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# Dynamic interactions at the mineral-organic matter interface

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### **Abstract**

 Minerals are widely assumed to protect organic matter (OM) from degradation in the environment, promoting the persistence of carbon in soil and sediments. In this Review, we describe the mechanisms and processes operating at the mineral-organic interface as they relate to OM transformation dynamics. A broad set of interactions occur, with minerals adsorbing organic compounds to their surfaces and/or acting as catalysts for organic reactions. Minerals can serve as redox partners for OM through direct electron transfer or by generating reactive oxygen species, which then oxidize OM. Finally, the compartmentalization of soil and sediment by minerals creates unique microsites that host diverse microbial communities. Acknowledgement of this multiplicity of interactions suggests the general assumption that the mineral matrix provides a protective function for organic matter is overly simplistic. Future work must recognize adsorption as a condition for further reactions instead of as a final destination for organic adsorbates, and should consider the spatial and functional complexity that is characteristic of the environments where mineral-OM interactions are observed.

## [H1] Introduction

The vast majority of organic carbon in the biosphere (5410 - 6545 PgC; including vegetation, permafrost, soils, coastal systems, and marine sediments<sup>1</sup>) occurs at solid earth interfaces in close spatial proximity to some form of mineral matter. For this reason, carbon flowing through the biosphere will at some point almost inevitably come into contact with mineral surfaces as it undergoes the many transformations inherent in the carbon cycle. Consequently, the interactions between organic and mineral phases have long been the subject of scientific inquiry. For example, most soil organic carbon is associated with colloidal [G] -sized minerals <sup>2,3</sup>, prompting early suggestions that this association is critical for soil fertility and, by extension, is "vital to life" <sup>4</sup>. When toxic organic chemicals were found to be accumulating in the biosphere <sup>5</sup>, mineral phases were investigated for their ability to break down organic contaminants in the environment <sup>6-8</sup>. More recently, interest has turned towards using soils and sediments as repositories for excess atmospheric carbon <sup>9,10</sup>, resulting in intensive research into the mechanisms determining the formation, strength, and durability of mineral-OM associations <sup>10-13</sup>. These latter research efforts have confirmed that minerals can protect OM from degradation. However, observed correlations between individual predictor values such as clay content <sup>14</sup> or abundance of poorly crystalline [G] minerals <sup>15</sup> tend to be specific to certain situations and soil types. To date, no carbon cycle model has succeeded in predicting carbon turnover dynamics based on a generalised, broadly applicable set of mineral phase parameters.

Several principles are critical to understanding mineral-OM interactions. First, soils and sediments originate from a combination of destructive and constructive processes. Weathering processes and organic matter decomposition disassemble rocks and organic debris into basic molecular units, which combine to create micron-sized agglomerations <sup>16</sup>. Biotic and abiotic processes organise the resulting microfabric into a porous, internally-structured medium <sup>17</sup>. Second, minerals generally dominate over OM in soils and sediments in terms of mass proportions, with organic materials (including living organisms) rarely contributing more than single-digit percentages to total mass <sup>18-20</sup> (although exceptions are found in organic soils and organic surface horizons). Owing to the quantitative dominance of the mineral phase, particle size distribution serves as a major constraint on the physical shape of the pore system as it evolves over time <sup>21</sup>. Third, liquid water is the most abundant phase in pore systems<sup>22</sup>, enabling geochemical and biological processes<sup>23,24</sup>. Therefore, it is useful to consider even unsaturated soils [G] and sediments as fundamentally aqueous systems. Finally, the mineral phase, the organic phase, and the pore system are habitats for biota, which constantly modify both physical shape of pores and biogeochemical activities within the system <sup>25</sup>

In this Review, we undertake a broad appraisal of the complexity of the mineral organic interface and derive a perspective for future research efforts. The diverse roles of minerals in the environment are described (**Fig. 1a**), including **OM adsorption** [**G**], catalysis, chemical reactions participation, and reactive oxygen species [**G**] (ROS) generation. The role of the mineral matrix in creating distinct reaction spaces, and its importance of this compartmentalization [**G**] in the

fate of OM, are examined. Finally, we urge the field to move towards considering mineral-OM interactions as multidimensional and multifaceted, and to reconsider the current notion of mineral protection of OM.

