**Climate, soil mineralogy, and mycorrhizal fungi influence soil organic matter fractions in eastern US temperate forests**

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**Abstract**

1. Identifying the primary controls of particulate (POM) and mineral-associated organic matter (MAOM) content in soils is critical for determining future stocks of soil carbon (C) and nitrogen (N) across the globe. However, drivers of these soil organic matter fractions are likely to vary among ecosystems in response to climate, soil type, and the composition of local biological communities.
2. We tested how soil factors, climate, and plant-fungal associations influenced the distribution and concentrations of C and N in MAOM and POM in seven temperate forests in the National Ecological Observatory Network (NEON) across the eastern United States. Samples of upper mineral horizon soil within each forest were collected in plots representing a gradient of dominant tree-mycorrhizal association, allowing us to test how plant and microbial communities influenced POM and MAOM across sites differing in climate and soil conditions.
3. We found that concentrations of C and N in soil organic matter were primarily driven by soil mineralogy and climate, but the relative abundance of MAOM *vs.* POM C was strongly linked to plot-level mycorrhizal dominance. Further, the effect of dominant tree mycorrhizal type on the distribution of N among POM and MAOM fractions was sensitive to local climate: in cooler sites, an increasing proportion of ectomycorrhizal-associated trees led to lower proportions of N in MAOM, but in warmer sites, we found the reverse. As an indicator of soil carbon age, we measured radiocarbon in the MAOM fraction, but found that within and across sites, Δ14C was unrelated to mycorrhizal dominance, climate, or soil factors, suggesting that additional site-specific factors may be primary determinants of long-term SOM persistence.
4. **Synthesis**: Our results indicate that while soil mineralogy and climate primarily control SOM C and N concentrations, the distribution of SOM among density fractions depends on the composition of vegetation and microbial communities, with these effects varying across sites with distinct climates. We also suggest that within biomes, the age of mineral-associated soil carbon is not clearly linked to the factors that control concentrations of MAOM C and N.

**Introduction**

Given the vast size of the global carbon (C) pool [(Scharlemann et al., 2014)](https://paperpile.com/c/5pSury/P7oL), and the desire to promote additional soil C storage, the factors that control the persistence of soil C are of critical concern [(Amelung et al., 2020; Bossio et al., 2020; Lal, 2004; Paustian et al., 2019)](https://paperpile.com/c/5pSury/vYYA+YcGf+KJZS+1ZNu). However, not all components of soil organic matter (SOM) have an equal potential to influence carbon-climate feedbacks, as certain soil pools are often more vulnerable to decay and removal than others [(K. Heckman et al., 2022; Torn et al., 1997)](https://paperpile.com/c/5pSury/TrxT+nj7X). Roughly two-thirds of terrestrial organic C is contained in mineral soils as either particulate organic matter (POM) or mineral-associated organic matter (MAOM; [(Sokol et al., 2022)](https://paperpile.com/c/5pSury/RJbS). Formed from partly-decayed plant and faunal biomass, POM is often characterized as vulnerable to continued decomposition, but remains a critical pool of nutrients and carbon in ecosystems where decomposition rates are limited either by climate or the chemical complexity of POM [(Haddix et al., 2020; Lugato et al., 2021)](https://paperpile.com/c/5pSury/k1rx+sg5p). In contrast, MAOM is composed of organic compounds that are associated with minerals through physical and chemical bonds [(Lehmann & Kleber, 2015; Sokol et al., 2018)](https://paperpile.com/c/5pSury/SFRW+nuLe), which may confer some protection against microbial access and further decomposition [(Lugato et al., 2021)](https://paperpile.com/c/5pSury/k1rx). MAOM is therefore a potentially important long-term reservoir of soil C, and there have been many efforts to quantify and characterize MAOM in ecosystems across the globe.

The primary environmental conditions that explain POM and MAOM pool sizes vary widely across study systems [(Craig et al., 2022; Haddix et al., 2020; K. Heckman et al., 2022; Mikutta et al., 2019; Sokol et al., 2022)](https://paperpile.com/c/5pSury/sg5p+EAT6+tlne+RJbS+nj7X), suggesting that there may be context-specific soil conditions and biological communities that influence their formation and persistence. Suggested drivers of POM and MAOM formation in soil include the quantity and chemistry of organic matter inputs from roots and aboveground litter [(Cotrufo et al., 2013; Craig et al., 2022; Crow et al., 2009; Keller et al., 2021)](https://paperpile.com/c/5pSury/vt6U+U3vT+TLSG+tlne), microbial community composition and activity [(Cotrufo et al., 2019; Craig et al., 2018; Frey, 2019; Sokol et al., 2022; Sollins et al., 2009)](https://paperpile.com/c/5pSury/bINK+IiiU+RJbS+YBKx+rJ1j), and soil texture and mineralogy [(K. A. Heckman et al., 2015; Kögel-Knabner et al., 2008; Schöning et al., 2005; Sollins et al., 2009; Swenson et al., 2015; Torn et al., 1997; Weng et al., 2018)](https://paperpile.com/c/5pSury/IiiU+TrxT+j6as+RcCA+M9tZ+XJCD+kXrd). Further, local climate may govern the strength of these drivers across ecosystems [(Kramer & Chadwick, 2018)](https://paperpile.com/c/5pSury/pCzP). Given this lack of consensus and the wide variation in the spatial influence of these drivers, it is likely that cross-scale interactions (*sensu* [Soranno et al., (2014)](https://paperpile.com/c/5pSury/PKPa)) between soil mineralogy, microbial communities, and organic matter quantity and quality collectively determine MAOM pool sizes, wherein the strength of each factor in a particular ecosystem is dependent on the underlying characteristics of the ecosystem. For example, the availability of mineral surfaces for organic matter sorption may serve as a primary determinant of the MAOM formation potential of an ecosystem, and plant or microbial community composition might therefore influence MAOM formation only where the soil mineral composition is suitable for organic matter sorption [(Jones et al., 2015; Kögel-Knabner et al., 2008; Mikutta et al., 2019; Slessarev et al., 2022)](https://paperpile.com/c/5pSury/lg96+EAT6+RuTC+j6as). This context-dependency is common among drivers of large-scale ecosystem processes [(Catford et al., 2022; Hakkenberg et al., 2021; Tedersoo et al., 2016)](https://paperpile.com/c/5pSury/GCmd+Llw0+yniV), including soil organic matter dynamics [(Hoffland et al., 2020; Kramer & Chadwick, 2018; Rasmussen et al., 2018)](https://paperpile.com/c/5pSury/zyCN+bk5v+pCzP), reinforcing the need to test these interactions across scales [(Nave et al., 2021)](https://paperpile.com/c/5pSury/xTBs).

Plant-mycorrhizal associations can also drive within-ecosystem variation in soil C and N cycling and affect the distribution of organic matter as POM and MAOM, particularly in forests. Recent work indicates that in temperate forests where the majority of trees associate with arbuscular mycorrhizal (AM) fungi *vs.* ectomycorrhizal (ECM) fungi, more of the total soil C is stored as MAOM rather than POM [(Cotrufo et al., 2019; Craig et al., 2018)](https://paperpile.com/c/5pSury/bINK+rJ1j). This pattern has been attributed to covarying traits of AM and ECM-associated tree species, such as the typically higher decay rates of AM-associated tree leaf litter [(Keller & Phillips, 2019)](https://paperpile.com/c/5pSury/xUy5), and to the higher rate of root-derived organic matter inputs in AM-dominated forests [(Keller et al., 2021)](https://paperpile.com/c/5pSury/vt6U). Therefore, in AM-dominated stands, fast-decomposing litter and large belowground inputs of labile C may stimulate microbial activity and turnover and lead to faster MAOM production than is observed in ECM-dominated stands [(Craig et al., 2018)](https://paperpile.com/c/5pSury/bINK). Further, differences in the productivity and activity of mycorrhizal fungi themselves, including hyphal production and turnover, and exoenzyme production and organic matter decay, may also influence the formation and stability of POM and MAOM [(Frey, 2019)](https://paperpile.com/c/5pSury/YBKx). Regardless of the mechanism for MAOM formation in ecosystems with distinct mycorrhizal types, it is unclear whether this pattern—larger proportions of C in MAOM under AM *vs* ECM trees—is consistent across forest types or dependent on local climate and soil mineralogy.

