



Formation of mineral-associated organic matter via rock weathering: an experimental test for the organo-metallic glue hypothesis

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Abstract. Mineral-associated organic matter, the dominant form of relatively stable carbon (C) in soil, often co-occurs with reactive iron (Fe) and aluminum (Al) phases across soils. Yet, how organo-metallic associations at the molecular scale give rise to emergent soil properties such as aggregate formation and the persistence of organic matter (OM) in soil remains unclear. The organo-metallic glue hypothesis proposes that dissolved metal released from weathering and microbially processed OM form cohesive organo-metallic phases that bind other particles into stable assemblages. We tested this concept using an artificial soil system comprising crushed rocks (fine basalt: 20–38 μm , coarse basalt and granite: 38–75 μm , and river sand), mixed with leaf compost and microbial inoculum, subjected to eight wet-dry cycles using artificial rainwater (pH 4.7) over 55 d. Sequential density fractionation after the incubation revealed the formation of meso-density, organo-mineral assemblages (1.8–2.4 g cm^{-3} ; MF) in the following order: fine basalt > coarse basalt > granite > sand. The accretion of C and oxalate-extractable Fe, Al, and Si in MF generally followed the same pattern. Fine basalt showed the strongest increase in extractable metals, especially Fe, in MF and the highest leaching of Fe and base cations (esp. Na and Ca). Enrichment of extractable Fe, Al, and Si in MF and their slight depletion in the high-density fraction (> 2.4 g cm^{-3}) suggest that weathering-derived metals first associated with OM, forming organo-metal-rich phases that subsequently bound other particles to form organo-mineral assemblages. MF formed in fine basalt treatment had the C : (Fe+Al) molar ratio of 0.6, consistent with organo-metal coprecipitates. Preferential incorporation of microbially-processed, N-rich OM into MF in the two basalt treatments was indicated by lower C : N ratios by 23–25 units and enrichment of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ by 0.9‰–1.2‰ and 0.6‰, respectively, relative to low-density fraction (< 1.8 g cm^{-3}). SEM and STXM/NEXAFS analyses of limited MF materials confirmed the presence of shaking-resistant microaggregates and the co-localization of microbially altered C with Fe and Al. Collectively, these results provide experimental evidence supporting the organo-metallic glue hypothesis and demonstrate that basaltic rock weathering can promote organo-mineral assemblage formation. This mechanism links microbial processing, mineral weathering, and reactive metal dynamics, offering insights into early pedogenesis and soil OM formation under rock amendment conditions.

1 Introduction

Organic matter (OM) in soil plays a fundamental role in global carbon cycling, soil fertility, and ecosystem functioning (Lehmann et al., 2020; Friedlingstein et al., 2022; Angst et al., 2023). Its long-term persistence strongly depends on its physicochemical associations with reactive minerals such as aluminosilicate clays and iron (Fe) and aluminum (Al) oxides with high surface area and reactivity (e.g., Sørensen, 1972; Baldock and Skjemstad, 2000; Wattel-Koekkoek et al., 2003; Saïdy et al., 2015; Hemingway et al., 2019). Among these, pedogenic Fe-, Al-, and Si-bearing phases, formed during mineral weathering, are especially reactive. They occur as organo-metal complexes, short-range-ordered (SRO) aluminosilicates and metal oxides, and, to a less extent, crystalline metal oxides (e.g., Harter and Naidu, 1995; Cornell and Schwertmann, 2003; Parker, 2005; Ashida et al., 2021; Watanabe et al., 2023).

At larger scales, soils rich in pedogenic metals generally store more OM. For instance, basaltic and andesitic soils often accumulate greater OM stocks than granitic soils (Orgill et al., 2017; Angst et al., 2018; Rasmussen et al., 2005; Mao et al., 2020), while OM composition also exerts some control depending on ecosystem type and climate (Sollins et al., 1996; Hall et al., 2020). Pedogenic Fe and Al contents often correlate positively with soil OM stock and, to some extent, with C persistence inferred from radiocarbon analysis (e.g., Torn et al., 1997; Percival et al., 2000; Masiello et al., 2004; Lawrence et al., 2015; Shimada et al., 2022; von Fromm et al., 2025). Their protective effects on OM may persist even under decades of intensive cultivation in humid tropics (Arai et al., 2025). At the molecular scale, these metals form various associations with OM, including hydrophobic interactions, cation bridging, adsorption, organo-metal complexation, and coprecipitation, that enhance OM resistance to microbial degradation and dissolution (Keil and Mayer, 2014; Kleber et al., 2015). Recent spectroscopic studies revealed diverse modes of OM-mineral and OM-metal/metalloid associations. Those include coprecipitates of metal and metalloid oligomers with organics (Basile-Doelsch et al., 2015; Tamrat et al., 2018; Jamoteau et al., 2023; Jamoteau et al., 2025), calcium complexation (Rowley et al., 2023), and nanoscale organic layering on mineral surface (Possinger et al., 2020; Underwood et al., 2024). However, how these molecular-scale interactions give rise to emergent soil properties such as OM persistence and hierarchical aggregate/pore structure remains a fundamental question (Asano and Wagai, 2014; Regelink et al., 2015; Rabot et al., 2018; Totsche et al., 2018; Yudina and Kuzyakov, 2023; Amelung et al., 2024). These hierarchical soil structures regulate microbial and plant root activities and thus link mineral reactivity to ecosystem functioning (Chorover, 2022; Hartmann and Six, 2023; Philippot et al., 2024).

The *organo-metallic glue* hypothesis (Wagai et al., 2020) provides a conceptual bridge between nanoscale mineral-organic interactions and macroscale aggregation, and may offer more mechanistic interpretation on previous observations (e.g., C protection within aggregates, C-metal correlations across bulk soils and physical fractions). It proposes that Fe and Al dissolved from minerals complex or coprecipitate with decomposing OM to form cohesive organo-metallic phases that act as “glue”, linking mineral and organic particles into stable microaggregates. Support for this mechanism includes the consistent co-occurrence of extractable Fe and Al with nitrogen (N)-rich OM in meso-density fractions (1.8–2.4 g cm⁻³) across diverse soils and the relatively constant OC : metal ratios among soil physical fractions within each soil (Wagai et al., 2020). Yet, testing this mechanism in natural soils is challenging because of their compositional heterogeneity and the influence of multiple pedogenic processes. An artificial soil system, which consists of defined mineral and OM components, would be suitable to assess specific mechanisms under controlled conditions. Previous experiments have shown relatively rapid development of various mineral-organic associations (e.g., Vogel et al., 2014; Pronk et al., 2017; Bucka et al., 2021). However, their formation mechanisms remain poorly understood.

This study aimed to evaluate key aspects of the *organo-metallic glue* hypothesis using an artificial soil system composed of crushed rock and leaf compost. We hypothesized that the differences in rock weathering rate, and thus the supply rate of Fe and Al, govern the formation of organo-mineral assemblages in the presence of OM. To test this, we incubated two end-member igneous rocks (granite and basaltic rock), or quartz sand (as a control), with leaf compost and a soil inoculum, subjecting the rock-OM mixtures to weekly leaching-drying cycles for 55 d. Formation of organo-mineral assemblage was assessed via sequential density fractionation, elemental and extractable metal analyses, and, to a limited extent, microscopic characterization (SEM and STXM-NEXAFS). We expected meso-density fractions to show enrichment in extractable metals relative to low- and high-density fractions, reflecting the role of weathering-derived metals in binding OM into physically-stable microaggregates.

2 Materials and Methods

We prepared four types of rock-organic matter (OM) mixtures using crushed rock and river sand with leaf compost, followed by 55 d aerobic incubation with eight wet-dry cycles (Fig. 1). Microbial respiration was not measured. Leachates were collected for chemical composition. The mixtures after the incubation were destructively sampled and fractionated by density. Each fraction was analyzed for total C and N as well as extractable metals and metalloids. For se-

lected samples and fractions, we also conducted microscopic/spectroscopic and microbial characterization.

2.1 Source materials and rock-organic matter mixture preparation

We selected two rocks with contrasting chemical composition and weatherability, and a river sand as a low-reactivity control for comparison. Basalt, a mafic igneous rock, weathers faster and releases rock-forming elements including Fe and Al more rapidly than felsic granite due to the difference in silicate mineral composition. In addition to feldspars, basalt comprises minerals of lower degree of polymerization (e.g., pyroxene), whereas granite consists mainly of more-polymerized mineral (e.g., quartz) (Gray and Murphy, 2002; Chapman et al., 2009). Basaltic rock from Akita, Japan (“dust” type, < 2500 μm , Horie-Kenzai Co.) used in this study was mainly composed of plagioclase and pyroxene, with minor magnetite, olivine, smectite, and quartz (Fig. S1 in the Supplement). The quartz detected likely resulted from the nearby lithologies. Further characterization of the basaltic rock sample is reported elsewhere (Yang et al., 2026). This rock is classified as basaltic andesite based on the Total Alkali-Silica diagram; however, for simplicity, the two particle-size treatments are referred to as “fine” and “coarse basalt”. Granite used in this study was from Hiroshima, Japan (“pan” type, $\leq 75 \mu\text{m}$, Yoko Bussan Co.) and had the elemental and mineralogical composition typical of granite (Table S1, Fig. S1 in the Supplement).

These rocks were sorted for 38–75 μm and, additionally, for 20–38 μm (only basaltic rock). The source granite was ground in a blender (WB-1, Osaka Chemical Co., Japan) followed by wet sieving to isolate the 38–75 μm particle-size class. Coarse basalt (38–75 μm) was isolated by wet sieving. Fine basalt (20–38 μm) was isolated by repeated sedimentation of < 38 μm materials in deionized water until the supernatant was free of visible particles < 20 μm based on Stokes law. Preferential loss of some fine-grain minerals (e.g., mica) and enrichment of coarser ones (e.g., quartz) during the sorting are possible. Mineralogical data, nevertheless, showed the size-sorted materials fall within a common compositional range (Fig. S1). River sand (mean particle size: 100–300 μm , Toyoura Keiseki Kogyo Co., Yamaguchi, Japan) served as a low-reactivity reference. All materials were oven-dried at 50 $^{\circ}\text{C}$. The elemental compositions of the four rock materials (granite, two sizes of basaltic rocks, and sand) were determined by X-ray fluorescence (NEX CG, Rigaku, Tokyo, Japan, Table S1). Mineralogy of granite and two basaltic rocks was identified by X-ray diffraction instrument equipped with a Cu tube (MiniFlex600-C, Rigaku, Tokyo, Japan, Fig. S1), scanned under XRF reduction mode from 3 to 90 $^{\circ}$ (2θ) with a step size of 0.01 $^{\circ}$, at a speed of 5 $^{\circ} \text{min}^{-1}$ (basaltic rock) and 10 $^{\circ} \text{min}^{-1}$ (granite). The diffraction patterns were matched to the Crystallography Open Database. Smectite identified may include vermiculite

because no phyllosilicate pre-treatment was applied. The two sizes of basaltic rock used in this study showed similar mineral composition to those reported in Yang et al. (2026). Particle size distributions (LA-920, Horiba Ltd., Kyoto, Japan) confirmed target ranges of the four rock materials (Fig. S2). The specific surface area (Table S2) was measured by N₂ gas sorption Autosorb iQ (Quantachrome Corp., Boynton Beach, FL, USA) using a multi-point BET approach (partial pressures of < 0.3) after outgassing under a vacuum at 150 $^{\circ}\text{C}$. Total C concentration of the four rock materials was < 0.3 mg g⁻¹. This trace level of C derived more likely from organic matter (e.g., dust, microbial debris) rather than inorganic C because, at least for the basaltic rock, calcite was not detected by XRD and XANES analysis (Yang et al., 2026).