## [H1] Fundamental properties

Mineral-organic interactions depend on several characteristic properties of the individual mineral and organic phases, which are briefly described below.

## [H2] Organic phase

Organic molecules can be produced by abiotic processes <sup>26,27</sup> and by chemotrophic microorganisms <sup>28</sup>, but phototrophic [G] algae in the oceans and vascular plants on land are responsible for the bulk of modern biomass production, in roughly equal proportions <sup>29</sup>. Primary production in the oceans creates mostly lipids, proteinaceous materials, and carbohydrates other than cellulose <sup>30</sup>, whereas lignin and cellulose together account for around half of primary production on land <sup>18</sup>.

Heterotrophic [G] organisms use the resulting biomass as a combined source of carbon, energy, and nutrients. Because cell wall passage is restricted to molecules with relatively small molecular masses (< 600-1000 Da) <sup>30</sup>, the size of biomacromolecules must be reduced before assimilation into the cell is possible. Depolymerization [G] and oxidation reactions inherent to extracellular molecular disassembly add ionizable oxygen-containing functional groups to decomposition products <sup>3</sup>. The resulting increasing abundances of polar functional groups (Fig. 2) enhance the aqueous solubility of the products, as well as their chemical reactivity towards metal cations and mineral surfaces, particularly when these functional groups are ionized. Moreover, the smaller molecular size facilitates diffusion <sup>31</sup> of solubilized, reactive organic fragments towards mineral microstructures where they can become entrapped and adsorbed. Once an organic molecule comes close to a mineral surface, the abundance and type of organic functional groups constrain options for further reactions between matrix surfaces and mobile organic adsorbates.

## [H2] Mineral phase

A diverse suite of minerals occur in soils and sediments, including phyllosilicates and aluminosilicates (collectively referred to as silicates); metal oxides, hydroxides, and oxyhydroxides (collectively referred to as oxides); and metal carbonates and sulfides (**Figs. 3 & 4**). In most soils and sediments, phyllosilicates are the predominant component of the finegrained fraction [G] <sup>32,33</sup>. Metal oxides are an order of magnitude less abundant than phyllosilicates except in highly weathered systems and metal-dominated marine benthic habitats (such as hydrothermal vent deposits and ferromanganese crusts), but they contribute distinct reactivity <sup>22,34</sup> (**Fig. 4**).

Minerals possess both permanent (structural) and variable (pH-dependent) surface charge distributions. Subsequent coulombic interactions [G] dictate many physical and chemical properties, such as dispersion [G] behavior, coagulation [G], colloidal stability, solubility, and adsorptive bonding mechanisms <sup>35-37</sup>. Mineral surface reactivity is also driven by surface topography <sup>38</sup>, which results from deviations in atomic crystal structure (such as steps or vacancies), in tandem with particle shape and size. In altering the configuration and number of reactive surface groups per unit mass, topographic surface drivers (steps, edges, and kinks, for example) can thus influence the composition, abundance, and bond strength of compounds at the mineral surface. Surface defects can facilitate the formation of stronger surface complexes or reveal edges that offer unsaturated atoms available for direct complexation <sup>39-42</sup>, and particle size modulates the relative number of reactive surface groups per unit mass of mineral. Finally, the presence and size distribution of pores within mineral grain assemblages also affects the available surface area for organic ligands, as well as exposure kinetics through steric constraints [G] and diffusion limitation <sup>43-46</sup>.

Beyond surface charge and topography, the key properties of fine-grained minerals that determine their interactions with OM (that is, mineral size, shape, charge, and covalent reactivity) are fundamentally derived from their crystal structures (**Fig. 3**). Notably, interfacial energies [**G**] of the different crystal facets [**G**] have a strong influence on particle size and shape <sup>47-49</sup>. For layer-type minerals (phyllosilicates, gibbsite, and layered manganese (Mn) oxides such as birnessite), one facet carries only surface O atoms or OH groups that are fully saturated <sup>50</sup>, <sup>51</sup>. The low interfacial energy of this crystal facet explains the tendency of the associated minerals to adopt a lamellar shape. In turn, this lamellar shape minimizes the tendency towards crystal growth [**G**] and hence favors the persistence of small particles <sup>52</sup>.

