy-hydroxides was done using dithionite–citrate–bicarbonate (DCB) at pH 8 (Mehra and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic organo-mineral associations and oxy-hydroxides was done using dithionite–citrate–bicarbonate (DCB) at pH 8 (Mehra and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture. Elemental composition of the soil samples was determined using inductively coupled plasma atomic emission spectrometry (ICP-AES) after borate fusion (Chao and Sanzolone, 1992). Extraction of pedogenic oxides, soil pH, cation exchange capacity (CEC), and soil texture.

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The spectroscopic analysis was carried out with a Bruker IFS 66v/S Vacuum FT-IR spectrometer (Bruker Biosciences Corporation, Billerica, MA, USA) to collect diffuse mid-infrared reflectance with a wavenumber range from 4000 to 400 cm⁻¹ (2500–25 000 nm wavelength). In our analysis, 100 co-added scans were performed on each ground sample and averaged for further analysis. Spectra were normalized with the standard normal variate (SNV) method for scatter correction and a potassium–bromide (KBr) background for baseline correction. Subsequently, absorbance peak areas were integrated using a local baseline defined as a straight line connecting the absorbances at the upper and lower wavenumber limits.
of each functional group (Demyan et al., 2012). Negative peak areas were attributed to noise due to low C stocks and not to excessive signal strength as they only occurred in the subsoil. To account for potential variation in signal strength due to varying C stocks, the peak areas were divided by the respective C stocks. The treatments of the spectra were performed using the OPUS software (version 7.5.18; Bruker Optik GmbH, Ettlingen, Germany).

We differentiated three functional groups, based on extensive previous research (Ellerbrock et al., 2004; Demyan et al., 2012; Vranova et al., 2013; Parikh et al., 2014; Ryals et al., 2014; Viscarra Rossel et al., 2016; Fissore et al., 2017; Hall et al., 2018): aliphatic C–H bonds, aromatic C = C bonds, and C = O bonds (see Fig. 2). These three functional groups are known to be indicative of three groups of SOM compounds with strongly differing chemical properties and consequently differing bioavailability to microorganisms in soil systems (see Table 1). The wavenumber ranges centred at 2925 (2976–2898 cm\(^{-1}\)) and 2850 cm\(^{-1}\) (2870–2839 cm\(^{-1}\)) are attributed to antisymmetric and symmetric aliphatic C–H stretching (Parikh et al., 2014) and represent simple plant-derived organic compounds (S-POM), e.g. originating from leaf compounds (i.e. waxes), as well as certain organic acids that are secreted by plant roots (i.e. citrate and oxalate) (Vranova et al., 2013). The second wavenumber range, located at 1512 cm\(^{-1}\) (1550–1500 cm\(^{-1}\)), represents aromatic C = C compounds (Parikh et al., 2014). This functional group is related to lignin-derived, complex plant-derived compounds (C-POM) present in lignin constituents of wooden plant parts and aromatic organic acids of root exudates (Ryals et al., 2014). A large wavenumber range at 1620 cm\(^{-1}\) (1660–1580 cm\(^{-1}\)) represents a carboxylic C = O stretch of amides, quinones, and ketones, with possible shares of aromatic compounds (Parikh et al., 2014). This peak area is interpreted as microbial-derived organic compounds (MOM) and jointly represents amides and quinones, which constitute microbial cell walls, as well as a ketone contribution attributed to long-chained lipids that can also be found in plant materials (Kögel-Knabner and Amelung, 2014). Wavenumber ranges that overlap with signals from mineral compounds, i.e. between 1400 and 400 cm\(^{-1}\) (Parikh et al., 2014; Margenot et al., 2015, 2017), were excluded from our analysis.

### 2.4 Statistical analysis

Statistical analysis was carried out using R (Version 4.0.1, R Core Team, 2020). Two-way analysis of variance (ANOVA) was carried out to assess differences across soil age (0.1, 3, 19, 295, 3000 ka) and soil depth (topsoil: 0–10; and subsoil: 10–30 cm) to identify significant trends of bulk SOM composition parameters \(n = 30\). The significance level of the ANOVA results was at \(p < 0.05\). Prior to the ANOVA, residuals were tested for normality with a Shapiro–Wilk test, and the homogeneity of variances were checked with Levene’s test.
Figure 2. Baseline-corrected DRIFTS spectra of each soil age for 0–10 (a) and 10–30 cm (b) soil depth. The wavenumber ranges across which absorbance peak areas were integrated for functional group assignment are indicated with grey bars in the background. The two wavenumber ranges around 2900 cm\(^{-1}\) stand for simple plant-derived OM (S-POM), the one around 1620 cm\(^{-1}\) for microbially associated OM (MOM), and the one at 1525 cm\(^{-1}\) for complex plant-derived OM (C-POM).

test. We used Tukey’s honestly significant difference (HSD) as a post hoc test for pairwise comparison of homogeneous subgroups across soil ages. For the mineralogy and texture data taken from Doetterl et al. (2018), no ANOVA was carried out due to the low number of observations (\(n = 10\)).