Leaf compost, the sole OM source, was produced mainly from tree leaves with one year of composting. Compost was air-dried, ground, and dry-sieved to a size of 100–250 μm . The compost had a C content of 408 mg g⁻¹ with a C : N ratio of 41. The pH of the compost was 5.92 (H₂O) and 5.45 (KCl). The C structure of the compost had carbonyl-C 7.7 %, aromatic-C 9.6 %, O-alkyl-C 63.3 %, and alkyl-C 19.4 % (Fig. S3), which was assessed by solid-state ¹³C nuclear magnetic resonance (NMR) spectrometry using an FT NMR system (JNM-ECA600II, JEOL Ltd., Tokyo, Japan) according to the procedure by Hiradate et al. (2004). The NMR spectrum was divided into chemical shift regions representative of the four major types of C present: 0–45 ppm (alkyl C), 45–110 ppm (O-alkyl C), 110–165 ppm (aromatic C), 165–210 ppm (carbonyl C) (Golchin et al., 1994). The total signal intensity and the proportion contributed by each C type were determined by the integration of the spectral regions.

The rock-OM mixtures were prepared by combining 20.00 g of each rock material with 2.50 g of the leaf compost under dry conditions in three replicates (Table S3). The rock-OM mixtures had an initial C content of 4.5 %, comparable to C-rich cultivated topsoils. For the incubation experiment, the mixtures were placed in 50 mL plastic columns fitted with two layers of nylon mesh ($\approx 6 \mu\text{m}$) at the bottom.

2.2 Aerobic incubation with repeated leaching

The rock-OM mixtures were leached weekly with artificial rainwater over a 55 d experimental period. The rainwater composition was based on the typical Japanese precipitation data from the national acid deposition survey in 2014 (Horie et al., 2016): pH of 4.73 and contained the following ionic concentrations: SO₄²⁻ 18.1 $\mu\text{mol L}^{-1}$; NO₃⁻ 15.4 $\mu\text{mol L}^{-1}$; Cl⁻ 90.2 $\mu\text{mol L}^{-1}$; NH₄⁺ 17.9 $\mu\text{mol L}^{-1}$; Na⁺ 76.6 $\mu\text{mol L}^{-1}$; K⁺ 2.62 $\mu\text{mol L}^{-1}$; Ca²⁺ 5.37 $\mu\text{mol L}^{-1}$; and Mg²⁺ 9.04 $\mu\text{mol L}^{-1}$. The solution was prepared using analytical-grade reagents: HCl, KCl, CaCl₂, MgSO₄·7H₂O, NaNO₃, (NH₄)₂SO₄, and NaCl.

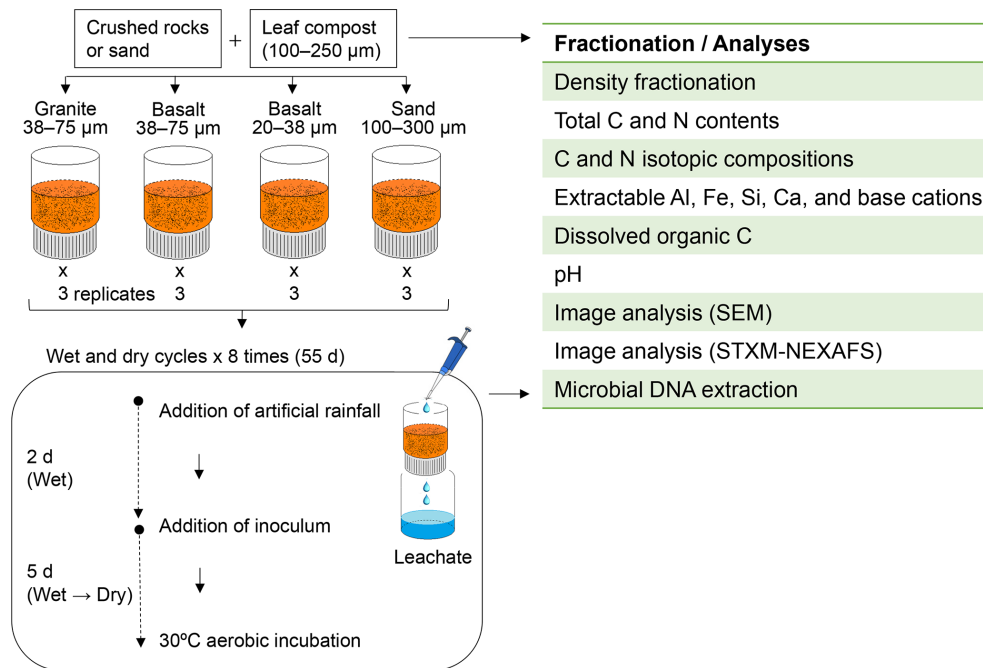


Figure 1. Schematics of the 55 d incubation experiment, including eight wet-dry cycles.

The microbial inoculum was prepared from 0–5 cm surface soil of a long-term experimental field under no-till management with annual leaf compost input at the Institute for Agro-Environmental Sciences, Tsukuba, Japan (Wagai et al., 2013a). The same leaf compost served as the sole organic matter source for the rock-OM mixtures. The soil is classified as a Hydric Hapludand (Soil Survey Staff, 2014) and a Hydric-Silic Andosol (IUSS Working Group WRB, 2015). The microbial inoculum was produced by mixing field-moist soil with deionized water (1 : 15, *w/v*) and shaking the suspension with glass beads for 30 min. The mixture was then filtered through a 5 μm membrane filter (modified from Wagai and Sollins, 2002) to retain bacteria and fungal spores/fragments while removing major grazers. The inoculum was stored at 4 °C for no longer than 24 h before use.

The incubation started with 40 mL of rainwater applied to each column, left at room temperature overnight, and the leachate was collected. This weekly volume corresponded to Japan's mean annual precipitation assuming an even rainfall distribution (Horie et al., 2016). After leachate collection, 1 mL microbial inoculum was added. Water content was 52%–63% on an air-dried basis (Table S4). The columns were then incubated in the dark at 30 °C for 5 d until nearly complete drying, forming partially cemented material. Before re-wetting each time, the hardened material was gently loosened with a spatula to approximately half the column depth to facilitate infiltration of rainwater and inoculum. Subsequent cycles received 40 mL of rainwater and 0.1 mL of inoculum (reduced from 1 mL in the first cycle). Eight wet-dry cycles were completed in 55 d (Fig. 1). After in-

cubation, the rock-OM mixtures were gently disaggregated, sieved (2 mm) using a spatula, homogenized in a plastic bag, and split for further analyses (Table S5). The leachates collected after each cycle were combined for each replicate and filtered for analyses of dissolved organic C (DOC), Al, Fe, Si, base cations (Na, Ca, K, and Mg), and pH.

2.3 Density fractionation before and after the incubation

The rock-OM mixtures after the incubation were fractionated by density with a moderate dispersion by mechanical shaking to distinguish organo-mineral assemblages as a meso-density fraction (1.8–2.4 g cm⁻³; MF) from particulate OM as a low-density fraction (< 1.8 g cm⁻³; LF) and crushed rock as a high-density fraction (> 2.4 g cm⁻³; HF) as described in Wagai et al. (2015). Briefly, 10.0 g of the rock-OM mixture after the incubation were mixed with 30 mL of sodium polytungstate (SPT-0 grade, TC-Tungsten Compounds GmbH, D-96271 Grub am Forst, Germany) with a final density of 1.8 g cm⁻³ and shaken at 120 rpm for 30 min. The suspension was then centrifuged at 2330 *g* for 20 min, and the floating material was collected on a 0.45 μm membrane filter. These steps were repeated at least three times to maximize the recovery. The material on the filter was washed with deionized water until the electric conductivity was below 50 mS cm⁻¹; this operation produced the LF fraction. The residue was re-suspended in SPT solution adjusted to 2.4 g cm⁻³ at the sample: extractant ratio of 10 g : 30 mL, shaken, and centrifuged. The floating material (1.8–2.4 g cm⁻³) was transferred to a 250 mL centrifuge bottle, mixed with deionized water, and

centrifuged at 17 000 g for 20 min. The supernatant was decanted, and this rinsing step was repeated until the electric conductivity of the supernatant became $< 50 \text{ mS cm}^{-1}$. The final material was recovered as the MF fraction of 1.8–2.4 g cm^{-3} . The remaining residue ($> 2.4 \text{ g cm}^{-3}$) was rinsed with deionized water in the same way as above to isolate the HF fraction of $> 2.4 \text{ g cm}^{-3}$. Recovered LF was oven-dried at 80 °C, and MF and HF were freeze-dried. The materials recovered in MF by this operation are, therefore, organo-mineral aggregates that are physically resistant to the disruptive forces from the fractionation steps (esp. repeated mechanical shaking).

The initial four rock materials were fractionated at a density of 2.4 g cm^{-3} to distinguish MF and HF. Assuming that the initial leaf compost entirely consists of LF, we calculated C, N, and extractable Al, Fe, Si levels in each fraction for the rock-OM mixtures on Day 0. Masses recovered in each density fraction from the rock-OM mixtures on Day 0 and Day 55 are shown in Table S6.

2.4 Chemical analyses of the mixtures

Total C and N contents in the density fraction were determined by an elemental analyzer (vario MAX cube, Elementar Analysensysteme GmbH, Langenselbold, Germany). The degree of microbial alteration of OM can be assessed by C and N isotopic compositions (e.g., Sollins et al., 2009). Thus, initial leaf compost and density fractions under the two basalt treatments after the incubation were analyzed by a continuous-flow stable isotope mass spectrometer (Delta V Advantage, Thermo Fisher Scientific Inc., USA) coupled with an elemental analyzer (Flash EA 1112 Series). To minimize the N blank, a gas-tight automatic sampler was used (Zero Blank Autosampler, Costech Analytical Technologies Inc., USA). To increase the sensitivity for analyzing the N isotopes, the combustion and reduction tubes (both 18 mm outer diameters) were exchanged with thinner ones (18–10 and 10–6 mm outer diameters, respectively; Ogawa et al., 2010; Koba et al., 2021). The obtained $^{13}\text{C}/^{12}\text{C}$ and $^{15}\text{N}/^{14}\text{N}$ ratios are shown in terms of the δ value ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) relative to Vienna Pee Dee Belemnite and the air dinitrogen, respectively. Their analytical precisions are $\pm 0.2\%$ except for the $\delta^{15}\text{N}$ values of the HF ($\pm 0.4\%$).