Despite the generally longer residence time of MAOM compared with POM [(K. Heckman et al., 2022; Swanston et al., 2005)](https://paperpile.com/c/5pSury/u52e+nj7X), observations of rapid MAOM formation and destabilization under conditions representative of rhizosphere soil suggest that MAOM is composed of some fast-cycling organic matter and some older, more tightly bound organic matter [(Fossum et al., 2022; K. A. Heckman, Swanston, et al., 2021; Jilling et al., 2021; Keiluweit et al., 2015; Neurath et al., 2021; Swanston et al., 2005)](https://paperpile.com/c/5pSury/VAMJ+XBoo+juKu+1qKW+u52e+6Q1E). Because MAOM persistence is dictated both by rates of formation and destabilization, concentrations of MAOM C and N in soil do not necessarily reflect MAOM persistence. The potential for MAOM destabilization may be primarily determined by the same factors that influence MAOM formation, including conditions in the soil matrix [(Kramer & Chadwick, 2018; Waring et al., 2020)](https://paperpile.com/c/5pSury/0uia+pCzP) or the particular features of the organo-mineral associations by which MAOM is formed, including both the type and crystallinity of soil minerals as well as the chemical and physical structure of the organic compounds [(K. Heckman et al., 2018; Kögel-Knabner et al., 2008)](https://paperpile.com/c/5pSury/j6as+ZraT). Alternatively, MAOM persistence may be driven by a separate suite of ecosystem properties than MAOM formation, including the rate of organic acid production and carbon exudation by roots and microbes [(Jilling et al., 2021; Keiluweit et al., 2015)](https://paperpile.com/c/5pSury/XBoo+VAMJ), decoupling the rates of formation and destabilization and leading to patterns in MAOM persistence dictated by local conditions.

To assess the strength of mineralogical, biological, and climatic drivers of POM and MAOM C and N content and MAOM C persistence in soils across the eastern U.S., we analyzed soil collected from plots representing gradients of tree-mycorrhizal associations within seven forested sites in the National Ecological Observatory Network (NEON) that span a range of climate conditions and vary in soil mineralogy. We then compared the roles of climate, soil oxalate-extractable iron content, and mycorrhizal dominance (% basal area of trees associating with either AM or ECM fungi) in driving POM and MAOM C and N concentrations, the distribution of soil C and N among POM and MAOM fractions, and measured δ13C and Δ14C to assess the mechanisms of MAOM C formation and persistence. We hypothesized that tree-mycorrhizal associations would influence the distribution of C and N among the POM and MAOM fractions, but that the effect of dominant tree mycorrhizal type on MAOM and POM would be smaller than the effects of soil mineralogy and climate. Further, we expected that higher concentrations of MAOM C and N would generally correspond with longer MAOM residence times (estimated using MAOM Δ14C) and a higher degree of microbial processing (i.e. higher δ13C) in forest soils, but that this relationship may be dependent on climate.

**Materials and methods**

*Soil sampling*

We obtained mineral horizon soil samples from study plots located within seven forests in the National Ecological Observatory Network (Figure 1) from the NEON Biorepository at Arizona State University (<https://biorepo.neonscience.org/portal/>). Soil samples were collected during the growing season between 2014 and 2020 following standard NEON protocols [(Hinckley et al., 2016)](https://paperpile.com/c/5pSury/E41E). Samples were collected from the upper 30 cm of the soil profile, unless rocks or other site characteristics prevented coring to this depth; in such cases, the maximum accessible depth was used. Within each plot, cores were collected in a number sufficient to provide the necessary soil mass for chemical and physical analyses. Organic and mineral horizon soils were separated, and in cases where multiple cores were collected within a plot, samples were aggregated and homogenized by horizon to represent the soil conditions at the plot scale. Samples were air-dried and sieved to 2mm prior to archiving. Forest sites were chosen for this study based on the availability of physical soil samples suitable for representing within-site variation in tree mycorrhizal type (for site descriptions, see Table 1). Plots within each site (TREE: n=6, all others: n=7) were selected to represent the largest possible range of dominance by AM- vs. ECM-associated tree species at the plot level within each site (Figure 1b).

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*Figure 1. (a) Locations of seven NEON temperate forest sites used in this study (b) Proportion of basal area attributed to arbuscular mycorrhizal (AM)- and ectomycorrhizal (ECM)-associated tree species within the plots where soil was collected in each site.*

*Data selection and processing*

The mycorrhizal dominance (i.e. %ECM and %AM tree basal area) of each study plot was calculated using tree basal area and species identity data from the Vegetation Structure data set on the NEON Data Portal [(NEON, 2022b)](https://paperpile.com/c/5pSury/7TfS). To assess tree species composition, we used woody vegetation data for the most recent year of sampling at the time of data download from each study site, excluding years where sampling efforts were interrupted by external factors or where data collection was otherwise incomplete. We filtered these data to include only live individuals with measurements of stem diameter and with either a species or genus level identification. Woody vines were excluded from the dataset. Mycorrhizal association of each species was determined using the USDA PLANTS database [(USDA, NRCS. National Plant Data Team., 2021)](https://paperpile.com/c/5pSury/99Ka), and plot-level AM and ECM dominance was calculated by dividing the total basal area of all AM-associated trees and all ECM-associated trees from the total basal area in each plot. Where tree species were not known, we used genus to assign dominant mycorrhizal type to individual stems. Trees associated with ericoid mycorrhizal fungi were present in eight of the study plots, and constituted no more than ~0.5% of the basal area in any one plot (Table S1). To assess associations between tree mycorrhizal type and leaf litter production, we used data from the Litterfall and fine woody debris production and chemistry data set from the NEON data portal [(NEON, 2022a)](https://paperpile.com/c/5pSury/OBig). These data were filtered to include only leaf and needle litter, and only data collected in the most recent years when sampling was not disrupted by external factors to maximize the available dataset. These years ranged from 2016 to 2019. Mean annual litterfall mass for each study plot was calculated as the mean value of the annual sum of all leaf and needle litter mass from each study plot. NEON datasets were accessed and merged using the neonUtilities package in R [(Lunch et al., 2021)](https://paperpile.com/c/5pSury/C8La).

*Soil chemical analyses*

We separated air-dried mineral soil samples into particulate and mineral-associated forms of organic matter with density fractionation [(Sollins et al., 2009)](https://paperpile.com/c/5pSury/IiiU) using sodium polytungstate (SPT) adjusted to a density of 1.7 g/mL. Briefly, approximately 5 g of mineral horizon sample was submerged in SPT solution and centrifuged to isolate loose, floating organic material (hereafter the free particulate organic matter [fPOM] fraction). Then, particulate matter contained within soil aggregates was separated from the remaining material by shaking the sample to disrupt aggregates for 18 hours at 200 oscillations per minute on a shaker table and centrifugation to isolate material with a density less than 1.7 g/mL (hereafter the occluded particulate organic matter [oPOM] fraction). The remaining organic matter was considered mineral-associated organic matter (hereafter MAOM). Following separation, all organic matter fractions were rinsed of residual SPT using Nanopure water, oven dried, and analyzed for percent carbon and nitrogen using an elemental combustion system (Costech Analytical Technologies, Valencia, CA, USA). The SPT solution was standardized to a density of 1.7 g/mL for all samples, which was deemed sufficient for clear separation of SOM pools with centrifugation following visual inspection.