Linear models were formulated to explain the influence of mineralogy and texture variables on the SOM composition (dependent variables: C : N ratio, \(\delta^{15}N\), \(\delta^{13}C\), S-POM, C-POM, MOM) independent of soil age and soil depth. We controlled for multicollinearity among all variables by calculating variance inflation factors (VIFs) and excluded all variables with VIFs above 5 (Kuhn, 2020). VIFs quantify the collinearity of a variable with the model (Fox and Monette, 1992; Fox and Weisberg, 2019). The VIF of each variable was computed as follows (James et al., 2013) (Eq. 3):

\[
\text{VIF} = \frac{1}{1 - R^2_{X_j}X_j}
\]

Out of the variables presented in Table 2, the Fe\(_{\text{total}}\) : Si\(_{\text{total}}\) ratio, clay content, Fe\(_{\text{DCB}}\), and Al\(_{\text{DCB}}\) were selected as explanatory variables as their inflation factor was below 5. Based on these variables, linear least squares regressions were calculated to explain SOM properties (C : N ratio, \(\delta^{13}C\), \(\delta^{15}N\), S-POM, C-POM, and MOM). To assess the variable importance of the model parameters we used the absolute \(t\) values which are calculated by dividing the parameter estimate (\(\beta_i\)) by the standard error of estimation (SE\(_i\)) (James et al., 2013; Kuhn, 2020). The \(t\) value is a measure to determine...
Table 1. Overview of wavenumber ranges used to calculate absorbance peak areas from DRIFTS, the respective functional groups related to specific biochemical properties of SOM compounds and compiled information on the origin (Demyan et al., 2018; Hall et al., 2018), and the respective SOM properties (C : N ratio, δ¹³C and δ¹⁵N values) of each functional group (Benner et al., 1987; Austin and Vitousek, 1998; Dijkstra et al., 2006; Xu et al., 2010; Vranova et al., 2013).

<table>
<thead>
<tr>
<th>SOM type</th>
<th>Wavenumber centre (range) [cm⁻¹]</th>
<th>Functional group</th>
<th>Origin</th>
<th>SOM properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>S-POM</td>
<td>2925 (2976–2898) + 2850 (2870–2839)</td>
<td>Aliphatic C–H stretch</td>
<td>Constituent of litter, component of many root exudates (i.e. citrate, oxalate)</td>
<td>Intermediate C : N ratio, δ¹³C and δ¹⁵N depleted compared to soil but higher compared to C-POM</td>
</tr>
<tr>
<td>C-POM</td>
<td>1525 (1550–1500)</td>
<td>Aromatic C = C stretch</td>
<td>Mostly lignin, related to wooden plant parts, also found in root constituents and aromatic root exudates (i.e. phthalic or vanillic)</td>
<td>High C : N ratio, δ¹³C and δ¹⁵N depleted</td>
</tr>
<tr>
<td>MOM</td>
<td>1620 (1660–1580)</td>
<td>Amide, quinone, ketone C = O stretch, aromatic C = C, and/or carboxylate C–O stretch</td>
<td>Constituents of microbial cell walls (amides and quinones) and long-chained lipids (ketones)</td>
<td>Low C : N ratio, δ¹³C and δ¹⁵N enriched</td>
</tr>
</tbody>
</table>

the effect that each parameter has on the regression slope (Grömping, 2015). To give information on the significance of the impact, we added the p value attributed to each predictor (Bring, 1996). The goodness of model fit was quantified with the R² value. The root mean squared error (RMSE), as a measure of prediction error, and the mean absolute error (MAE), as a measure of model bias, were quantified by a Monte Carlo cross-validation following a leave-one-group-out principle. Thus, the RMSE and MAE were calculated as the difference between predicted values based on a randomly selected training set (80 % of the observations) in relation to a randomly selected validation set (20 % of the observations) and relative to the sample size. The RMSE, the MAE, and the R² values were averaged, and respective standard deviations were calculated based on 100 iterations of the cross-validation. In addition, we calculated the relative RMSE (rRMSE) and relative MAE (rMAE). They were calculated as the quotient of the RMSE and the mean of the response variable and multiplied by 100 (to get percentages).