Extractable Al, Fe, Si, and Ca in the density fraction and bulk samples were quantified by the sequential extraction with sodium pyrophosphate (PP), oxalate-extractable (OX), and dithionite-extractable (DC), as described in Wagai et al. (2018). These extractions have been widely used as a practical method to estimate the concentrations of organo-metal complexes, short-range-ordered minerals, and pedogenic Fe oxides, respectively (Shang and Tiessen, 1998; Heckman et al., 2018; Ashida et al., 2021; Hall and Thompson, 2022). However, cautions are required for their interpretation due to the limited selectivity in extracting target phases as discussed previously (Wagai et al., 2013b; Rennert, 2019; Fuku-

masu et al., 2025). Briefly, each sample was extracted with 0.1 M sodium pyrophosphate (pH 10) at a sample: extractant ratio of 0.1 g : 10 mL at 120 rpm for 16 h at room temperature, followed by high-speed centrifugation at 29 000 g for 40 min. After collecting an aliquot of the extract (approximately 7 mL) and discarding the remaining supernatant, the residue was extracted with 10 mL of 0.2 M acidified sodium oxalate solution (pH 3) at 150 rpm for 4 h at 25 °C in the dark and centrifuged in the same way. After collecting an aliquot of the extract and discarding the remaining supernatant, the residue was added 0.167 g of sodium dithionite and then extracted with 10 mL of 0.646 M sodium citrate at 120 rpm for 16 h at room temperature and centrifuged in the same way.

The amounts of Al, Fe, Si, and base cations (Na, Ca, K, and Mg) after each of the three extractions were measured by an inductively coupled plasma optical emission spectrometer (700 series ICP-OES, Agilent Technologies, Inc., CA, United States). The combined leachate from the eight wet-dry cycles for each column of all four treatments was also analyzed in the same way. The leachate was also analyzed for DOC by a TOC analyzer (TOC-L, Shimadzu, Kyoto, Japan) using non-pergeable organic C mode and for pH.

The pH of the bulk samples from all four treatments after the incubation was measured using a glass electrode in ultra-pure water (0.4 g : 4 mL) after 1 h of shaking at 150 rpm. Subsequently, we added 1.6 mL of 3.5 M KCl to the water suspension to measure pH in the 1 M KCl conditions.

2.5 Imaging analyses of selected samples

Microscopic observation of the initial crushed rocks (granite, coarse basalt, and fine basalt) as well as the MF of their mixtures after the incubation was done using a scanning electron microscope (SU1510, Hitachi High-Tech Corporation, Tokyo, Japan). The subsets of MFs (stored in a suspension at ca 5 °C after the density fractionation without freeze-drying) were diluted in ultra-pure water and deposited on a carbon tape, air-dried, and coated with palladium before the observation.

The subsets of MF from selected treatments (granite and coarse basalt) were assessed to examine the spatial arrangement of C, Al, and Fe, as well as C functional group within the meso-density materials (largely present as shaking-resistant microaggregates) in MF using scanning transmission X-ray microscopy (STXM) and near-edge X-ray absorption fine structure (NEXAFS). Another subset of suspended MF was diluted with ultra-pure water and weakly sonicated ($< 10 \text{ J mL}^{-1}$) followed by deposition on a Si_3N_4 window (50 nm thick, window size 500 μm). To minimize the re-aggregation on the Si_3N_4 window upon drying, the samples were slowly dried at room temperature. Carbon, Al, and Fe were recorded by a compact STXM at BL-19A beamline in the Photon Factory of the High Energy Accelerator Research Organization, Ibaraki, Japan (Takeichi et al., 2016). The STXM data analysis was carried out with the IDL pack-

age aXis 2000 (Hitchcock, 2023). The distribution maps of C, Al, and Fe were obtained by subtracting the post-edge optical density (OD) image from the pre-edge OD image, respectively. Specifically, we collected images in the X-ray energy region of 280.0 and 300.0 eV for C K-edge adsorption, 700 and 709.5 eV for Fe L-edge adsorption, and 1550.0 and 1567.0 eV for Al K-edge adsorption (Solomon et al., 2012). The spatial resolution of the images was 200 nm. Deposited materials inevitably have sample thickness variation. Interpretation of the spectral results therefore requires cautions because the obtained signal from each pixel is intensified in the thicker region of the deposited sample (Wan et al., 2007).

2.6 Microbial community analyses

Bacteria may exert stronger control on rock weathering than fungi under our short-term conditions with repeated physical mixing. We thus assessed the changes in bacterial community 16S amplicon sequencing using the initial leaf compost ($n = 3$, air-dry conditions), bulk samples from all four treatments ($n = 3$, air-dry conditions), and MF from selected treatments (granite, coarse basalt, and fine basalt) ($n = 1$, wet conditions) after the incubation. The latter was a composite of three reps due to the limited mass recovery of MF. The DNA was extracted from 0.4 g of each sample by the FastDNA[®] SPIN Kit for Soil (MP Biomedicals, California, USA) according to the manufacturer's instructions with a few modifications. Quantitative PCR (qPCR) assays were conducted using the fluorescent dye SYBR Green (THUNDERBIRD Next SYBR qPCR mix, Toyobo) by a QuantStudio 3.0 real-time PCR System (Applied Biosystems/Thermo Fisher Scientific). 16S rRNA genes quantified using the primer pairs Bact1369F/ProK1492R (Suzuki et al., 2000) for the V3-V4 region of the 16S rRNA. The PCR reactions for 16S rRNA started with an initial denaturing step at 95 °C for 30 s, followed by 40 cycles at 95 °C for 5 s and 60 °C for 30 s. Melting curve analyses involved a denaturing step at 95 °C for 15 s, annealing at 65 °C for 1 min, and melting in 0.1 °C steps up to 95 °C. Standard curves for each assay were generated by serial dilutions of linearized plasmids with cloned fragments of environmental DNA. Amplification efficiencies were 96.4%. 16S rRNA amplicon sequencing and subsequent bioinformatics analysis are the same as the previous studies (Bamba et al., 2024; Hara et al., 2024). DNA sequencing data (16S rRNA gene amplicon) are available at the NCBI Sequence Read Archive (SRA) under BioProject ID PRJDB18777. For the 16S rRNA amplicon, taxonomy was assigned to ASVs using the SINTAX algorithm (Edgar, 2016) implemented in USEARCH (v11.0.667) against the RDP database v18 (Cole et al., 2014).

2.7 Statistics and calculations

The effects of rock treatment on the measured variables were tested by one-way ANOVA, followed by a Tukey's post hoc

test. The changes in variables before and after the incubation were evaluated using paired *t*-tests. All statistical analyses were performed in Microsoft Excel for Microsoft 365 MSO and Excel Toukei (BellCurve for Excel, Social Survey Research Information, Tokyo, Japan).

We estimated the total amounts of extractable metals in the rock-OM mixtures after the incubation by summing up the metals in LF, MF, and HF instead of the bulk sample values because we expected less sub-sampling error from the former. Specifically, roughly 44% of the mass of the incubated mixture was used for the density fractionation analysis, whereas only 13% was allocated for bulk sample chemistry. The high mass recovery of the density fractions (range: 99%–101%, Table 1) suggests that this approach was appropriate.

Alpha and beta diversities of the bacterial communities in the initial leaf compost and post-incubation samples were calculated using the “phyloseq” pipeline (McMurdie and Holmes, 2013) and the “microeco” pipeline (Liu et al., 2021), respectively. Principal coordinate analysis (PCoA) of unweighted UniFrac distance was used to visualize beta diversity between the treatments. We estimated the influence of the mineral type in the rock-OM mixtures on the beta diversity by permutational multivariate analysis of variance (PERMANOVA) using the “adonis” function in the “vegan” package (Oksanen, 2013). We further analyzed the correlations between the bacterial composition and the property variables of MF and bulk samples on Day 55 and initial compost by the Mantel test using the “mantel” function in the “ecodist” package (Goslee and Urban, 2007). All the microbial analyses were performed using R version 4.2.0 (<https://www.r-project.org>, last access: 1 April 2026).

We presented and discussed the mass, C and N, and extractable metal data of bulk and density fractions in multiple ways (e.g., per fraction vs. per bulk mass basis). Tables and figures of these data are summarized in Table S7.

3 Results

3.1 Recovery after density fractionation

Density fractionation of the rock-OM mixtures at the end of the 55 d incubation experiment showed reasonable recoveries for all the rock treatments: 99%–101% for mass, 86%–107% for C, 90%–109% for N, 88%–95% for extractable Al, 99%–108% for extractable Fe, and 85%–93% for extractable Si (Table 1). Metal recovery data is not available for granite and sand treatment as the metal extraction of LF and MF was not possible due to the limited mass. Some variations among the rock treatments were found. The coarse basalt treatment had less complete C and N recoveries than the other treatments and less complete extractable Al and Si recoveries relative to the fine basalt treatment. Overall, we considered that these recoveries were at acceptable levels

Table 1. Recovery of mass, C, N, and extractable Al, Fe, and Si after the density fractionation of the rock-OM mixtures on Day 55.

	Mass	C	N	Extractable Al	Extractable Fe	Extractable Si
	(%)	(%)	(%)	(%)	(%)	(%)
Granite 38–75 μm	100 \pm < 1	95 \pm 12	100 \pm 6	N.A.	N.A.	N.A.
Basalt 38–75 μm	99 \pm 1	86 \pm 8	90 \pm 8	88 \pm 7	99 \pm 6	85 \pm 3
Basalt 20–38 μm	100 \pm < 1	102 \pm 1	106 \pm 1	95 \pm 3	108 \pm 1	93 \pm < 1
Sand 100–300 μm	101 \pm < 1	107 \pm 14	109 \pm 18	N.A.	N.A.	N.A.

Al, Fe, and Si were extracted sequentially with sodium pyrophosphate, acid oxalate, and dithionite-citrate reagents. Value shows mean \pm standard deviation ($n = 3$). N.A. Not analyzed due to the limited mass recovery of the meso-density fraction.

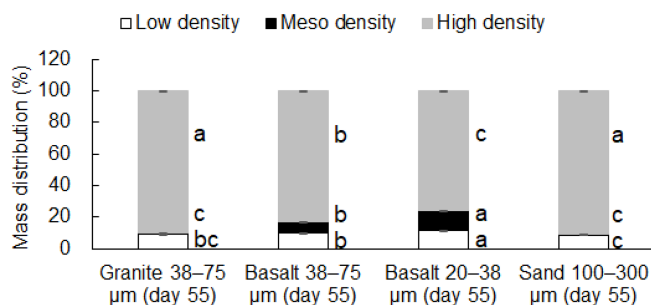


Figure 2. Mass distributions of the four rock-OM mixtures across low-, meso-, and high-density fractions on Day 55. For each density fraction (i.e., low-, meso-, or high-density), significant differences among the four rock treatments were shown by different letters at $P < 0.05$ (Tukey's test; $n = 3$).

for further assessment of the elemental distribution patterns across the density fractions.