Isolated MAOM from mycorrhizal gradient plots in six of the NEON sites was analyzed for δ13C and Δ14C. Samples were analyzed for δ13C at the Stable Isotope Research Facility in the Department of Earth and Atmospheric Sciences at Indiana University. Samples were run on an elemental combustion system (Costech ECS 4010, Costech Analytical Technologies, Valencia, CA, USA) attached to an isotope ratio mass spectrometer (Thermo Finnigan DELTA Plus XP IRMS; San Jose, CA, USA) with a gas bench interface. Δ14C measurements were conducted at the Center for Accelerator Mass Spectrometry at Lawrence Livermore National Laboratory (as per McFarlane, et al. 2013). Samples were not pre-treated to remove carbonates due to the relatively acidic conditions observed in soils from these sites; mean soil pH in each site ranged from 4.33 (BART) to 5.54 (DELA). Samples from ORNL were not included in isotope analyses due to 14C contamination from a nearby hazardous waste incinerator [(Trumbore et al., 2002)](https://paperpile.com/c/5pSury/wVAC). Samples were combusted and converted to graphite (Vogel 1984) and analyzed on the NEC 1.0 MV Tandem AcceleratorMass Spectrometer [(Broek et al., 2021)](https://paperpile.com/c/5pSury/Z61D). The 14C content of each sample was reported in Δ14C notation, corrected for mass-dependent fractionation with measured δ13C values, and then corrected to the year of measurement (2021) for 14C decay since 1950 [(Stuiver & Polach, 1977)](https://paperpile.com/c/5pSury/mmkn).

To determine oxalate-extractable iron content, a 0.2 g subsample of bulk mineral soil from each study plot was submerged in 20 mL of a 0.2 M oxalate extracting solution, shaken at low speed for four hours in the dark, centrifuged for 15 min at 2500 rpm, and filtered through grade 1 qualitative filter paper (Whatman, Buckinghamshire, UK). Extracts were then analyzed for iron content on a mass percent basis with atomic absorption spectrometry (PerkinElmer Instruments, Waltham, MA, USA).

*Climate Decomposition Index*

We extracted monthly temperature and precipitation data for each NEON site for the year 2017 using the Parameter-elevation Regressions on Independent Slopes Model (PRISM) dataset [(PRISM Climate Group, Oregon State University, n.d.)](https://paperpile.com/c/5pSury/vER1). We then calculated the climate decomposition index (CDI) of each site to represent the overall climate using the temperature functions described by Lloyd and Taylor [(1994)](https://paperpile.com/c/5pSury/pfEy) following [Adair et al. (2008)](https://paperpile.com/c/5pSury/Rmsy). The CDI of each site is a function of minimum and maximum monthly temperature and monthly precipitation, and represents an integrated index of conditions important for influencing decomposition. Higher CDI values indicate warmer or wetter sites; for the seven forests studied here, mean annual precipitation was less variable than mean annual temperature, so CDI values primarily reflect differences in site temperature.

*Statistical analysis*

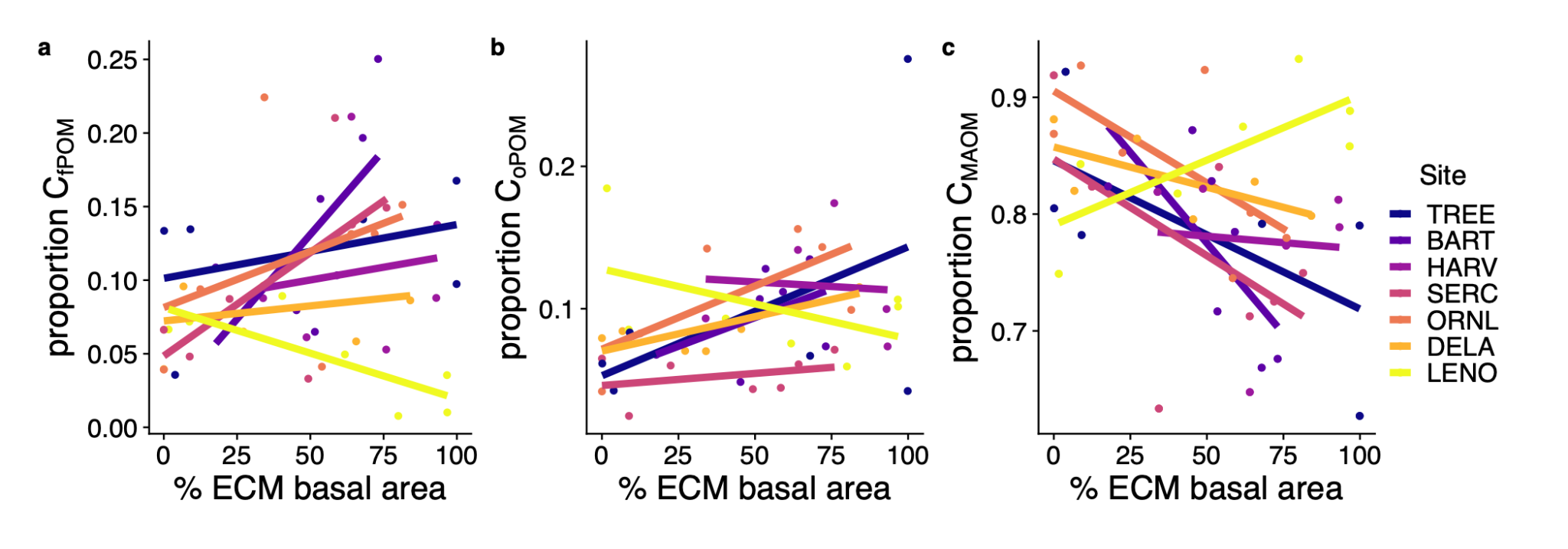
To address our hypothesis that climate, soil mineralogy, and mycorrhizal dominance influence SOM interactively, we constructed a suite of linear models for the proportions of total soil C and N stored in each density fraction as well as the concentrations of MAOM and POM C and N per gram of mineral soil. These models include plot-level mycorrhizal dominance and soil FeOx content and site-level CDI as fixed effects, with study site as a random intercept [(Bates et al., 2015)](https://paperpile.com/c/5pSury/t1pY). Because we expected the effects of these variables to change across climate and soil conditions, we also tested all pairwise interactions between our fixed effects, but included interaction terms in the final models only when they significantly influenced SOM fractions. Finally, we compared models of MAOM C and N content using each site’s mean annual temperature in place of CDI; both methods yielded the same patterns, so we present models using CDI to represent variability in both temperature and precipitation across sites. One sample from Harvard Forest (HARV) with an exceedingly high value of fPOM C and N concentration (> 20 standard deviations above mean fPOM C concentration; > 6 standard deviations above mean fPOM N concentration) was removed from these analyses for a total of 47 observations. Additionally, we constructed linear models with the same fixed and random effects to test how the C:N of each density fraction, as well as the δ13C and Δ14C of the MAOM fraction, responded to mycorrhizal associations, FeOx content, and climate. We also used a series of simple linear models to assess whether the MAOM C concentrations or the proportions of C in the MAOM fraction were associated with MAOM C isotope ratios. Linear mixed effects models were constructed with the lme4 package in R (v. 4.0.2) [(Bates et al., 2015)](https://paperpile.com/c/5pSury/t1pY).