3 Results

3.1 Changes in soil properties and SOM composition along the chronosequence

With increasing soil age, the soil mineral matrix was progressively weathered. Weathering and leaching of cations were reflected in decreasing trends of the CEC from 15.2 to 4.5 at 0–10 cm depth (difference between youngest (0.1 ka) and oldest (3000 ka) terrace) and from 14.2 to 2.1 at 10–30 cm depth (see Table 2). Similarly, soil pH showed a trend to decrease from 6.8 to 4.7 in 0–10 cm and 6.8 to 4.3 at 10–30 cm depth (see Table 2). Yet, the gradient observed for CEC and soil pH was not found in base saturation (Bsat) values which did not show any clear pattern with increasing soil age but were consistently higher at the 10–30 cm compared to 0–10 cm depth (except for the youngest 0.1 ka terrace). We observed an increase in Si_total from 241.5 to 393.4 g kg⁻¹ at 0–10 cm and from 280.3 to 389.4 g kg⁻¹ at the 10–30 cm depth with increasing soil age. Al_total and Fe_total contents showed the opposite trend, decreasing with soil age in both depths. However, this decrease was not displayed in the fractions of DCB-extractable Al (AlDCB) and Fe (FeDCB). While AlDCB stagnated along the age gradient at values ranging from 0.7 to 1.3 g kg⁻¹ at 0–10 cm depth and 0.6 to 1.4 g kg⁻¹ at 10–30 cm depth, FeDCB exhibited an increasing trend in both depths. At 0–10 cm depth the values increased from 6.2 to 13.6 g kg⁻¹ and at 10–30 cm depth from 6.1 to 13.5 g kg⁻¹. The higher FeDCB values among the DCB-extracted pedogenic oxides reflect the high potential of Fe to be quantitatively important for SOM binding. The Fe_total : FeDCB ratio showed the highest values in the soils of the 19 ka terrace (7.6 at 0–10 cm and 6.4 at 10–30 cm depth). The Al_total : AlDCB ratio was highest in soils of the 295 ka terrace with values of 88.5 at 0–10 cm depth and 106.8 at 10–30 cm depth (see Table 2). Soil texture showed no clear patterns with soil age but instead differed between the 0–10 and 10–30 cm depths (Table 2). Except for the youngest terrace, silt and clay fractions were more abundant at 10–30 cm depth, while the proportion of sand was mostly lower relative to the 0–10 cm depth.

Overall, bulk total C decreased significantly with soil depth (ANOVA results were significant at p<0.05) and with...
Table 2. Overview of soil mineralogy and texture changes with increasing soil age at 0–10 and 10–30 cm depth (n = 10). The mineralogy was measured using inductively coupled plasma atomic emission spectrometry (ICP-AES). Texture was quantified with a laser diffraction particle size analyser. Pedogenic oxides were extracted in duplicates with dithionite–citrate–bicarbonate (DCB).

<table>
<thead>
<tr>
<th>Soil age [ka]</th>
<th>0.1</th>
<th>3</th>
<th>19</th>
<th>295</th>
<th>3000</th>
<th>0.1</th>
<th>3</th>
<th>19</th>
<th>295</th>
<th>3000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si total [g kg(^{-1})]</td>
<td>82.7</td>
<td>65</td>
<td>81.1</td>
<td>63.9</td>
<td>29.3</td>
<td>92.2</td>
<td>73.4</td>
<td>87.1</td>
<td>64.9</td>
<td>36.7</td>
</tr>
<tr>
<td>Fe total [g kg(^{-1})]</td>
<td>36.5</td>
<td>35.3</td>
<td>38.4</td>
<td>18.2</td>
<td>23.1</td>
<td>37.1</td>
<td>39.6</td>
<td>43.7</td>
<td>18.4</td>
<td>25.3</td>
</tr>
<tr>
<td>Fe (_{\text{DCB}}) [g kg(^{-1})]</td>
<td>6.2</td>
<td>9.5</td>
<td>6.6</td>
<td>6.2</td>
<td>13.6</td>
<td>6.1</td>
<td>10.4</td>
<td>7.2</td>
<td>6.5</td>
<td>13.5</td>
</tr>
<tr>
<td>Al (_{\text{DCB}}) [g kg(^{-1})]</td>
<td>5.9</td>
<td>3.7</td>
<td>7.6</td>
<td>2.9</td>
<td>1.7</td>
<td>6.1</td>
<td>3.8</td>
<td>6.4</td>
<td>2.8</td>
<td>1.9</td>
</tr>
<tr>
<td>Ti : Zr</td>
<td>21.68</td>
<td>19.62</td>
<td>26.85</td>
<td>13.82</td>
<td>11.1</td>
<td>4.9</td>
<td>12.9</td>
<td>17.4</td>
<td>9.9</td>
<td>18.3</td>
</tr>
<tr>
<td>Clay [%]</td>
<td>58.4</td>
<td>62.6</td>
<td>64.3</td>
<td>50.3</td>
<td>68.1</td>
<td>63.1</td>
<td>69.2</td>
<td>69.3</td>
<td>53.3</td>
<td>69.2</td>
</tr>
<tr>
<td>Silt [%]</td>
<td>26.4</td>
<td>28.2</td>
<td>24.5</td>
<td>36</td>
<td>20.8</td>
<td>32</td>
<td>17.9</td>
<td>13.4</td>
<td>21.7</td>
<td>12.5</td>
</tr>
<tr>
<td>CEC [meq/100 g]</td>
<td>15.2</td>
<td>13.4</td>
<td>24.7</td>
<td>4.6</td>
<td>4.5</td>
<td>14.2</td>
<td>9.9</td>
<td>20.8</td>
<td>2.1</td>
<td>2.1</td>
</tr>
<tr>
<td>pH</td>
<td>86.6</td>
<td>109.4</td>
<td>81.1</td>
<td>103.7</td>
<td>58.3</td>
<td>74.3</td>
<td>116.6</td>
<td>95.5</td>
<td>131.2</td>
<td>85.3</td>
</tr>
</tbody>
</table>