3.2 Mass, C and N, and $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$

Mass proportion in MF after the incubation was the highest in the fine basalt (13 %), followed by the coarse basalt (7.1 %), granite (0.57 %), and sand (0.17 %) treatments (Fig. 2). The increases in MF largely corresponded to the reduction in the mass of HF. However, the initial granite, coarse basalt, and fine basalt contained small amounts of MF, accounting for 0.30 %, 6.8 %, and 6.3 % of the initial mass of respective minerals (Fig. 3a). After accounting for the initial MF in these treatments, we detected a significant increase in the mass of MF after the incubation for the granite and fine basalt treatments (Fig. 3a). The largest increase in MF mass was found in the fine basalt treatment (65 mg g^{-1} bulk) whereas that in the coarse basalt and sand treatments changed little ($0.26\text{--}3.0 \text{ mg g}^{-1}$ bulk) over the 55 d duration.

The C concentration in LF on Day 0, the initial leaf compost, was 408 mg g^{-1} fraction (Table S8-1), whereas that on Day 55 ranged from 378 to 431 mg g^{-1} fraction (Table S9-1). The C distribution to LF differed among the treatments in the following order: fine basalt > coarse basalt > granite > sand (Table S9-2). The same pattern was found for MF and HF. Because the initial crushed rocks contained only small

amounts of MF (Fig. 3a, Table S6) that had only trace level of C (Fig. 3b; Tables S8-1, S8-2), the C increase in MF after the incubation was significant for all four treatments ($P < 0.01$, Fig. 3b). The largest increase was shown in fine basalt ($1.50 \pm 0.05 \text{ mg C g}^{-1}$ bulk) followed by the coarse basalt ($1.03 \pm 0.07 \text{ mg C g}^{-1}$ bulk) and then the granite ($0.60 \pm 0.06 \text{ mg C g}^{-1}$ bulk) treatments. Even in the sand treatment, we detected a significant increase in the amount of C in MF ($0.10 \pm 0.01 \text{ mg C g}^{-1}$ bulk). When summing up the C and N in all three fractions, the total amounts of C and N left after the incubation showed a significant difference among the treatments in the following order: fine basalt > coarse basalt > granite > sand (Table S9-2).

The C : N ratio of the initial leaf compost and that of LF after the incubation were similar in all four treatments (Fig. 4a, Table S9-1) because the majority of C was present in LF. Among the density fractions, the C : N ratio progressively declined with increasing density. In other words, more C was lost relative to N from LF towards HF during the incubation. The isotopic values ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) of initial leaf compost and those of LF after the incubation were similar in both basalt treatments (Fig. 4b, c). The value of $\delta^{13}\text{C}$ significantly increased from LF towards HF, whereas $\delta^{15}\text{N}$ showed a significant increase from LF to MF and a slight decline from MF to HF (Fig. 4b, c).

3.3 Extractable Al, Fe, and Si and pH

Initial sand contained only negligible amounts of OX-extractable metals (metal_{OX}), and initial granite contained $< 1 \text{ mg g}^{-1}$ bulk of metal_{OX} , whereas initial coarse and fine basalts contained much greater extractable metals (Table S2). Between the two basalt size treatments, fine basalt had a higher amount of PP-extractable Al, Fe, and Si (Al_{PP} , Fe_{PP} , Si_{PP}) as well as Al_{OX} , Fe_{OX} , and Si_{OX} compared to coarse basalt.

After the 55 d incubation, the metal concentrations in LF consistently increased for Fe_{OX} , Fe_{DC} , Fe_{PP} , and Al_{PP} among the four rock treatments, but especially for the basalt treatments (Tables 2, S10-1). For the basalt treatments where enough MF mass recovery allowed the metal extractions (Fig. 2), a major increase in metal concentration was shown

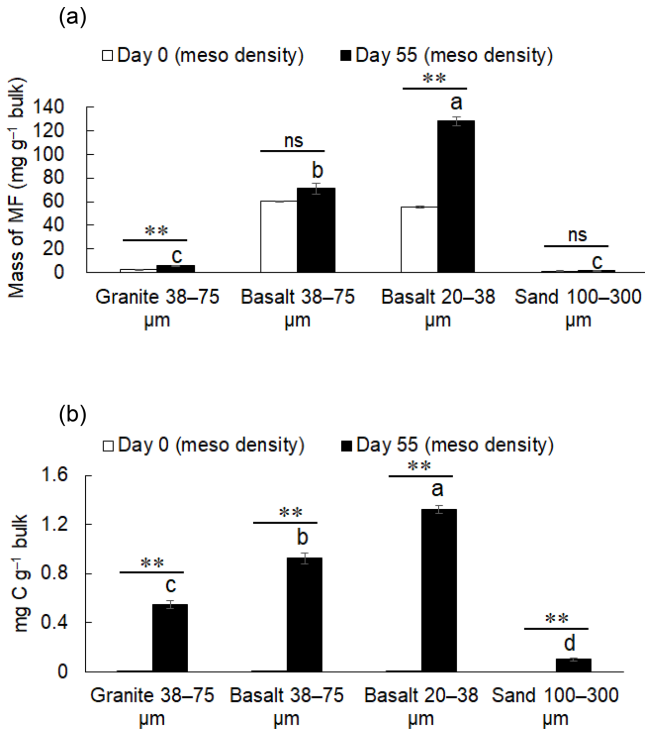


Figure 3. The mass of meso-density fraction (MF) in the rock-OM mixtures before (Day 0) and after the incubation (Day 55) (a). The amounts of C in MF in the bulk mixtures on Day 0 and Day 55 (b). Significant differences among the rock treatments on Day 55 are shown by different letters at $P < 0.05$ (Tukey's test; $n = 3$). The increase in MF from Day 0 to Day 55 is shown with asterisks (** significance at $P < 0.01$; ns, not significant).

in Fe_{OX} , Al_{OX} , Si_{DC} , and Al_{PP} for fine basalt, and Si_{DC} and Al_{PP} for coarse basalt. In HF, major changes were the decline in Fe_{OX} and Al_{OX} concentration for coarse basalt (Tables 2; S10-1).

The distribution of these metal phases across the density fractions, accounting for their masses, showed that the initial rock-OM mixtures held most of the extractable metal phases in HF (Table S10-2) and that the incubation led to enrichment of Fe, Al, Si, especially oxalate-extractable phases, in MF for the two basalt treatments (Table S11). Overall, the distribution of total extractable Fe in MF changed from 11%–12% to 16%–27% of the respective bulk samples (i.e., LF+MF+HF at Day 0 and 55) (Tables S10-2, S11). Similarly, total extractable Al also increased from 15%–19% to 23%–38% and Si showed a similar pattern. Among the three extractions, major increases over the incubation occurred in Fe_{OX} , Al_{OX} , and Si_{OX} phases in MF particularly for fine basalt treatment, which was in part balanced by their decreases in HF (Tables S10-2, S11). Similar but weaker trends were shown for Fe_{DC} , Al_{DC} , and Si_{DC} phases, while the changes in pyrophosphate-extractable phases remain minor. On a bulk sample basis, total extractable Fe in MF increased from 3.6–4.2 to 4.9–11 $mg\ g^{-1}$ and Al from 0.88–1.0

Table 2. Concentrations ($mg\ g^{-1}$ per fraction) of Al, Fe, and Si dissolved by sequential extractions in each density fraction from the rock-OM mixtures on Day 55.

	Sodium pyrophosphate (PP)				Acid oxalate (OX)				Dithionite-citrate (DC)				Total												
	Al	Fe	Si		Al	Fe	Si		Al	Fe	Si	Al	Fe	Si											
Low-density fraction ($< 1.8\ g\ cm^{-3}$)																									
Granite 38–75 μm	0.33	± 0.03	0.25	$\pm < 0.01$	3.0	± 0.1	1.0	$\pm < 0.1$	1.7	$\pm < 0.1$	0.51	± 0.01	0.30	± 0.03	1.5	± 0.1	2.7	± 0.2	1.6	± 0.1	3.4	± 0.1	6.2	± 0.3	
Basalt 38–75 μm	0.48	± 0.01	0.53	± 0.02	2.0	± 0.1	1.4	± 0.1	3.3	± 0.1	0.67	± 0.02	0.37	± 0.01	2.1	$\pm < 0.1$	4.0	± 0.1	2.2	± 0.1	5.9	± 0.1	6.6	± 0.1	
Basalt 20–38 μm	0.52	± 0.02	0.71	± 0.02	1.6	$\pm < 0.1$	2.0	± 0.1	6.3	± 0.1	1.2	$\pm < 0.1$	0.48	± 0.01	3.3	± 0.1	6.4	± 0.1	3.0	$\pm < 0.1$	10	$\pm < 1$	9.2	± 0.1	
Sand 100–300 μm	0.39	± 0.01	0.24	$\pm < 0.01$	3.1	± 0.2	1.1	± 0.1	1.6	$\pm < 0.1$	0.49	± 0.03	0.26	± 0.04	1.1	± 0.1	2.4	± 0.3	1.7	± 0.1	3.0	± 0.1	6.0	± 0.4	
Meso-density fraction ($1.8\text{--}2.4\ g\ cm^{-3}$)																									
Granite 38–75 μm	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.
Basalt 38–75 μm	0.44	± 0.01	0.76	± 0.04	1.7	$\pm < 0.1$	13	± 3	51	± 11	6.1	± 1.3	2.7	± 0.1	16	± 1	9.6	± 0.2	17	± 3	68	± 11	17	± 1	
Basalt 20–38 μm	0.44	$\pm < 0.01$	0.69	± 0.02	1.7	$\pm < 0.1$	18	± 1	74	± 1	9.4	± 0.2	2.4	± 0.1	14	$\pm < 1$	9.2	± 0.1	21	± 1	89	± 1	20	$\pm < 1$	
Sand 100–300 μm	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.
High-density fraction ($> 2.4\ g\ cm^{-3}$)																									
Granite 38–75 μm	0.0	$\pm < 0.1$	0.0	$\pm < 0.1$	0.0	$\pm < 0.1$	0.12	± 0.02	0.68	± 0.04	0.13	± 0.02	0.18	± 0.01	1.8	± 0.0	0.73	± 0.02	0.30	± 0.03	2.5	± 0.1	0.86	± 0.04	
Basalt 38–75 μm	0.071	± 0.011	0.23	± 0.01	0.36	± 0.02	3.7	± 0.1	25	± 1	2.2	± 0.1	0.72	± 0.04	4.8	± 0.3	8.9	± 0.2	4.5	± 0.1	30	± 1	11	$\pm < 1$	
Basalt 20–38 μm	0.15	$\pm < 0.01$	0.28	$\pm < 0.01$	0.57	± 0.02	4.7	± 0.1	34	± 1	3.1	$\pm < 0.1$	0.59	± 0.05	4.9	± 0.6	9.2	± 0.1	5.4	± 0.1	39	± 1	13	$\pm < 1$	
Sand 100–300 μm	0.0	$\pm < 0.1$	0.0	$\pm < 0.1$	0.0	$\pm < 0.1$	0.015	± 0.008	0.14	± 0.10	0.0	$\pm < 0.1$	0.087	± 0.014	1.3	± 0.3	0.080	± 0.021	0.10	± 0.01	1.4	± 0.4	0.080	± 0.021	

Al, Fe, and Si were extracted sequentially with sodium pyrophosphate, acid oxalate, and dithionite-citrate reagents. Value shows mean \pm standard deviation ($n = 3$). N.A. = Not analyzed due to the limited mass recovery of the meso-density fraction. Concentrations (per fraction) on Day 0 are represented in Table S10-1.