To further investigate how patterns in leaf litter production may have influenced our results, we constructed simple linear models of leaf litter mass across AM-ECM tree gradients within each NEON study site. For this analysis, we gathered data from all plots for which appropriate litter mass data were available from the NEON data portal; therefore, these plots do not necessarily correspond to those from which we were able to request soil samples for density fractionation, instead representing the general patterns in litter production with respect to tree mycorrhizal associations within each site.

**Results**

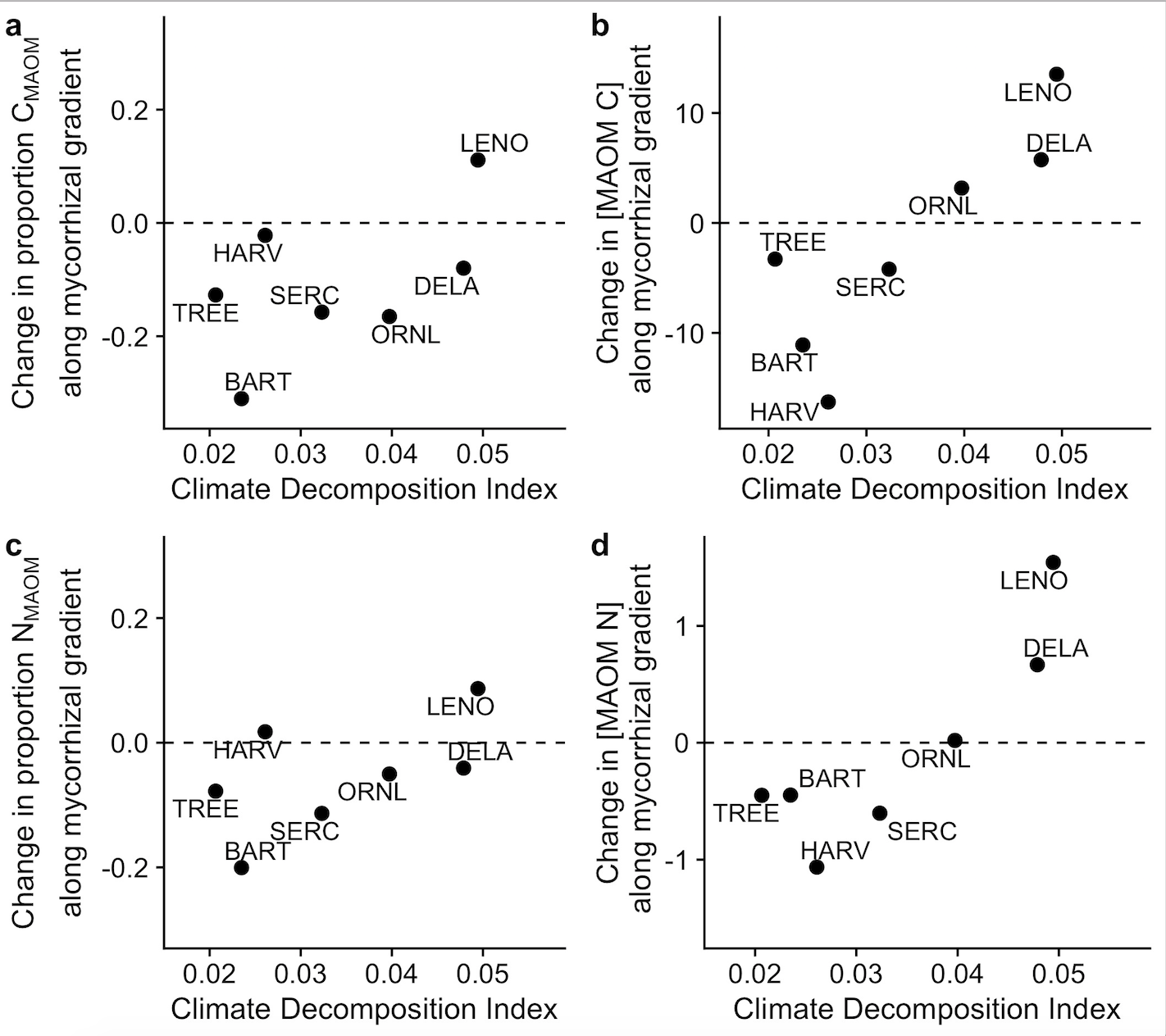
*Distribution of soil C and N among density fractions*

To test our hypothesis that tree mycorrhizal associations influence the formation of POM and MAOM, we assessed how the proportion of C and N in MAOM vs POM varied with the dominant mycorrhizal type of the trees in our study plots. The mean percentage of total soil C [80 ± 1.1 %] and N [89 ± 0.9] contained in the MAOM fraction was generally consistent across study sites, but varied within sites along gradients of AM to ECM tree dominance in the forest plots (Table 2). With increasing ECM-associated tree basal area, we found that the proportion of C in particulate fractions (both fPOM and oPOM) increased (fPOM: t41= 2.21, p= 0.033; oPOM: t41= 2.30, p= 0.027), and in turn, the proportion of C in MAOM decreased (t41= -3.29, p= 0.002) within the majority (6 out of 7) of our study sites (Figure 2). In one site (LENO), we observed the opposite pattern, where a higher proportion of C in the MAOM was associated with increasing ECM tree dominance (F1,5=6.554, p= 0.0506). However, combining all sites, the mean percentage of total soil C in the MAOM fraction decreased by 8.0% across a gradient of 100% AM to 100% ECM tree basal area.



*Figure 2. Proportions of total soil C in the a) free particulate fraction (CfPOM), b) occluded particulate fraction (CoPOM, and c) mineral-associated organic matter (CMAOM) fraction in plots situated along a gradient of tree mycorrhizal dominance in seven forested NEON sites (see Table 1 for site definitions). Points represent individual plots within each study site (n=46). Lines reflect the best fit from linear models specific to each study site.*