soil age at the 0–10 cm depth (see Fig. 3a). However, the decrease was not linear, and the highest C stock was measured at 3 ka (3.6 kg m\(^{-2}\); see Fig. 3a). Furthermore, the C : N ratio was lower at 10–30 cm depth and in the 19, 295, and 3000 kyr old soils compared to the 0.1 and 3 kyr old soils (see Fig. 3b). In contrast, \(\delta^{13}C\) values showed no significant pattern with soil age but increased with soil depth (see Fig. 3c). The \(\delta^{15}N\) values increased significantly with soil age but showed no significant difference from 0–10 to 10–30 cm depth (see Fig. 3d). Similarly, the peak areas of S-POM were significantly lower at the 10–30 cm depth and remained stable with increasing soil age (see Fig. 4a). Conversely, peak areas related to C-POM and MOM significantly increased with age at 0–10 cm depth and 10–30 cm depth. Moreover, peak areas were significantly higher at 10–30 cm (see Fig. 4b and c).

### 3.2 Link between soil geochemistry and SOM composition

The DRIFTS peak areas related to S-POM decreased significantly with soil depth, while the C-POM and MOM signals rose significantly with increasing soil age and soil depth (see Fig. 4). This raised the question of whether there were identifiable soil properties driving the relative changes in S-POM, C-POM, and MOM. Therefore, we used a modelling approach to predict bulk SOM composition using the soil mineralogy and texture parameters which were not affected by multicollinearity (see “Methods”). In Table 3, the mean model fit (RMSE and \(R^2\)), respective relative measures (rRMSE, rMAE), and variable importance coefficients are presented based on the cross-validated results of the linear models with randomly divided training and control sets (see “Methods”).