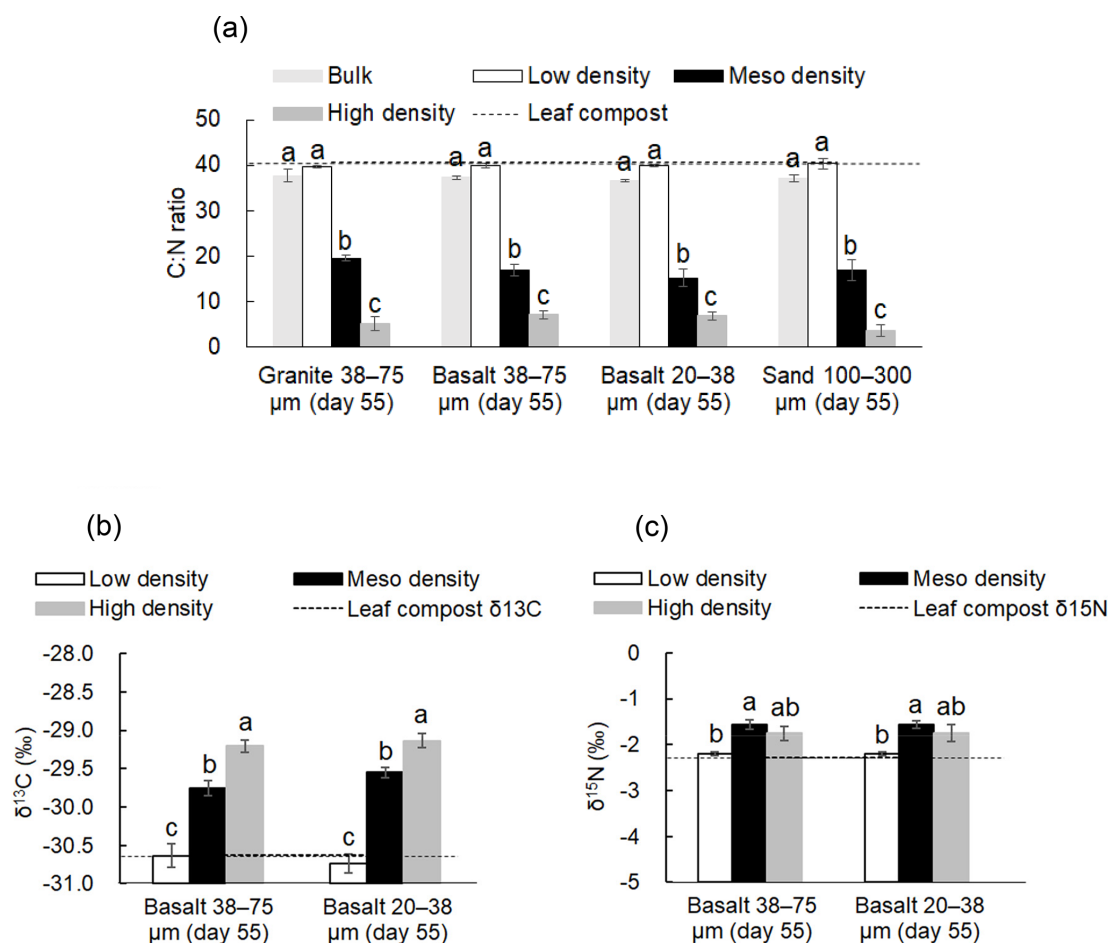


Figure 4. C : N ratio (a), $\delta^{13}\text{C}$ (b), and $\delta^{15}\text{N}$ (c) among the density fractions from the four rock treatments on Day 55 and those of initial leaf compost as a horizontal dotted line. No $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ analyses were done for granite and sand treatment due to the limited mass of MF. Significant differences among the density fractions are shown with different letters at $P < 0.05$ (Tukey's test; $n = 3$).

to 1.2–2.7 mg g^{-1} during the incubation. Concurrently, the total extractable Fe in HF slightly decreased from 28–30 to 25–29 mg g^{-1} and Al from 4.2–4.9 to 3.7–4.1 mg g^{-1} during the incubation. These changes mainly occurred in oxalate-extractable phases: Al_{OX} and Fe_{OX} in MF after the incubation accounted for 82%–89% and 76%–86% of the total extractable metal pools, respectively (Table S11). As a result, MF showed statistically detectable increases (thus accumulation) of Al, Fe, and Si, especially in OX-extractable pools, for fine basalt (Fig. 5a–c), whereas a small but significant increase in Al_{PP} phase was shown for coarse basalt. Meso-density fraction also showed significant enrichment of Fe_{OX} for both basalt treatments and that of Al_{OX} for the coarse basalt treatment relative to Si_{OX} (Table 3). In addition, Fe_{OX} phase appeared to be enriched relative to Al_{OX} phase in fine basalt treatment. The carbon-to-extractable metal molar ratios ranged from 0.5 to 2.3 (Table S12).

When summing up the total extractable Al and Fe among the three fractions, a significant increase after the incubation period was detected for Fe (7.2 mg g^{-1} bulk) and Al

(1.7 mg g^{-1} bulk) only in fine basalt treatment (Tables S10–2, S11). Thus, the observed changes in the extractable metals among the fractions likely resulted from both basalt weathering and redistribution in fine basalt and only from the redistribution in the other treatments.

The $\text{pH}(\text{H}_2\text{O})$ of the bulk samples of all treatments on Day 55 ranged between 6.5 and 7.6, with the highest pH in the fine basalt treatment, followed by the coarse basalt, while the granite and sand treatments were slightly acidic (Table 4). The difference in $\text{pH}(\text{KCl})$ among the treatments was less clear.

3.4 Leaching loss of elements and leachate chemistry

The total amount of DOC released from the rock-OM mixtures during the eight repeated leaching events was 13–15 mg C per incubation column (Table 5), which was equivalent to 1.3%–1.5% of the total C in the initial mixtures. No significant difference in DOC release was found among the four treatments. For the major rock-forming elements, Fe

Table 3. Molar ratios of acid oxalate extractable Al, Fe, and Si in meso-density fraction in the two basalt treatments before and after the 55 d incubation.

		Meso-density fraction (1.8–2.4 g cm ⁻³)								
		Al : Si		Fe : Si		Fe : Al				
Basalt 38–75 μm	Day 0	2.0	± < 0.1	3.5	± < 0.1	1.8	± < 0.1			
	Day 55	2.3	± < 0.1	**	4.2	± 0.1	**	1.8	± < 0.1	ns
Basalt 20–38 μm	Day 0	2.0	± < 0.1		3.6	± < 0.1		1.8	± < 0.1	
	Day 55	2.0	± < 0.1	ns	4.0	± < 0.1	**	2.0	± < 0.1	**

Value shows mean ± standard deviation (*n* = 3). The increase in molar ratio from Day 0 to Day 55 is shown with asterisks (** significance at *P* < 0.01; ns, not significant, *t*-test).

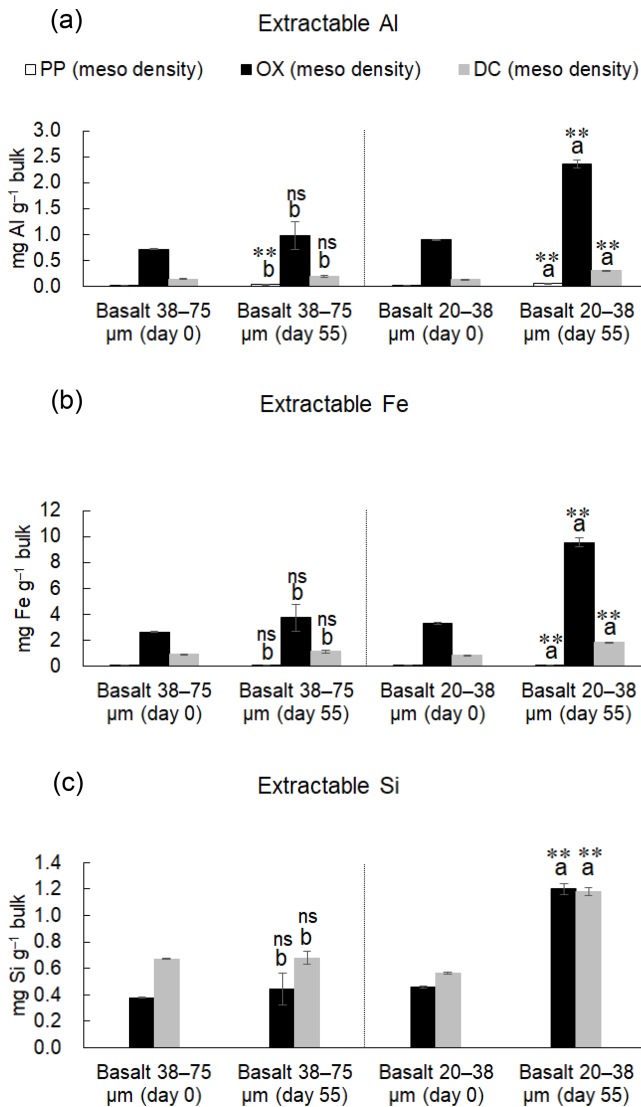


Figure 5. Amounts of extractable Al (a), Fe (b), and Si (c) in MF from the rock-OM mixtures on Day 0 and Day 55. See Fig. 3 legend for statistical difference signs.

Table 4. pH of the rock-OM mixtures after the 55 d incubation.

	pH(H ₂ O)		pH(KCl)			
Granite 38–75 μm	6.7	± < 0.1	c	6.1	± < 0.1	a
Basalt 38–75 μm	7.3	± < 0.1	b	5.9	± < 0.1	bc
Basalt 20–38 μm	7.6	± < 0.1	a	6.0	± < 0.1	ab
Sand 100–300 μm	6.5	± < 0.1	d	5.8	± < 0.1	c

Value shows mean ± standard deviation (*n* = 3). Significant differences among the rock treatments on Day 55 are shown by different letters at *P* < 0.05 (Tukey’s test; *n* = 3).

leaching was the highest in the fine basalt, followed by the coarse basalt treatment, and Si leaching showed similar patterns (Table 5). Conversely, the total amount of Al leached was the highest in the sand and the lowest in the coarse basalt treatment. The sum of base cations leached was the highest in the fine basalt treatment, followed by the coarse basalt, sand, and granite treatments. Leaching of base cations during the incubation followed the order: Ca > Na > K > Mg. Greater leaching of Na from fine basalt relative to coarse basalt was consistent with (Yang et al., 2026), which likely indicates rapid dissolution of Na plagioclase from the basaltic rock. The pH of the leachates was neutral to alkaline (7.1–9.1), with the highest pH in the fine basalt, followed by the coarse basalt treatment, while the other two treatments had nearly neutral pH (Table 4). The observed pH difference largely corresponded to the variation of the leachate chemistry among the treatments (e.g., Na, Ca, and Al).