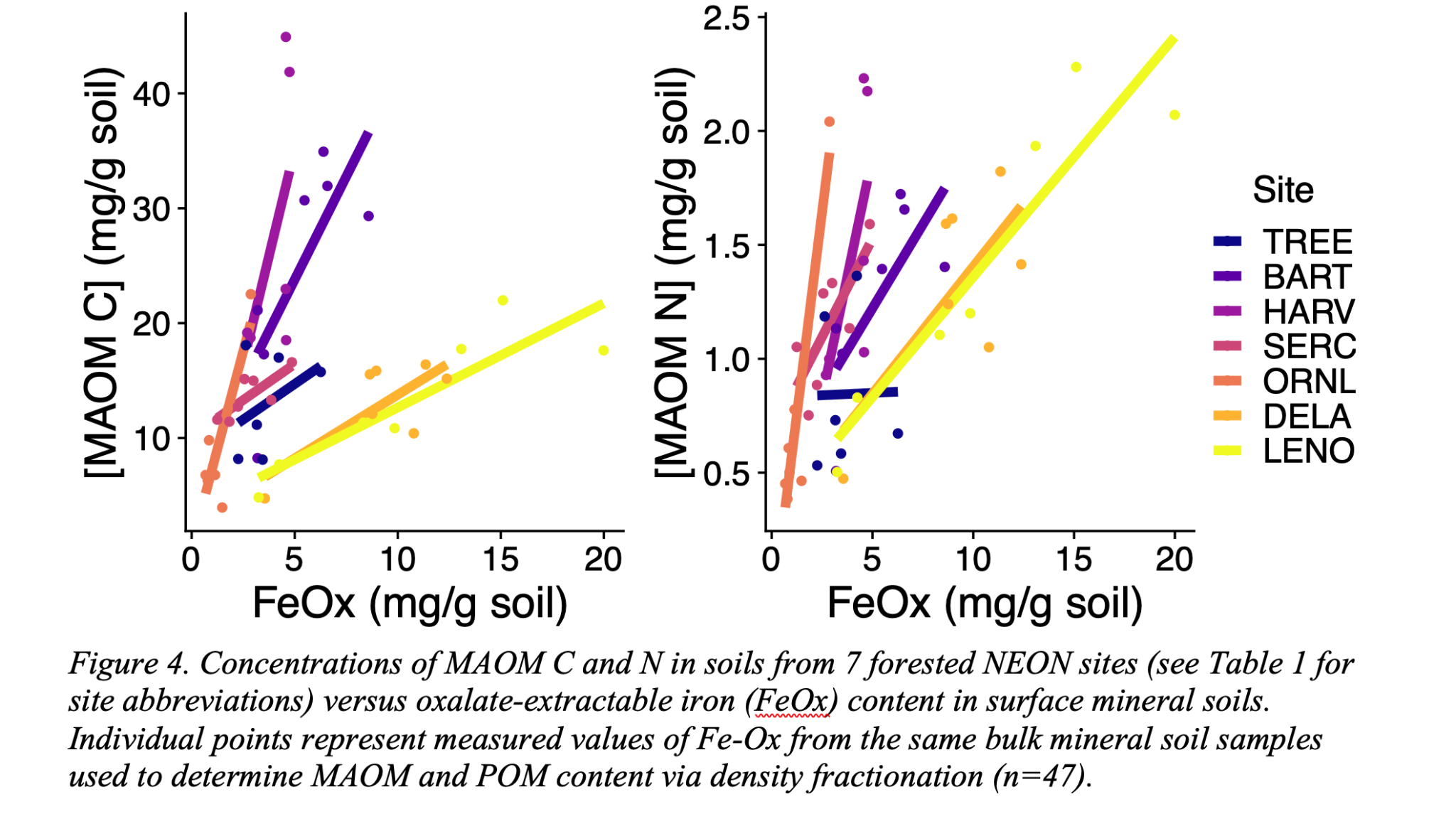
The proportion of soil N in the MAOM fraction responded differently to mycorrhizal dominance under different climate conditions (t40= 2.17, p= 0.036; Table 2); in cooler sites, the proportion of soil N in the MAOM fraction declined with increasing dominance of ECM-associated tree species, but increased with ECM dominance in warmer sites (Figure 3). The proportion of N in the oPOM fraction showed the reverse, with stronger positive relationships with ECM dominance in colder sites (t40= -2.17, p= 0.036; Table 2). To better assess how differences in organic matter inputs may have influenced the patterns we observed along AM to ECM-dominated gradients, we compared leaf and needle litter production with tree mycorrhizal type in all seven forests; the dry mass of leaf and needle litter was negatively associated with plot-level ECM tree dominance in only two sites: TREE (F1,18= 22.65; p < 0.001) and BART (F1,18= 8.95; p = 0.008).



*Figure 3. The influence of tree mycorrhizal dominance (%ECM tree basal area) on MAOM C (a-b) and N (c-d) proportions and concentrations in seven forested NEON sites with respect to climate decomposition index (CDI). CDI represents the mean values and variability of monthly temperature and precipitation in each site (see Methods for details and calculation). Each point represents the slope of the relationship between MAOM C or N and plot-level ECM dominance within a site (as in Figure 2). Panels a and c show the strength and direction of changes in the proportion of C and N in MAOM with increasing ECM tree dominance in each site, while panels b and d show the strength and direction of changes in the concentrations of mineral-bound C and N with increasing ECM dominance. Points above the dotted line represent positive slopes (i.e. soils in these sites show an increase in MAOM C and/or N with increasing ECM dominance; n=7 for all sites except TREE and HARV [n=6]).*

*Mineralogical and climatic controls on SOM C and N concentrations*

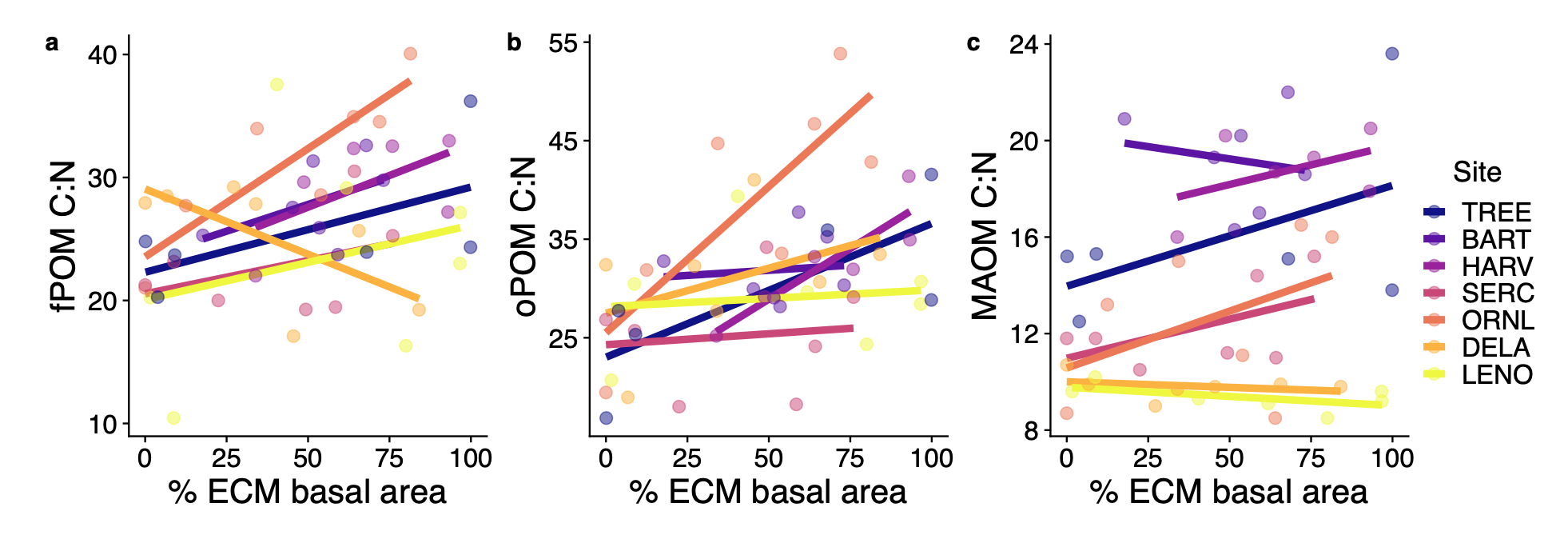
Because soil mineralogy and climate-driven weathering are recognized as key determinants of mineral-organic matter relationships [(Rasmussen et al., 2018; Slessarev et al., 2022)](https://paperpile.com/c/5pSury/zyCN+RuTC), we tested how bulk soil oxalate-extractable iron (FeOx) concentration and site-level climate decomposition index (CDI) influenced the distribution of C and N among soil density fractions. Bulk soil FeOx concentration was strongly associated with higher concentrations of MAOM C (t41= 4.11, p< 0.001) and N (t41= 5.88, p< 0.001 ; Figure 4) but was not an important predictor of oPOM or fPOM concentrations (Table 2). Concentrations of C and N in nearly all soil fractions were negatively associated with CDI; in other words, warmer sites tended to have lower C and N concentrations in both POM fractions and lower MAOM C concentration (Table 2). FeOx concentration was also positively associated with the proportion of C in MAOM (t41= 3.13, p= 0.003), but not with the proportion of N in MAOM (Table 2).