The C : N ratio was best explained by the Fe\(_{\text{total}}\) : Si\(_{\text{total}}\) ratio which was the most important variable contributing significantly to the regression slope (\(p<0.01\); see Table 3). However, the \(R^2\) showed high variability and was comparatively low in relation to the other models (\(R^2\): 0.32 mean \(\pm\)0.27 SD). The rRMSE of 17.9 % and the rMAE of 16.1 % indicated a good prediction error compared to the other models (see Table 3). The \(\delta^{13}N\) values were most influenced by clay content, the Fe\(_{\text{DCB}}\), the Fe\(_{\text{total}}\) : Si\(_{\text{total}}\) ratio, and Al\(_{\text{DCB}}\). However, despite the high model fit, no model parameter was significant. Similar results were obtained for the \(R^2\) values of \(\delta^{13}C\). In this model all parameters were significant (\(p<0.001\)), and based on the absolute values of the variable importance coefficients, Fe\(_{\text{DCB}}\), the Fe\(_{\text{total}}\) : Si\(_{\text{total}}\) ratio, and clay were more important than Al\(_{\text{DCB}}\) (see Table 3). The rRMSE and rMAE were slightly higher for \(\delta^{15}N\) compared to the C : N ratio model (rRMSE: 24.2 %; rMAE: 20.3 %) and low for \(\delta^{13}C\) (rRMSE: −2.2 %; rMAE: −2.2 %). Regarding the peak areas related to S-POM, the most important significant variable was clay (\(p<0.01\); see Table 3). Yet, the explained variance was low and had a high standard deviation (\(R^2\): 0.39 \(\pm\)0.33). This is also reflected in a high rRMSE and rMAE of >100 % (see Table 3). Conversely, the linear models explaining the variance of the peak areas linked to C-POM and MOM had high \(R^2\) values (\(R^2\): 0.78 \(\pm\)0.10 and 0.57 \(\pm\)0.27, respectively) indicating a good fit. In both cases, clay content was the only significant model parameter (\(p<0.01\)). Despite the high variable importance of the Fe\(_{\text{total}}\) : Si\(_{\text{total}}\) ratio in the C-POM model, this parameter was not significant (see Table 3). The rRMSE (C-POM): 56.3 %;
Figure 3. Mean ± standard deviation of site triplicates of total C (a), the C : N ratio (b), δ\textsuperscript{13}C (c), and δ\textsuperscript{15}N (d) for 0–10 (black) and 10–30 cm soil depth (grey). Significant differences between the soil ages calculated with Tukey’s HSD test are indicated with letters (\(p<0.05\)). Differences across soil age are indicated with lowercase letters and differences across soil depth in capital letters. Soil age is spaced based on the log-transformed age differences.

Table 3. Overview of the linear regression models describing SOM composition (C : N ratio, δ\textsuperscript{15}N, and δ\textsuperscript{13}C), as well as peak areas of functional groups related to simple plant-derived OM (S-POM), complex plant-derived OM (C-POM) and mainly microbial-derived OM (MOM) based on \(n=30\) observations. Explanatory variables related to the mineral matrix and soil texture were selected (Al\textsubscript{DCB}, Fe\textsubscript{DCB}, Fe\textsubscript{total} : Si\textsubscript{total} ratio, clay content). Absolute and relative model fit parameters (root mean squared error (RMSE and rRMSE), mean absolute error (MAE and rMAE), and \(R^2\)) with standard deviations were computed following 100 iterations of a Monte Carlo cross-validation. Higher coefficients of variable importance indicate a higher importance of the variable to explain the impact on the regression slope. The significance levels of each model parameter on the linear model are denoted as \(*\) \(p<0.001\) and \(*\) \(p<0.01\).

<table>
<thead>
<tr>
<th></th>
<th>C : N ratio</th>
<th>δ\textsuperscript{15}N</th>
<th>δ\textsuperscript{13}C</th>
<th>S-POM</th>
<th>C-POM</th>
<th>MOM</th>
</tr>
</thead>
<tbody>
<tr>
<td>RMSE</td>
<td>2.0 ± 0.5</td>
<td>1.2 ± 0.5</td>
<td>0.6 ± 0.2</td>
<td>2.9 ± 1.0</td>
<td>3.2 ± 0.7</td>
<td>20.3 ± 5.7</td>
</tr>
<tr>
<td>rRMSE</td>
<td>17.9</td>
<td>24.4</td>
<td>-2.2</td>
<td>128.9</td>
<td>56.3</td>
<td>61.7</td>
</tr>
<tr>
<td>MAE</td>
<td>1.8 ± 0.4</td>
<td>1.0 ± 0.4</td>
<td>0.6 ± 0.1</td>
<td>2.5 ± 0.9</td>
<td>2.6 ± 0.3</td>
<td>18.1 ± 5.5</td>
</tr>
<tr>
<td>rMAE</td>
<td>16.1</td>
<td>20.3</td>
<td>-2.2</td>
<td>111.1</td>
<td>45.8</td>
<td>55.1</td>
</tr>
<tr>
<td>(R^2)</td>
<td>0.32 ± 0.27</td>
<td>0.64 ± 0.24</td>
<td>0.61 ± 0.32</td>
<td>0.39 ± 0.33</td>
<td>0.78 ± 0.10</td>
<td>0.57 ± 0.27</td>
</tr>
</tbody>
</table>