3.5 Microscopic observations of organo-mineral assemblage by SEM and STXM-NEXAFS

We observed some shaking-resistant, organo-mineral assemblages in the post-incubation MF (Fig. 6). The observed assemblages were present as microaggregates of 100–200 nm in diameter, consisting of crushed rocks (Fig. 6b, d, f) and OM, including fungal hyphae (Fig. 6d). Greater aggregate sizes were apparent in the two basalt treatments compared to the granite treatment (Fig. 6d, f). Much finer subunits (rock

Table 5. Amounts of DOC, Al, Fe, Si, base cations, and pH in leachates released from the rock-OM mixtures during the 55 d incubation (the sum of the eight repeated leaching events).

	DOC			Al			Fe			Si		
	(mg)			(mg)			(mg)			(mg)		
Granite 38–75 μm	14	± 1	a	0.0068	± 0.0019	ab	0.015	± 0.001	c	1.5	± 0.1	b
Basalt 38–75 μm	13	$\pm < 1$	a	0.0044	± 0.0007	b	0.028	± 0.001	b	3.5	$\pm < 0.1$	a
Basalt 20–38 μm	15	$\pm < 1$	a	0.0060	± 0.0009	ab	0.036	$\pm < 0.001$	a	4.5	± 0.2	a
Sand 100–300 μm	13	± 1	a	0.010	± 0.001	a	0.013	$\pm < 0.001$	c	1.7	± 0.4	b

	Sum of base cations			Na	Ca	K	Mg	pH						
	(mg)			(mg)	(mg)	(mg)	(mg)							
Granite 38–75 μm	7.7	± 0.3	c	1.6	$\pm < 0.1$	4.2	± 0.2	0.94	± 0.06	1.0	± 0.1	7.2	$\pm < 0.1$	c
Basalt 38–75 μm	9.8	± 0.3	b	2.0	± 0.1	5.5	± 0.2	1.7	$\pm < 0.1$	0.62	± 0.04	7.9	± 0.1	b
Basalt 20–38 μm	14	$\pm < 1$	a	6.1	± 0.2	5.4	± 0.1	1.6	$\pm < 0.1$	0.49	± 0.02	9.1	$\pm < 0.1$	a
Sand 100–300 μm	8.9	± 0.7	bc	1.2	± 0.2	3.3	± 0.4	3.3	± 0.3	1.1	± 0.1	7.1	± 0.1	c

Value shows mean \pm standard deviation ($n = 3$). Significant differences among the rock treatments on Day 55 are shown by different letters at $P < 0.05$ (Tukey's test; $n = 3$).

fragments) were confirmed in the fine basalt aggregates compared to the coarse basalt aggregates.

The subsamples of dispersed MF material from the selected rock-OM mixtures, the granite and coarse basalt treatments (see Fig. S4 for SEM images), were further characterized to assess the chemical nature of the organo-mineral assemblages by STXM-NEXAFS (Fig. 7). We first obtained the spatial distribution of C, Al, and Fe for a large area ($35 \times 35 \mu\text{m}$, Fig. 7A, B). As the C-rich zones in the targeted assemblage were patchy, we selected one of these patches and the adjacent mineral-rich zones as a focused area ($5 \times 5 \mu\text{m}$) to examine the distribution of C functional groups (Fig. 7a, b). In the granite sample, the C-rich zone (shown as ROI in Fig. 7a) was relatively enriched in aromatic and carboxylic C compared to the average C K-edge NEXAFS spectra of the entire $5 \times 5 \mu\text{m}$ area (Fig. 7a1, a3). While some overlaps of C with Fe and Al (Fig. 7a4, a5) were shown, the metal-rich patches appeared to be present adjacent to, rather than overlapped with, the C-rich zone. The dominance of Fe over Al shown in the coarse basalt treatment (Fig. 7B) was in line with the higher content and increase of Fe_{OX} relative to Al_{OX} phase (Fig. 5a, b). In the coarse basalt sample, the C-rich zone (ROI in Fig. 7b) was enriched in ketonic C in contrast with the granite sample (Fig. 7b2). Aromatic C and carboxylic C appeared to be distributed separately (Fig. 7b1, b3). While the three major C peaks were found in MF from both granite and coarse basalt, the latter appeared to have a higher relative abundance of ketonic C (Fig. 7, middle left spectra).

3.6 Bacterial community composition after the 55 d incubation

We assessed the effect of the rock treatment on the microbial community composition at the end of the incubation using

bacterial amplicon sequencing. The bacterial copy number ranged from 3.7×10^8 to 1.5×10^9 , and the initial compost tended to have a higher number than the bulk samples after the incubation (Table S13). Among the post-incubation bulk samples, the bacterial number decreased in the following order: sand > granite > two basalt treatments, with no significant difference among the treatments (ANOVA, $P = 0.08$). The alpha diversity indices, including Chao1, Shannon index, and evenness, showed no significant differences among the treatments (Table S13; ANOVA, $P = 0.28, 0.24, 0.10$, respectively). Notably, the beta diversity (PCoA plot; Fig. 8a) was different among the treatments (PERMANOVA, $F = 5.63$, $P = 0.001$), while the top three phyla – Proteobacteria, Actinobacteria, and Firmicutes – were commonly observed across the samples (Fig. 8b). The sum of the relative abundance of the three ASVs belonging to *Neobacillus* (Fig. 8c), the most dominant genus, was higher in MF compared to the bulk sample ($15.5 \pm 8.4\%$ and $3.5 \pm 1.7\%$, respectively, t -test, $P = 0.0036$). The mantel results using the data of MF and the bulk samples on Day 55 further showed significant correlations between the microbial composition (PCoA score) and selected soil properties including total C, total N, $\text{pH}(\text{KCl})$, $\text{pH}(\text{H}_2\text{O})$, Fe_{PP} , and Al_{PP} ($R^2 = 0.86, 0.86, 0.82, 0.80, 0.30$ and 0.22 respectively, $P < 0.05$; Table S14).

4 Discussion

4.1 Formation of organo-mineral assemblages via weathering-derived metal supply

We conducted a simple incubation experiment to test whether Fe and Al released from rock weathering promote the formation of organo-mineral assemblage, represented by the meso-density fraction (MF), and to assess the concurrent changes in both organic matter (OM) and rock-derived el-

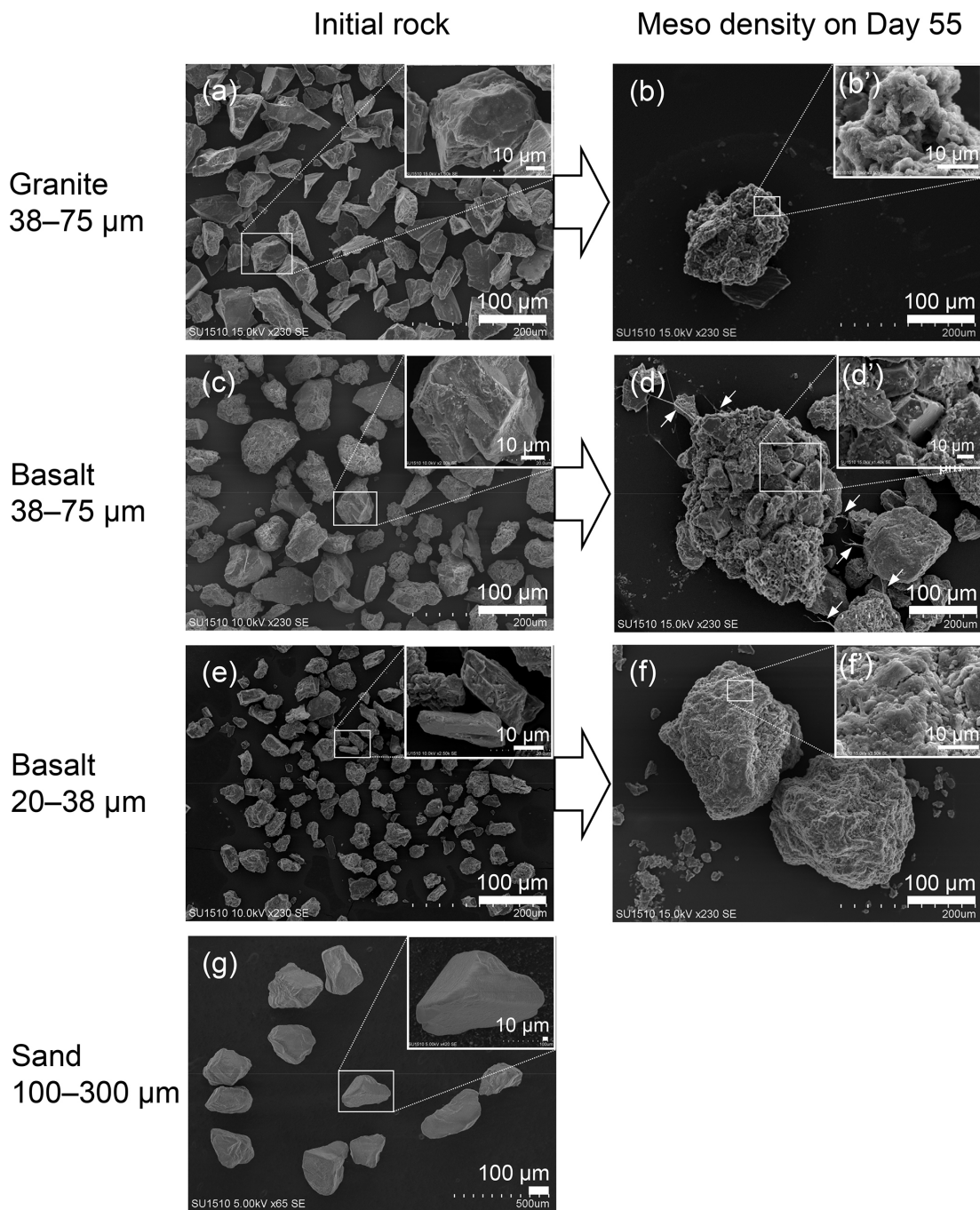


Figure 6. SEM images of the crushed rock minerals and the isolated density fraction. (a) initial granite, (b) MF of granite, (c) initial coarse basalt, (d) MF of coarse basalt, (e) initial fine basalt, (f) MF of fine basalt, and (g) initial sand. Fungal hyphae were shown with arrows (d).

ements. Among the three crushed rocks used, fine basalt showed the highest increases in MF mass (Fig. 3a), C content (Fig. 3b), and extractable metals and metalloids particularly FeO_x , AlO_x , and SiO_x (Fig. 5). Over the 55 d incubation period, C accumulation in MF was measured at $1.3 \pm 0.1 \text{ mg C g}^{-1}$ bulk for fine basalt, $0.9 \pm 0.1 \text{ mg C g}^{-1}$ bulk for coarse basalt, and $0.6 \pm 0.1 \text{ mg C g}^{-1}$ bulk for gran-

ite (Tables S8-2, S9-2). Extractable Fe, Al, and Si in MF also showed the same trend for the basalt treatments (Tables S10-2, S11). The concurrent enrichment of mineral-associated OM (MAOM) and extractable metals supports the idea that metal release from basalt weathering directly facilitates the formation of organo-mineral assemblages, consistent with the *organo-metallic glue* hypothesis (Wagai et al., 2020).