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*Figure 4. Concentrations of MAOM C and N in soils from 7 forested NEON sites (see Table 1 for site definitions) versus concentrations of oxalate-extractable iron (FeOx) in bulk surface mineral soils. Individual points represent measured values of FeOx from the same bulk mineral soil samples used to determine MAOM and POM content via density fractionation (n=46).*

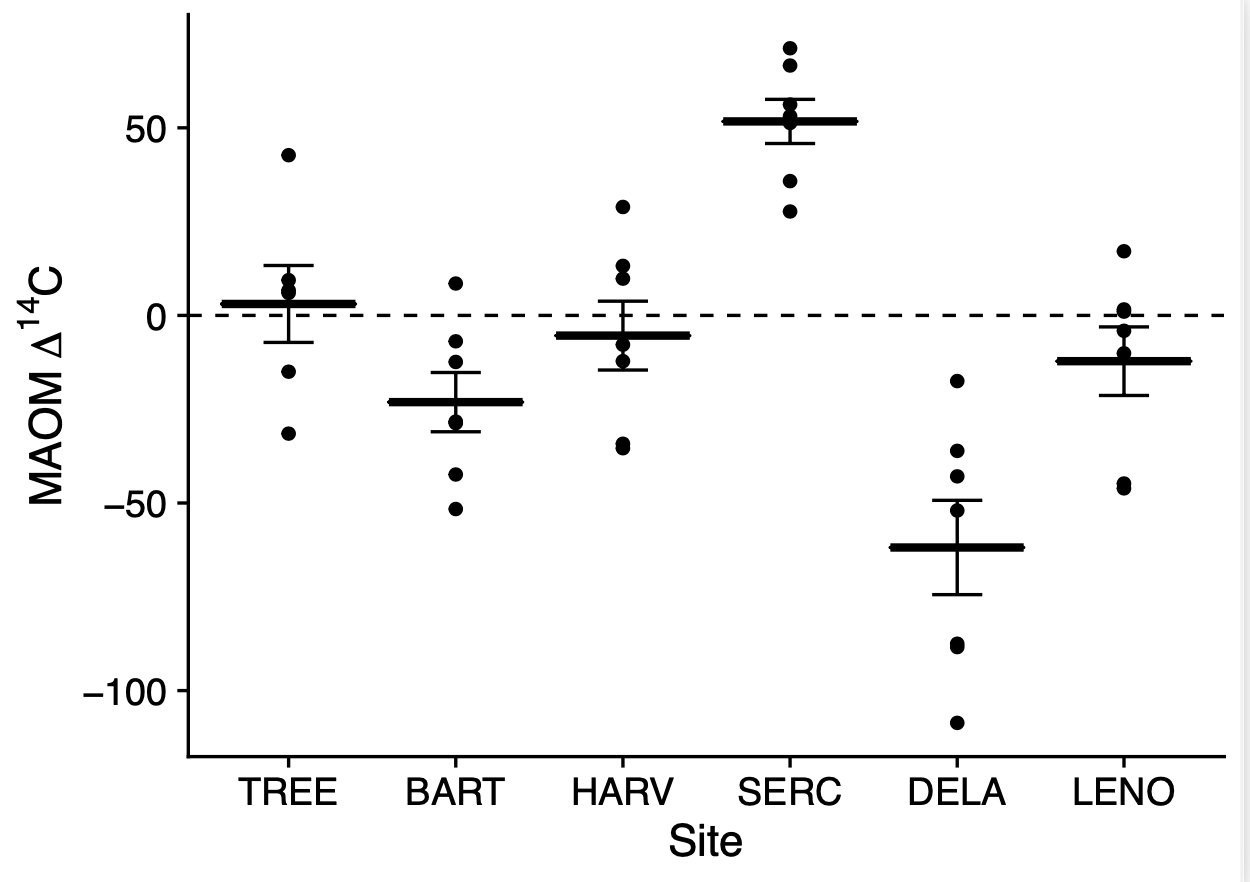
*Patterns in SOM C:N and MAOM carbon isotopes*

To better understand the pathways of MAOM formation, turnover, and SOM origin, we assessed how mineralogical, climatic, and microbial drivers impacted SOM fraction C:N and the isotopic composition of MAOM C. The C:N of all three density fractions generally increased with ECM dominance, but this effect was strongest in the POM fractions (Figure 5; Table 3). In addition, MAOM C:N was significantly lower in warmer sites (t41= -4.42, p< 0.001), while neither the fPOM nor oPOM fraction was associated with site climate (Table 3). Finally, the fPOM C:N was negatively associated with FeOx concentration (t41= -2.62, p= 0.012; Table 3).



*Figure 5. SOM C:N in soils from 7 forested NEON sites (see Table 1 for site abbreviations) along gradients of tree-mycorrhizal associations ranging from AM-dominated to ECM-dominated forest plots within each site. Individual points represent measured ratios of carbon to nitrogen in density fractionated fPOM, oPOM, and MAOM samples (n=47).*

We measured radiocarbon content of the MAOM fraction as an estimate of the carbon age to illuminate possible drivers of MAOM persistence. The Δ14C of the soil MAOM fraction varied substantially across six of the forest sites (ORNL omitted from this analysis due to environmental 14C contamination; Figure 6; Table 4) but was not correlated with any of the soil, climate, or plant-fungal variables that we tested (Figure S1), nor was it related to the MAOM C concentrations (F1,39= 0.215; p = 0.65) or proportions of C in MAOM (F1,39= 0.044; p = 0.83). MAOM in most sites was near modern in origin, with the least modern C in the MAOM fraction at Dead Lake, AL (DELA) and the most modern C incorporated into MAOM at the Smithsonian Environmental Research Center (SERC). MAOM δ13C signatures were positively associated with soil FeOx content, though this relationship was driven primarily by samples from Lenoir Landing, AL (LENO), which had the highest values of oxalate-extractable iron content of any site (Table 4). We also observed a negative association between climate decomposition index and MAOM δ13C, but the size of this effect was minimal (-0.04 ‰ across the range of CDI values represented by our sites; Figure S1). Finally, MAOM δ13C signatures were not associated with either MAOM C concentrations (F1,39= 0.004; p = 0.94) or the proportion of C in the MAOM fraction (F1,39= 0.47; p = 0.50).