Coefficients of variable importance

<table>
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<tr>
<th></th>
<th>Al\textsubscript{DCB}</th>
<th>Fe\textsubscript{DCB}</th>
<th>Fe\textsubscript{total} : Si\textsubscript{total}</th>
<th>Clay</th>
</tr>
</thead>
<tbody>
<tr>
<td>C : N ratio</td>
<td>1.3</td>
<td>1.8</td>
<td>3.2**</td>
<td>0.4</td>
</tr>
<tr>
<td>δ\textsuperscript{15}N</td>
<td>1.3</td>
<td>0.6</td>
<td>5.2***</td>
<td>1.5</td>
</tr>
<tr>
<td>δ\textsuperscript{13}C</td>
<td>3.3**</td>
<td>5.0***</td>
<td>4.1***</td>
<td>4.1**</td>
</tr>
<tr>
<td>S-POM</td>
<td>2.3</td>
<td>2.1</td>
<td>1.4</td>
<td>2.2 **</td>
</tr>
<tr>
<td>C-POM</td>
<td>1.0</td>
<td>1.0</td>
<td>5.0</td>
<td>2.8**</td>
</tr>
<tr>
<td>MOM</td>
<td>1.1</td>
<td>1.6</td>
<td>0.8</td>
<td>3.7**</td>
</tr>
</tbody>
</table>
4 Discussion

Our study assessed to what extent SOM composition changes as driven by soil weathering. We found that soil mineralogical and texture properties were able to explain a large part of variance in SOM composition parameters (Table 3). SOM was increasingly processed as indicated by a significant decrease in the C:N ratio and $\delta^{15}$N over time and significant decreases in $\delta^{13}$C with soil depth. These developments were accompanied by significant increases in the peak areas related to complex plant-derived OM (C-POM) and microbial-derived OM (MOM), as well as constant peak areas related to simple plant-derived OM (S-POM) (see Fig. 5). This suggests that changes in soil mineralogy following weathering likely exert control on SOM composition.

4.1 Assessing chemical alteration of soil mineralogy, texture, and SOM composition over time and depth

Older soils are increasingly weathered, and soil fertility decreases, as indicated by the lower soil pH and CEC (see Table 2). Further, the increasing dominance of Si with ongoing mineral weathering indicates increasing amounts of low reactive clay-sized silicates, i.e. kaolinite, which offer fewer binding sites to SOM (White et al., 2005). At the same time, significantly lower total C in strongly weathered soils were accompanied by an increasing trend of pedogenic iron oxides, i.e. Fe$_{DCB}$, that likely contributed to the unaltered base saturation values. Similar effects were observed in previous soil chronosequence studies, highlighting the importance of iron-bearing mineral phases for SOM stabilization in strongly weathered soils (Chorover et al., 2004; Mikutta et al., 2009). In this study, the increasing trends in pedogenic iron oxides co-varied with proxies that indicate increasingly stabilized and microbially transformed SOM (decreasing C:N ratio, increasing $\delta^{13}$C and $\delta^{15}$N values) and a shift towards a C-limited soil environment (Coyle et al., 2009; Cotrufo et al., 2021). The stronger microbial-derived origin of SOM is reflected in the increasing $\delta^{13}$C and $\delta^{15}$N values that develop due to isotopic discrimination of microorganisms against heavier $^{13}$C and $^{15}$N (Dijkstra et al., 2006, 2008; Brunn et al., 2016; Kramer et al., 2017).

Soil fertility and SOM stabilization capacity of soils decreased along the chronosequence, thus leading to lower total C stocks on old terraces and stronger signs of microbially transformed SOM following reduced C input (Doetterl et al., 2018). Additionally, we observed significant changes in the peak areas of different functional groups of different origins and complexity regarding their chemical structures. While peak areas of simple plant-derived OM (S-POM) were less affected over time but significantly decreased with depth, peak areas of microbial-derived (MOM) and complex plant-derived OM (C-POM) significantly increased over time and with depth (see Fig. 4). We related S-POM to aliphatic compounds from litter and/or root exudates, C-POM compounds to aromatic structures of aboveground woody debris and/or aromatic root constituents, and MOM to long-chained lipids of predominantly microbial origin (see Table 1).