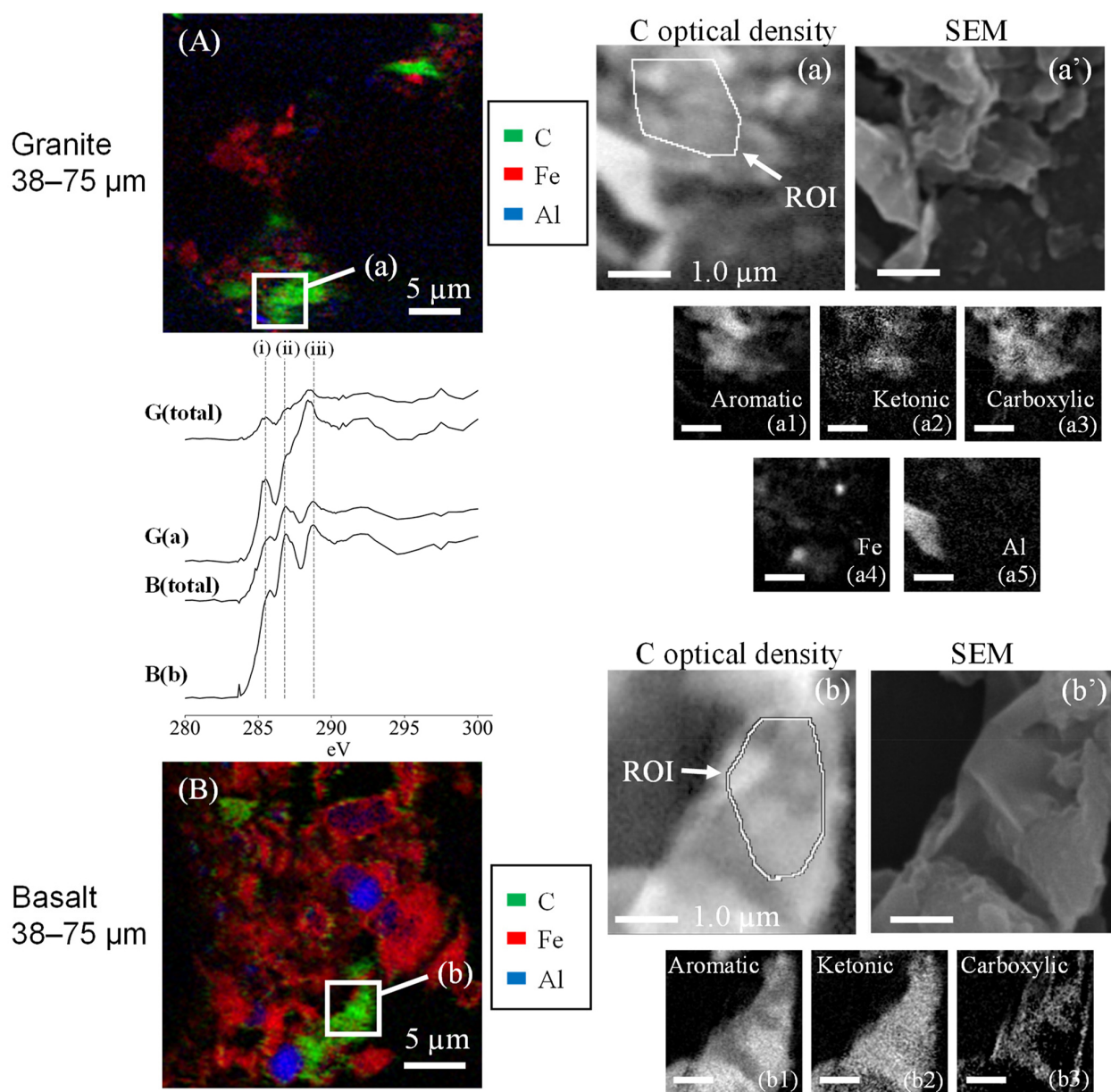


Figure 7. The elemental maps obtained with STXM-NEXAFS analysis. The images show the distribution of elements in the $35 \times 35 \mu\text{m}$ region of interest for MF of granite (A) and $30 \times 30 \mu\text{m}$ region of interest for coarse basalt treatments (B) after the 55 d incubation, respectively. C optical density (a, b) and SEM (a', b') images for a selected C-rich region at higher resolution ($5 \times 5 \mu\text{m}$ and $4 \times 4 \mu\text{m}$, respectively). The C K-edge NEXAFS spectra in the middle left panel show the image stacks for the $35 \times 35 \mu\text{m}$ region of granite, G(total), and coarse basalt, B(total), and for the C-rich regions of granite, G(a), and coarse basalt, B(b). Dashed lines (i), (ii), and (iii) represent the energy levels at 285.5, 286.8, and 288.8 eV corresponding to aromatic C, ketonic C, and carboxylic C. The images (a1), (a2), and (a3) and (b1), (b2), and (b3) show the spatial distribution of specific C type, aromatic C, ketonic C, and carboxyl C, respectively, in the higher resolution images. The distribution of Fe (a4) and Al (a5) was measured only for the granite sample due to beamtime limitations.

The hypothesis proposes that the assemblage (aggregation) occurs through organo-metallic binding agents that can form through organo-metal complexation and coprecipitation as well as dissolved OM sorption on secondary metal oxide phases. Identifying the sources for the metal enrichment in MF provides insight into the formation mecha-

nism. Mass balance of the extractable metals before and after the incubation is, however, quite challenging due to relatively high errors in metal extractions, especially when having multiple extractions and fractions. While our results are subject to these errors, we postulated potential dynamics of Fe and Al phases for the fine basalt treatment

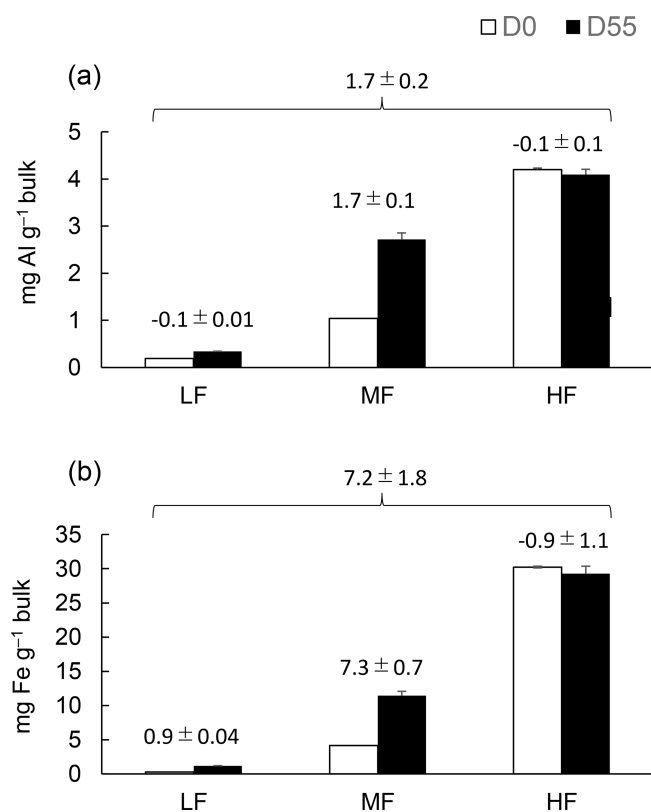


Figure 9. The changes in total extractable Al (a) and Fe (b) before and after the 55 d incubation in LF, MF, and HF on a bulk sample mass basis. Values \pm SD above the two columns represent the change from Day 0 to Day 55. The sum of the changes across LF, MF, and HF was shown on the top of the three fractions.

lease \rightarrow preferential association with OM \rightarrow aggregation – provides direct experimental support for the *organo-metallic glue* hypothesis, as depicted in Fig. 10.

4.2 Chemistry of the organo-mineral assemblages

Chemical characterization of MF provides further insights into the nature of the presumed *organo-metallic glue*. The dominance of Fe_{OX} and Al_{OX} relative to Fe_{PP} and Al_{PP} phases in the basalt treatments (Table S11) indicates that readily soluble organo-metal complexes were minor. The accreted OM in MF was thus more likely to be associated with Fe_{OX} and Al_{OX} phases through adsorptive or coprecipitation mechanisms. Microscopic observation of a representative MF microaggregate in the coarse basalt treatment (Fig. 7B) showed Fe dominance with patchy distributions of Al and C (rich in ketonic C). This pattern implies that Fe_{OX} phases (as inferred from Fig. 5b) might have entrapped small OM-rich domains during coprecipitation. Given that only a limited area was analyzed, broader generalization across all MF materials is not warranted.

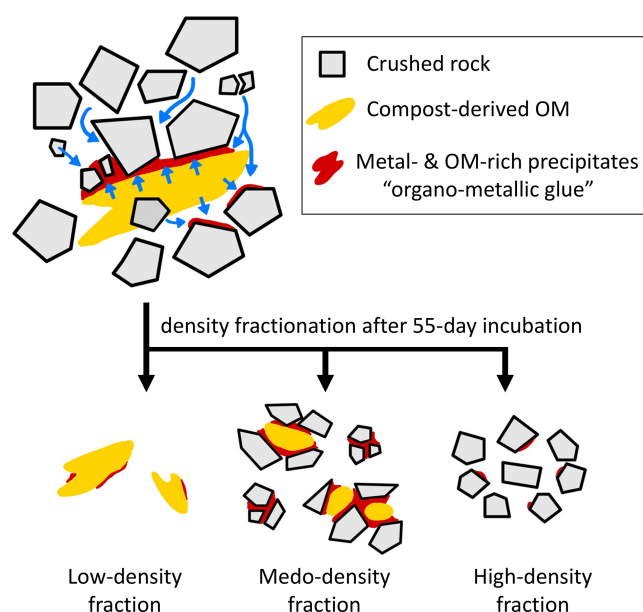


Figure 10. Conceptual synthesis of the experimental results in the context of the *organo-metallic glue* hypothesis. Metals (primarily Fe and Al) released from crushed rock under wet conditions preferentially associate with dissolved and particulate OM, forming organo-metallic precipitates on mineral and organic surfaces, particularly during drying. Following eight wet–dry cycles with physical mixing, these precipitates become enriched in the meso-density fraction, supporting their inferred role as binding agents that link mineral and organic particles and promote the formation of physically stable organo-mineral assemblages.