The iron (Fe) oxides and framework Mn oxides are exceptions to the condition presented above, in that all their crystal facets carry under- or over-coordinated surface O atoms <sup>53</sup>. One consequence of this imperfect surface O coordination state is that these minerals are primed for covalent reactions with appropriate organic functional groups. Another consequence is that these minerals tend to grow and dissolve readily, and form particles in a variety of shapes depending on the aqueous chemistry <sup>34</sup>. Finally, the relatively high interfacial energy of these minerals enables the coexistence of multiple Fe oxides in most soils, as differences in surface and bulk energies cause reversals in the order of thermodynamic stability as a function of particle size and aqueous chemistry <sup>47</sup>. For example, poorly-crystalline ferrihydrite is often the initial product of Fe<sup>III</sup> crystallization owing to its favorable surface energy. However, at sizes greater than ~5 nm ferrihydrite becomes unstable relative to more crystalline goethite and hematite owing to its less favorable bulk energy <sup>48</sup>

### [H1] Adsorption

Adsorption has long stood as an explanation for the spatiotemporal preservation of organic matter in soils and sediments <sup>54</sup>. However, mounting recognition of the dynamism occurring at the mineral-organic interface, buoyed by diversifying intersectionality in expertise

and analytical techniques<sup>14,55-58</sup>, has emerged in the last decade of biogeochemical research. Rather than a static and irreversible binding mechanism, it is increasingly clear that adsorption itself encapsulates a series of fundamentally kinetic processes: diffusion, competition, and exchange reactions are concurrent and continuous as organic matter accumulates at the mineral surface. The dynamic mineral-OM interface evolves continuously with changes in the chemistry of aqueous phases, OM, and mineralogy, and influences both magnitude and rate of OM adsorption.

Observed complexity in OM adsorptive processes is rooted in the inherent diversity of the reactants. Organic matter encompasses a complex, heterogeneous continuum of structure, processing, and functional diversity <sup>59-61</sup>. Even among small organic compounds, this variety enables adsorption through a combination of multiple anchoring points and mechanisms <sup>62,63</sup> (**Fig. 2**). In tandem, a diverse suite of mineral phases occurs in soils and sediments, varying in surface charge and distribution, topography, and particle size (**Fig. 4**). OM adsorption almost invariably involves a combination of several mechanisms. However, mechanisms of organomineral adsorption can be grouped by the surface chemistry of the mineral (**Fig. 5**), as it affects surface-ligand bond types, coordination state, and spatial organization. These surface chemistries fall into three general groups: variably-charged, non-charged, and negatively charged.

Variably-charged surfaces of metal oxides and phyllosilicate crystallite edges carry hydroxyl groups that are increasingly protonated with decreasing pH, thereby acquiring positive charge. This protonation, in turn, enables the retention of organic ligands through rapid ligand-exchange <sup>64,65</sup>. In many phyllosilicates, substitution of octahedral aluminum (Al) and tetrahedral silicon (Si) with cations of lesser charge will generate permanently charged surfaces. The prevailing negative charge of these surfaces allows for electrostatic attraction of metal cations <sup>66</sup>. When isomorphic substitution is absent, as in some 2:1 phyllosilicates and kaolin-group minerals, the resulting surfaces are electrostatically neutral and thus allow nonpolar organic molecules to accumulate via entropy-driven hydrophobic exclusion phenomena in combination with Lifshitz - van der Waals forces and H-bond formation <sup>3,67</sup> (**Fig. 5**).

As organic ligands partition from solution [G] to mineral surfaces, monolayer coverage of the mineral surface converges into a multilayer molecular architecture <sup>68,69,70</sup> (Fig. 5). Organic ligands tend to have a variety of functional groups (Fig. 2), thus there can be multiple bonding mechanisms between the mineral and OM. This diversity of bonding reactions, paired with the competition of organic compounds for sorptive sites both between themselves and with inorganic ions <sup>71</sup>, results in complex multidimensional structures at the mineral surface (Fig. 5). Ion concentrations are well known to modulate adsorption rates and extents, with variations amongst bonding mechanisms and ion composition <sup>72,73</sup>. However, the importance of competition between organic compounds in structure-selective adsorption (molecular fractionation) is less well-constrained.

Similarly, a plethora of recent work has probed the formation and structure of this multilayer over a range of time and length scales based on nanometer-scale microscopy and

spectroscopy <sup>74</sup>, temporal partitioning of organic ligands from solution onto the mineral surface <sup>75,76</sup>, and detection of uneven structures at the mineral surface <sup>77-79</sup>.