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*Figure 6. Mean (± SE) Delta 14C of MAOM from six NEON temperate forests (ORNL was omitted due to site-level 14C contamination). Values greater than zero indicate a modern origin of MAOM C with shorter turnover times; negative values indicate older C (i.e. C fixed pre- 1950). Individual points represent MAOM isolated from mineral horizon soil samples (0-30 cm) collected along tree-mycorrhizal gradients within each site. Sites ordered by increasing climate decomposition index.*

**Discussion**

*Tree mycorrhizal associations influence the relative abundance of MAOM and POM C within sites*

We show that the dominance of AM- *vs* ECM-associated tree species influences the relative abundance of MAOM C and POM C within eastern temperate forests: beneath AM trees, a higher proportion of the soil carbon was found in the MAOM fraction compared to the POM fraction, while the reverse was true beneath ECM trees. Notably, this pattern was consistent in all but one site (Figure 2), and the additional variance in MAOM and POM C proportions explained by other features of each site not measured in this study (i.e. the random effect of site) was small or negligible (Table S2). These results add to the body of evidence that the mycorrhizal communities present in soil may influence the likelihood that incoming organic matter becomes mineral-bound [(Cotrufo et al., 2019; Craig et al., 2018)](https://paperpile.com/c/5pSury/bINK+rJ1j). Differences between forests dominated by tree species with AM *vs.* ECM associations, in particular, may arise from variation in decomposition processes beneath AM- and ECM-associated trees. Previous studies show that both microbial community composition and activity can vary with tree mycorrhizal type [(Carteron et al., 2020; Cheeke et al., 2016; Fitch et al., 2020; Moore et al., 2015; Singavarapu et al., 2022)](https://paperpile.com/c/5pSury/SF4B+4ikG+FhGa+lgzB+0PaR), but the extent to which these changes influence MAOM formation is difficult to test. Mycorrhizal association type is often strongly associated with related factors that are likely to impact SOM decay and sorption, including include the quality of organic matter inputs [(Cornelissen et al., 2001; Huang et al., 2022; Keller & Phillips, 2019)](https://paperpile.com/c/5pSury/xUy5+eJZ7+WtGf), the composition of the free-living bacterial and fungal communities [(Carteron et al., 2020; Fitch et al., 2020; Singavarapu et al., 2022)](https://paperpile.com/c/5pSury/SF4B+4ikG+lgzB), and the production of organic compounds by roots and mycorrhizal fungi themselves [(Keller et al., 2021)](https://paperpile.com/c/5pSury/vt6U).

*Climate mediates the effect of mycorrhizal fungi on SOM nitrogen distribution*

The effect of tree-mycorrhizal dominance on the abundance of MAOM N *vs* POM N varied across sites, implying that the role of mycorrhizal fungi in soil nitrogen dynamics is context-dependent. Specifically, we noted that larger proportions of soil N were found in the MAOM fraction in AM-dominated forest plots in sites with cooler climates, but that in warmer sites, more of the total soil N was found in the MAOM fraction in ECM-dominated plots. One possible explanation for the differences in these patterns across sites may be that the primary origin of the mineral-bound organic matter may vary with site climate [(Angst et al., 2021; Liang et al., 2019)](https://paperpile.com/c/5pSury/0G3O+Q35C). Molecular analyses of MAOM from other NEON sites shows that MAOM in forests receiving >1200 mm precipitation annually with ratios of MAOM C to N greater than 15 may be primarily composed of plant-derived, rather than microbially-derived organic matter [(Yu et al., 2022)](https://paperpile.com/c/5pSury/yGhd). In our study, these conditions are found at BART and HARV (Figure 5; Table 1), where we also noted that leaf litter inputs are lower in ECM-dominated plots *vs.* AM-dominated plots (Figure S1). Therefore, the decline in MAOM N concentration with ECM dominance in these sites may be partly explained by a smaller contribution of plant-derived dissolved organic matter, owing to the generally slower decomposition rate of ECM leaf litter in temperate forests [(Keller & Phillips, 2019)](https://paperpile.com/c/5pSury/xUy5), smaller inputs of aboveground litter in ECM-dominated plots (Figure S1), or lower rates of root-derived organic matter input from ECM *vs*. AM trees [(Keller et al., 2021)](https://paperpile.com/c/5pSury/vt6U). This explanation is broadly supported by other studies showing larger MAOM pools beneath AM versus ECM vegetation in other temperate forests [(Cotrufo et al., 2019; Craig et al., 2018)](https://paperpile.com/c/5pSury/bINK+rJ1j).

In addition to factors that influence the rate of MAOM N accumulation, destabilization and removal of N from soil mineral surfaces may also contribute to patterns in MAOM N concentrations with tree mycorrhizal type. These mechanisms may be particularly important in cooler sites where decomposition rates are lower and N limitation of primary productivity is potentially greater than in warmer sites, leading to N mining by plant roots and mycorrhizae from mineral-bound organic matter [(Jilling et al., 2018; Lambers et al., 2008; Lovett et al., 2018)](https://paperpile.com/c/5pSury/5NdA+ZN2F+eJos). Given that some species of ECM fungi have retained the ability to depolymerize organic matter as a source of N for their plant hosts [(Pellitier & Zak, 2018)](https://paperpile.com/c/5pSury/xWF4), it is possible that N mining by ECM fungi may contribute to lower concentrations of MAOM N beneath ECM trees in cooler sites. Both of these processes—higher MAOM N inputs from AM-associated tree litter and potential removal by ECM fungal mining—may weaken in strength in warmer climates, where differences in leaf litter decay rates between AM and ECM trees are generally smaller [(Keller & Phillips, 2019)](https://paperpile.com/c/5pSury/xUy5), and where higher POM decomposition rates may reduce the need for N mining from mineral soil as a dominant pathway of plant N uptake.

The positive relationship we found between MAOM N proportions and ECM tree dominance in the warmer sites is not as widely supported by existing literature. This relationship is particularly strong at LENO and DELA, both located in central Alabama, USA; however, the potential mechanisms behind the patterns may be distinct at these two sites. At LENO, ECM tree dominance is positively correlated with soil FeOx content, so higher concentrations of MAOM N beneath ECM trees in this site are likely due to corresponding differences in soil mineral suitability [(Keiluweit et al., 2012; Swenson et al., 2015)](https://paperpile.com/c/5pSury/kXrd+0AMg). However, soil FeOx content was not an important driver of the proportion of soil N in the MAOM fraction, and yet the fraction of N in MAOM was positively associated with ECM dominance. At DELA, it is possible that tree species exhibit preferences for other soil conditions that also correspond to tree mycorrhizal type; for example, ECM-associated trees may be abundant in wetter microsites, where decomposition is limited by seasonal inundation and water-filled pore space allows for the rapid transport of low-molecular-weight organic compounds to mineral surfaces [(Waring et al., 2020)](https://paperpile.com/c/5pSury/0uia). However, MAOM in these sites also had the lowest C:N ratios, indicating a greater influence of microbially-derived organic matter in the mineral soil (Figure 5), and it has been suggested that the proportion of microbial vs. plant-derived contributions to MAOM are positively associated with mean annual temperature [(Angst et al., 2021)](https://paperpile.com/c/5pSury/0G3O). Therefore, we suggest that in these warmer sites, fungal tissues may constitute a larger proportion of the organic matter forming MAOM. Given that ECM-associated trees are thought to invest more in mycorrhizal tissue production than in root tissue production compared to AM-associated trees [(Jevon & Lang, 2022; Tedersoo & Bahram, 2019)](https://paperpile.com/c/5pSury/AgjR+gGDc), it is therefore possible that in warmer forests, soil beneath ECM-dominated stands stores more N in the MAOM fraction in part due to the turnover of chitin-rich ECM fungal hyphae [(Fernandez et al., 2016)](https://paperpile.com/c/5pSury/TuXm). If so, the primacy of fungal vs. plant-derived tissues as precursors to MAOM may also account for the fact that changing MAOM N proportions with increasing ECM dominance is not necessarily accompanied by changes in MAOM C proportions (Table 2).