The C : (Fe+Al) molar ratio of newly-formed MF is consistent with the coprecipitate interpretation. For the fine basalt treatment, the ratio of 0.58 ± 0.03 (mol : mol), after accounting for the total extractable metals originally present in the basalt, falls within the range reported for laboratory-synthesized Fe-OM, Al-OM, and Fe-Al-Si-OM coprecipitates (Kleber et al., 2015; Tamrat et al., 2019; Jamoteau et al., 2023). This ratio was substantially lower than those for MF in natural soils (Wagai et al., 2020) and for organo-metal complexes typical of Andisols and spodic horizons (Takahashi and Dahlgren, 2016; Lundström et al., 2000), suggesting that the MF formed in our experiment contained relatively metal-rich, OM-poor precipitates. The relatively low C : metal ratios coincide with enrichment of Fe_{OX} and, to some extent, Al_{OX} relative to Si_{OX} (Table 3), implying a greater role of Fe_{OX} phase for the organo-mineral assemblage formation. Assuming that the weathering-derived Fe_{OX} phase (Fig. 5b) consists entirely of ferrihydrite, the accreted OM in MF corresponds to 108 ± 6 mg C g⁻¹ oxide – a value comparable to the maximum sorptive capacity of ferrihydrite for dissolved OM under laboratory conditions (Tipping, 1981; Kaiser et al., 1997). This result suggests that the surface of Fe_{OX} mineral phase was nearly saturated by OM.

The metal- and OM-rich phases observed in MF may reflect either coprecipitation of metals with organic ligands or the spatial co-occurrence of metal oxides and OM (e.g., OM sorption on precipitated oxides or metal precipitation on OM surfaces). Because these two mechanisms cannot be distinguished with the available data, we collectively refer to the resulting phases as organo-metallic precipitates (Fig. 10). We made a first-order approximation of the organo-metallic precipitates, recognizing that the calculation relies on simplified assumptions that Fe and Al occurred as ferrihydrite and allophane and OM consisted of 50 % C. This component accounted for approximately a quarter of the total MF mass, suggesting that resultant Fe-Al-OM precipitates may be abundant enough to link the remaining solids (mainly basalt particles with minor POM) to form stable organo-mineral assemblages (Fig. 10). In addition, other major elements released from rocks (particularly Si and Ca) may synergistically contribute to the assemblage formation (Oades, 1988; Jamoteau et al., 2025).

4.3 Organic matter dynamics during the assemblage formation

Several lines of evidence point to preferential accretion of microbially-processed, N-rich OM during organo-mineral assemblage formation. First, the C : N ratio of OM in MF sample was significantly lower than that in LF in all four rock-OM mixtures on Day 55 (Fig. 4a), suggesting a greater contribution of microbially altered material in MF. Second, MF had higher $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values (+0.88‰–1.2‰ and +0.63‰–0.64‰, respectively) relative to LF in basalt treatments (Fig. 4b, c). The negative $\delta^{15}\text{N}$ values observed are within the common range for temperate forest leaves ($-2.8 \pm 2.0\%$; Martinelli et al., 1999) and for organic layers of forest soils in the temperate climate zone ($-0.3 \pm 0.3\%$; Choi et al., 2020), which likely reflects tree-leaf origin of the OM used in our study. Leaf compost itself contains a variety of plant- and microbially-derived compounds. Thus, the observed isotopic enrichment in MF cannot be fully attributed to microbial reactions that took place during the incubation. Our results are, nonetheless, consistent with previous field studies showing the increase in $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ with soil particle density (Sollins et al., 2009; Hatton et al., 2012; Moni et al., 2012). Importantly, HF contained small amounts of even stronger stoichiometric (Table S9-1, Fig. 4a) and isotopic (Fig. 4b) signatures than MF. These results suggest that some of decomposing POM fragments were likely occluded in MF, while HF primarily retained adsorbed microbial OM. The role of POM in MF formation aligns with its well-known role as hotspots for microbial activity and dissolved OM production, which may induce localized dissolution of minerals (e.g., Bölscher et al., 2025).

These findings are in line with the conceptual framework previously proposed (Fig. 7b in Wagai et al., 2020). STXM-NEXAFS spectra (Fig. 7) further suggest that C spectra of

MF resemble those of typical soil OM (Solomon et al., 2012; Chen et al., 2014; Prietzel et al., 2018), which is dominated by microbially-processed OM in mineral soils (Sollins et al., 2009; Heckman et al., 2022). Our incubation conditions (wet-dry cycles, high temperature, and inoculation at each cycle) likely promoted microbial turnover and supply of microbial residues such as cell wall fragments and extracellular polymers (Krause et al., 2019).

These results collectively indicate that organo-mineral assemblages formed in MF were enriched in microbially-processed, N-rich OM, linking microbial activity, chemical weathering, and aggregation. Microbes likely accelerated mineral weathering via disaggregation, hydration, and localized dissolution (Banfield et al., 1999; Burgehelea et al., 2015; Finlay et al., 2020), releasing Fe and Al that subsequently coprecipitate with, or were sorptively associated with, microbially derived compounds on the surface of cells or biofilms as low-crystallinity Al, Fe, Si mineral phases (Lybrand et al., 2019). This process may have created reactive Fe–Al–OM phases that acted as nucleation sites for aggregation. The observed bacterial community patterns may be linked to these processes. While the community composition differed markedly among the rock treatments, bacterial copy numbers tended to be lower in the basaltic treatments (Table S13) potentially reflecting the protective role of the extractable metal phases. The genus *Neobacillus*, known for its biomineralization potential (Farda et al., 2022; Bhattacharjee et al., 2023), appeared to be enriched in MF relative to the bulk fraction (Fig. 8c), supporting the idea of MF as a mineralosphere (Uroz et al., 2015) where organo-metallic precipitates and microbial residues interact to drive aggregation.

4.4 Influence of the size and chemical composition of crushed rock

Both particle size and chemistry of the rock materials affected the formation of organo-mineral assemblages. Fine basalt treatment produced significantly more MF than coarse basalt or other treatments (Fig. 3a), emphasizing the importance of mineral particle size in regulating reactivity and aggregation. However, the size effect is not straightforward. Using the basaltic rock from the same source, Yang et al. (2026) observed less formation of Fe_{Ox} and Al_{Ox} phases from finer basalt due to greater occlusion of basalt particles within meso-density aggregates where water access and weathering are likely to be limited. Thus, multiple factors such as the degree of drying and mixing, OM decomposability, incubation duration, and basalt particle size likely interact to produce size-dependent effects. Even small differences in pore structure or redox dynamics can strongly influence weathering and aggregation, for instance, via affecting Fe-OM interactions (e.g., Chen et al., 2020).

Rock chemistry also influenced OM retention. Despite the similar size distribution (Fig. S2), coarse basalt held greater OM in MF than granite (Fig. 3b), which is attributable

to more than one order of magnitude higher initial metal contents, especially in Fe_{OX} , Al_{OX} , and Si_{OX} (Table S2). When normalized to surface area (Table S2), OM accretion in coarse basalt relative to granite corresponds to additional $\approx 0.1 \text{ mg C m}^{-2}$, which lies within sorptive limit of reactive secondary minerals (Wagai and Mayer, 2007; Schneider et al., 2010) and is comparable to C loadings on goethite under in temperate field conditions (Bramble et al., 2024). The strong influence of pre-existing extractable metals (i.e., secondary minerals) in basaltic rock on organo-mineral associations is consistent with findings from a similar incubation experiment (Yang et al., 2026). Although our study lacked fine granite treatment for full factorial comparison, the present results indicate that both particle size and mineralogy jointly regulate organo-mineral assemblage formation and OM stabilization.

4.5 Implications

Our experiment, conducted under simplified conditions, may provide mechanistic insights relevant to early-stage pedogenesis. Weathering of crushed rock in the presence of OM and microbial activity under fluctuating water regimes likely occurs during pedogenesis (e.g., primary succession on glacial tills). Although the wet–dry cycles applied here were more intense than typical field conditions, such processes can also occur in agricultural topsoils, where physical mixing and periodic drying can promote mineral–organic interactions. The application of crushed basalt and other mafic rocks to soils has drawn increasing attention as a strategy for CO_2 removal through enhanced weathering (Beerling et al., 2020). Besides CO_2 removal, several pot- and field-scale studies have reported increases in soil OM following mafic rock addition (Buss et al., 2024; Xu et al., 2024), although others found no response (Yan et al., 2023; Sokol et al., 2024). In the current study, we did not detect the significant change in total C possibly due to the short incubation period. Another factor is carbonate formation as the incubation was done under relatively high pH conditions. However, the consistent density-dependent changes in C : N, $\delta^{13}\text{C}$, and $\delta^{15}\text{N}$ (Fig. 4, Sect. 4.3 above) suggest that total C mainly consists of organic C. Regardless, our results indicated that the incorporating crushed basalt can promote the formation of stable organo-mineral assemblages via *organo-metallic glue* and thus potentially increase OM persistence. On the other hand, basalt-OM aggregation may reduce the rate of basalt weathering by shielding basalt mineral surfaces from further dissolution, highlighting a possible trade-off between OM stabilization and CO_2 removal efficiency (Yang et al., 2026). Further mechanistic studies are important to better elucidate rock-OM-secondary mineral interactions and to optimize C balance within enhanced rock-weathering strategies.

Across the rock treatments, the degree of OM accretion in MF (fine basalt > coarse basalt > granite > sand, Fig. 3b) corresponded closely to the abundance of reactive Fe_{OX} ,

Al_{OX} , and Si_{OX} phases from both inherent rock composition (Tables S2, S10-2) and weathering during incubation (Table S11), suggesting that reactive metal phases controlled OM retention more strongly than pH (Table 4) or bacterial abundance (Table S13). These results are consistent with the observations at a field scale (e.g., Fukumasu et al., 2021), regional to continental scales (Hughes, 1982; Ashida et al., 2021; von Fromm et al., 2021), up to a global scale (Rasmussen et al., 2018; von Fromm et al., 2025). The current study demonstrated that organo-metallic binding and aggregation can develop rapidly through weathering and microbial processes, directly linking mineral transformation to OM stabilization and highlighting the intimate interactions among microbes, their metabolites, and minerals in soil (Chorover, 2022; Fang et al., 2023; Wagai et al., 2023).

5 Conclusions

The formation of organo-mineral assemblages (meso-density fraction) from the mixture of crushed rocks and leaf compost was observed within 55 d incubation subjected to eight wet-dry cycles. The extent of assemblage formation followed the sequence: fine basalt > coarse basalt > granite \geq sand. Assemblage development was driven by two coupled processes: (i) microbial transformation of organic matter, evidenced by lower C : N and higher $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ relative to the initial leaf compost, and (ii) the supply of Fe and Al from the rock weathering. The resulting meso-density materials were enriched in organo-metallic precipitates with a dominance of oxalate-extractable Fe and, to a less extent, Al phases. These findings provide experimental support for the *organo-metallic glue* hypothesis (Wagai et al., 2020), demonstrating that weathering-derived metals can rapidly induce aggregation and OM persistence. This mechanism links microbial activity, mineral weathering, and OM stabilization, suggesting that early-stage pedogenesis and soil OM formation are strongly controlled by the coupled action of microbial metabolism and reactive metal dynamics.

Data availability. Data are available upon request.

Supplement. The supplement related to this article is available online at <https://doi.org/10.5194/soil-12-521-2026-supplement>.

Author contributions. KM: conceptualization, investigation, methodology, formal analysis, writing (original draft preparation, review, and editing); JJ: investigation; HS: investigation and formal analysis; EM: investigation and formal analysis; RS: investigation and formal analysis; PTY: investigation, formal analysis, and writing (review and editing); RW: conceptualization, methodology, formal analysis, writing (original draft preparation, review and editing), funding acquisition, and supervision.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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