Throughout the quaternary period, the climate and vegetation undoubtedly varied. Nevertheless, we found that S-POM did not decrease along the chronosequence, thus indicating a steady supply by aboveground (light fraction of litter compounds) and belowground input (low-molecular weight root exudates; Nardi et al., 2005) particularly in the topsoil layer. The significant decreases with depth might be related to a lower stabilization potential of aliphatic organic acids with increasing soil depth (Vranova et al., 2013). Yet, it is note-
worthy that below the main rooting zone (>16 cm depth) increases in aliphatic compounds have been observed in similar grassland soils which was explained by increasing stabilization on mineral surfaces (Feng et al., 2007). The latter is further supported by the high affinity of S-POM, in particular citrate and oxalate, to bind with Fe and Al oxides (Fe\textsubscript{DCB} tended to increase with soil age) in acidic soils (Clarholm et al., 2015). The significant increase in MOM peak areas in parallel to the constant S-POM peak areas along the chronosequence could indicate that an increasing fraction of microbially processed and transformed OM gains importance in bulk SOM with soil age (Feng et al., 2007). Previous studies showed a strong correlation between the MOM peak area and organo-mineral associations (Kaiser et al., 2007; Demyan et al., 2012; Kaiser et al., 2012). Thus, the observed increases in our study hint towards an increasing importance of OMAs with progressing soil weathering. C-POM signals, interpreted as lignin compounds from woody plant parts, behaved similarly to MOM and significantly increased with soil age and depth. In grassland soils, increases in signals related to lignin compounds with depth were explained by the increasing dominance of root-derived OM input in subsoils (Feng et al., 2007). Yet, increases in lignin compounds were also observed following a preferential association with iron oxides which is supported by the increase in pedogenic Fe\textsubscript{DCB} with soil age (Hall et al., 2016; Zhao et al., 2016), as well as kaolinite which is supported by the increasing clay contents in our samples with soil depth (Li et al., 2019).

4.2 Mineralogical control of SOM dynamics over geological time

As indicated above, the changes in the degree of decomposition (as indicated by the C : N ratio, $\delta^{13}$C, and $\delta^{15}$N) and SOM composition (peak areas of S-POM, MOM, and C-POM) must be contemplated in the light of simultaneously occurring changes in the mineral matrix along the chronosequence. Therefore, here we discuss to which extent the relationship between soil mineralogy, texture, and SOM (de-)composition is reflected in linear models and how these findings can be embedded in our current understanding of SOM dynamics.

The properties related to SOM decomposition showed differences in the model fit (see Table 3). The C : N ratio showed the lowest model fit of all SOM composition properties. Low and variable $R^2$ values highlighted that the mineral matrix did not strongly affect the C : N ratio of bulk SOM even though SOM was progressively decomposed with increasing soil age. The only variable that contributed significantly to the explanation of the C : N ratio was the Fe\textsubscript{total} : Si\textsubscript{total} ratio. The decreasing trend of the Fe\textsubscript{total} : Si\textsubscript{total} ratio with soil age indicated that during soil weathering processes, low reactive silicates, i.e. quartz or 1 : 1 layer type of clay minerals (kaolinite), abound (White et al., 2005). The second most important predictor variable, Fe\textsubscript{DCB}, reflects an increasing importance of pedogenic iron oxides in SOM stabilization as these compounds retain very high reactive surface areas (Kleber et al., 2015).
Our findings show that the properties of the mineral matrix play a subordinate role in shaping the C : N ratio which is more strongly driven by the increasing microbial origin of the SOM (Sollins et al., 2009).

In contrast to the C : N ratio, linear models with $\delta^{13}$C and $\delta^{15}$N values as response variables showed a good model fit ($R^2 > 0.60$, Table 3). The $\delta^{13}$C values were strongly related to $Fe_{DCB}$, $Fe_{total} : Si_{total}$, and clay, while $\delta^{15}$N values were explained by clay, $Fe_{total} : Si_{total}$, and $Al_{DCB}$ (although not significantly). In both cases, the high importance of clay content as predictor supports empirical findings that SOM with a higher share of microbial-derived OM (high $\delta^{13}$N; high MOM signal) and increasingly decomposed OM (low C : N ratio; high $\delta^{15}$N, less negative $\delta^{13}$C) is predominantly found within the heavy (clay) fraction of soils (Conen et al., 2008; Sollins et al., 2009; Clemente et al., 2011; Lawrence et al., 2015). It has been previously demonstrated that SOM bound to iron oxides was enriched in $\delta^{13}$C and exhibited more MOM (Zhao et al., 2016). Likely the imprint of $\delta^{13}$C enrichment with increasing microbial transformation of the SOM manifested itself more clearly in bulk SOM as the overall C stocks decreased in the oldest soils (Yang et al., 2015).

Similar to the models explaining SOM decomposition variables, the linear models on DRIFTS peak areas had varying model fits (see Table 3). The model for S-POM had a lower model fit ($R^2 = 0.39$) compared to the models explaining C-POM and MOM ($R^2 = 0.78$ and 0.57, respectively). In all models, the clay content was the key explanatory variable which hints at different behaviours in top- and subsoils as consistently higher shares of clay were found at the 10–30 cm depth (see Table 2). The fact that S-POM was not significantly influenced by the $Fe_{total} : Si_{total}$ ratio suggests that the contribution of S-POM to OM is less related to weathering-induced changes in the mineral matrix but is rather determined by above- and belowground vegetation input, i.e. through litter components or root exudates (Vranova et al., 2013; Mueller et al., 2013). Lower signals of S-POM in subsoils compared to topsoils (see Fig. 4) indicate that the mineral stabilization of S-POM compounds in subsoils is lower or that S-POM compounds are less abundant at these depths (Vranova et al., 2013). Still, the higher variable importance of $Al_{DCB}$ and $Fe_{DCB}$ to explain S-POM potentially indicates that certain compounds, likely aliphatic organic acids derived from root exudates, are preferentially attached to minerals (Clarholm et al., 2015).