*Effects of soil mineralogy and climate on MAOM C and N*

Across our study sites, we found that the concentrations of MAOM C and N in soil were primarily explained by soil mineralogy and climate. The concentration of oxalate-extractable iron in soil (FeOx), representing the abundance of the poorly crystalline mineral phases best suited for forming organo-mineral associations [(Kleber et al., 2005)](https://paperpile.com/c/5pSury/TAQl), was the strongest driver of MAOM C and N concentrations. On average, a one percent increase in soil FeOx concentration led to an increase of 32 mg of MAOM C and 2.4 mg of MAOM N per gram of soil. This pattern has been observed previously, strengthening the evidence that mineralogy controls MAOM content despite the large variation in soil order, climate and forest species composition among our sites [(K. A. Heckman et al., 2015; Kögel-Knabner et al., 2008; Mikutta et al., 2006; Slessarev et al., 2022; Torn et al., 1997; Weng et al., 2018)](https://paperpile.com/c/5pSury/6kpw+TrxT+j6as+RcCA+M9tZ+RuTC). Further, climate strongly influenced the concentration of MAOM C. We expected that warmer or wetter conditions would accelerate litter leaching and POM decomposition, leading to faster production of the small-molecular weight compounds that form the bulk of MAOM. While this may be the case, we did not find a corresponding pattern with MAOM C concentration: instead, we found that warmer sites generally had lower overall concentrations of MAOM C (Table 2). This pattern may result from a shift in the balance of MAOM C inputs and losses with site climate. In warmer sites, where microbial activity is not as limited by cooler annual temperatures, microbes may be better able to overcome the energetic constraints of accessing MAOM C. If these microbes are indeed using MAOM C as an energy source, MAOM C gained by higher rates of POM decay and greater organic matter export to soil mineral surfaces may be offset by C losses from MAOM decomposition.

Across large spatial scales, the effects of climate on MAOM C may also be modified by concurrent changes in 1) soil weathering rate and, subsequently, the availability of poorly crystalline mineral phases [(Slessarev et al., 2022)](https://paperpile.com/c/5pSury/RuTC), or 2) primary productivity, which controls the rate of organic matter inputs to soil. While we found that plot productivity, represented by tree basal area, was not related to the climate decomposition index of our study sites (F1,46= 0.047; p =0.83), soil in warmer sites did have the largest concentrations of FeOx (F1,46= 13.44; p < 0.001). However, the concentration of MAOM C was negatively associated with climate decomposition index, despite these warmer sites having higher concentrations of FeOx, indicating suitable mineralogy for MAOM formation [(Slessarev et al., 2022)](https://paperpile.com/c/5pSury/RuTC). Therefore, we suggest that climate-driven variation in SOM C concentrations in our study is not likely due to covarying effects of climate on soil mineralogy or organic matter production.

*Carbon isotopes and MAOM decomposition and persistence*

We expected MAOM carbon in warmer sites to have more enriched δ13C and Δ14C signatures due to the generally faster rates of microbial decomposition and C cycling with increasing mean annual temperature [(Shi et al., 2020; Yang et al., 2014)](https://paperpile.com/c/5pSury/Rno9+cUSY). Instead, we found a minor decrease in δ13C with increasing CDI, and no effect of CDI on Δ14C (Table 4; Figure S1). Further, δ13C increased with the percentage of FeOx in our samples, though this trend was most evident at LENO, where the largest variation in FeOx was observed (Figure S1). Given the positive association between concentrations of MAOM C and N and FeOx, this pattern in δ13C may be due to overall faster rates of microbial activity and opportunity for organo-mineral interactions at this site, where both climate and mineralogy are well-suited for MAOM formation. As has been noted in other studies, soil radiocarbon content across our study sites were not necessarily correlated with the factors that controlled the concentrations of C and N in SOM fractions [(K. A. Heckman, Nave, et al., 2021; McFarlane et al., 2013)](https://paperpile.com/c/5pSury/1aG4+0v56). These patterns suggest that local conditions, perhaps land use history, hydrology, or soil disturbance, may be more important than climate, microbial communities or the availability of potential binding sites for explaining variation in MAOM turnover times within biomes. For example, the lowest Δ14C values, indicating older MAOM C, were measured at DELA, a seasonally flooded bottomland forest, where, despite a high mean annual temperature, soil inundation likely limits decomposition and supports the persistence of MAOM C, while inhibiting the formation of new MAOM. Conversely, the high Δ14C values at SERC may be due to regular mixing of new litter material into the mineral soil by the invasive earthworm *Lumbricus rubellus* [(Crow et al., 2009)](https://paperpile.com/c/5pSury/U3vT). The radiocarbon content of the MAOM at other sites in our study was relatively consistent, with most C reflecting a near modern origin, despite variation in climate and soil type. Though others have identified broad-scale patterns in soil Δ14C corresponding with climate and soil mineralogy [(Mathieu et al., 2015; Shi et al., 2020)](https://paperpile.com/c/5pSury/Rno9+rsTB), it is likely that the variation in these factors among the sites used in our study was insufficient to demonstrate these effects.

The site-specific conditions that may have influenced MAOM carbon age in our study may be difficult to identify in broad-scale analyses of soil carbon age. Therefore, to accurately predict fine-scale variation in MAOM persistence, we suggest that individually-derived models of MAOM turnover based on local conditions that influence SOM decomposition rates are needed. Additionally, it is generally accepted that MAOM pools are heterogeneous in nature, with fast-cycling and slow-cycling components that are governed by distinct environmental conditions and inputs and likely vary in their chemical compositions [(Neurath et al., 2021; Sokol et al., 2022)](https://paperpile.com/c/5pSury/RJbS+juKu). Therefore, it is possible that drivers of MAOM C and N concentrations in our sites also exert important controls on MAOM persistence in either the slow-cycling or fast-cycling fraction, but that these patterns were not detectable in our analysis of MAOM radiocarbon content. Further, because radiocarbon-based estimates of MAOM age represent the average persistence of both fast and slow-cycling MAOM fractions, the patterns observed here do not necessarily reflect the stability of the total MAOM pool across our study sites, but rather offer a general comparison of the average MAOM C turnover time.

**Conclusion**

We show that soil reactive iron content and climate were the best predictors of MAOM C and N concentrations in the upper mineral soils in eastern U.S. forests, but that tree mycorrhizal type influences the relative proportion of C and N in MAOM vs. POM. Additionally, climate influenced the effect of tree mycorrhizal type on the relative proportion of N in SOM density fractions, where forests dominated by ECM-associated trees stored relatively more soil N as MAOM in colder climates, but less soil N as MAOM in warmer climates, when compared with stands dominated by AM-associated trees. Radiocarbon content of the mineral-associated soil fraction did not help explain potential mechanisms for the patterns we found in MAOM C and N concentrations: we found that the MAOM fraction radiocarbon age was unrelated to soil mineralogy, tree mycorrhizal type, or climate, suggesting that other factors operating at local scales may be more important for explaining variation in MAOM turnover within biomes. These results support recent findings that mycorrhizal fungi influence the distribution of soil C and N among different SOM fractions, which vary in their vulnerability to decay and are thus critical for predicting soil C and N dynamics in a changing climate. Given the interactive effects of climate and tree mycorrhizal type on the proportion of soil N in MAOM vs. POM, we encourage models of the controls on SOM C and N pools across large spatial scales to more directly account for how the primary drivers of MAOM and POM pools vary across climate gradients.

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**Author contributions**

A.K.L., R.P.P., and J.P.R. conceived of and designed the study, R.P.P. and J.P.R. provided institutional support for analyses, A.K.L. and K.J.M. conducted laboratory analyses, A.K.L. wrote the manuscript, and all authors contributed substantially to edits and improvements.

**Data Availability Statement**

Woody vegetation data (DP1.10098.001) and leaf litter production and chemistry data (DP1.10033.001) for the NEON sites used in this study are available on the NEON Data Portal ( <https://data.neonscience.org/>). Other data and code for the analyses generated by this work are available at <https://github.com/ashleylang/NEON_MAOM>.

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