In the case of C-POM, the decreasing $Fe_{total} : Si_{total}$ ratio with increasing soil age had the highest variable importance coefficient but was not significant to the model. The high variable importance coefficient is in line with other studies that found functional groups that are considered chemically more complex (i.e. through an aromatic ring structure) to be increasingly stabilized relative to other functional groups as soils become more weathered, as overall SOM stocks decrease, and as protection through reactive mineral surfaces declines in strongly weathered soils (Angst et al., 2018). Lignin compounds represented by C-POM were shown to be better stabilized due to attachment on mineral surfaces, such as calcite (Grünewald et al., 2006), kaolinite (Li et al., 2019), and Fe oxides (Kramer et al., 2012; Hall et al., 2016; Zhao et al., 2016; Huang et al., 2019). Yet, the coefficients of variable importance of the C-POM model did not show a particular importance of $Fe_{DCB}$ and $Al_{DCB}$ (see Table 3). Potentially the more linear decrease in the $Fe_{total} : Si_{total}$ ratio might mask less consistent increases in $Fe_{DCB}$ or the varying behaviour of $Al_{DCB}$ (see Table 2). Clay content was the only significant predictor, and it was previously shown that kaolinite accumulates in the mineral matrix with increasing soil age and depth along the present chronosequence (White et al., 2005). Hence, the mineral stabilization of C-POM in heavy soil fractions might be more strongly linked to the presence of clay minerals than iron oxides (Li et al., 2019; Grünewald et al., 2006).

The peak areas related to MOM were significantly explained by the clay content (see Table 3). This is consistent with the significant model parameters for S-POM and C-POM. The clay fraction is known to harbour the vast majority of soil microorganisms and to stimulate their activity (i.e. Stotzky and Rem, 1966; Sollins et al., 2009). Therefore, it is logical that the increasingly microbial-derived imprint in the bulk SOM is significantly linked to the clay fraction. Further, the $Fe_{DCB}$ parameter had a higher variable importance coefficient which might be linked to the increasing importance of pedogenic iron oxides and MOM along the soil chronosequence. This might be caused by the increasing importance of metal surfaces in stabilizing bulk SOM but also soil microorganisms or microbial necromass in particular (Kaiser et al., 1997, Zhao et al., 2016). Yet, these interpretations are more speculative due to the heterogeneity of soil microorganisms, and the vast range of possible interactions requires further investigation.

5 Conclusions

The present observational study provides new information on qualitative changes in the SOM composition in relation to soil geochemical dynamics driven by mineral weathering from a soil chronosequence covering a gradient of 3 million years. With progressing soil development SOM stocks decreased, and we observed an increasingly microbially transformed bulk SOM (lower C : N ratio, higher $\delta^{13}$C and $\delta^{15}$N values). Simultaneously, the spectroscopic fingerprint of SOM (reflected in the peak areas related to S-POM, C-POM, and MOM) shifted as peak areas of structurally more complex compounds increased (C-POM and MOM). These qualitative changes in SOM composition co-occurred with changes in soil geochemistry and texture induced by soil formation. Hence, changes in soil texture (clay content) and an increasingly kaolinite- and iron-oxide-rich, strongly weathered mineral matrix might discriminate in favour of
structurally more complex SOM compounds (C-POM and MOM). Our study shows that soil mineralogy plays an important role in shaping SOM composition during soil weathering across large timescales. We therefore recommend further studies to assess this link in contrasting soil mineralogies or vegetation types in order to gain a better understanding of how mineral matrix and C input affect SOM composition across larger geographical areas.


Data availability. The dataset and the data description files underlying the statistical analysis can be accessed on Zenodo (https://doi.org/10.5281/zenodo.6366361, Mainka, 2022).

Author contributions. AAB and SD designed the experiments. MM performed the measurements, analysed the data, and wrote the manuscript. DW, GG, MG, and LS reviewed and edited the manuscript. LS created the map (Fig. 1). All authors contributed to writing the paper (lead author: MM).

Competing interests. The contact author has declared that neither they nor their co-authors have any competing interests.

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References