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Adsorption of OM at the mineral-water interface occurs at a rapid rate and reaches equilibrium with aqueous phase in 24 hours for many organic moieties and non-porous minerals <sup>80</sup>. The initial adsorption step is typically fast, and a majority of adsorption is completed within the first few hours after the solid-phase is exposed to aqueous OM. Slow adsorption can follow this first step and is most likely caused by the diffusion of molecules into the intra-particle regions [G] 45,65,81. Most kinetic datasets regarding OM sorption on soils and sediments come from studies conducted on xenobiotic compounds [G] 65. However, studies carried out on organic molecules isolated from soil and aquatic environments suggest that the variables that control the magnitude of sorption, also control its kinetics 82,83. Some of these variables include the size, charge, hydrophobicity, charge density, and concentration of OM, solution composition (pH, ionic strength, ion type), and mineral type (such as metal oxides versus silicates). For example, rapid adsorption is more common at lower pH values, high OM concentration, and in the presence of cations 82-85, where adsorption rates are faster under conditions that promote weaker adsorption affinity, such as in alkaline solutions <sup>83</sup>. Notably, newly adsorbed organics might compete with and actually induce the release of previously adsorbed organic compounds<sup>86</sup>, though this process is not well studied in situ.

Once bound to a mineral surface, an organic ligand [G] only rarely remains undisturbed and instead is vulnerable to desorption and exchange processes, mineralization, and catalytic transformations in situ (Fig. 6). During these processes, organic molecules on mineral surfaces might experience changes to their hydration, structure, isomerization, rotation, and electron delocalization, depending on the type of chemical bonding at mineral-water interface. The overall chemical or electronic state of the adsorbed molecule tends to be similar to the molecule in solution when the adsorbed molecules exhibit H-bonding, Van der Waals interactions, or retain their solvated water (Fig. 6). Conversely, the adsorbed molecule tends to experience substantial changes to its chemical state if it forms covalent or ionic interactions and directly coordinates to the metal atoms at the mineral-water interface, such as during ligand-exchange. These distinct responses are relevant to hydrolysis, electron transfer, and radical mediated reactions, as well as photochemical lability [G] 87,88. Although electron transfer amongst weaklycoordinated species can be limited in some of these complexes, for directly-bound covalent complexes, not only the chemical state of the organic molecule is changed but also the electronic state of the metal atom on the surface (**Fig. 6**). Unlike alkali and alkaline earth elements, transition metal ions in solution or at mineral-water interfaces, such as Mn, Fe, Ni, Cu, and Zn, play an important role in modifying the electronic states of the interacting organic functional groups 89-91.

Adsorption potentially impacts not just the adsorbed OM, but also the minerals: adsorption invariably lowers interfacial energy <sup>92</sup>, so accumulation of OM on mineral surfaces should both facilitate the nucleation [G] of new particles and inhibit the growth of existing particles. This should favor the formation and persistence of smaller particles, a phenomenon observed

experimentally in the apparent stabilization of short-range-ordered **[G]** (SRO) metal oxides (ferrihydrite) and aluminosilicates (proto-imogolite) <sup>93-96</sup>. A logical consequence of this is that correlations between OM and fine-grained minerals, though generally interpreted as reflecting the impacts of minerals on OM, should additionally reflect (to an unknown extent) impacts of OM on mineral nucleation, growth, and transformation.

The transformations summarized here, and the subsequent temporal persistence of adsorbed molecules, are critical to parameterization of organic C cycling in environmental systems. Yet, comparatively few studies have assessed the susceptibility of organic ligands to desorption and exchange despite widespread attribution of adsorptive temporal persistence <sup>12,58,97,98</sup>. Even relatively short desorption experiments using minerals and whole soils reveal partial removal of adsorbed ligands from all surfaces except variable-charge hydrous oxides. However it is likely that there is continuous exchange even with these minerals, masked by near-zero net solubility <sup>58,99</sup>. Mineral transformation and dissolution could also mediate release of adsorbed compounds into the aqueous phase. Anoxic events and biotic reductive dissolution of oxides <sup>100-104</sup>, acidity-driven deprotonation in low-Fe systems <sup>105</sup>, and secretion of root exudates <sup>106,107</sup> could all serve a weathering-like role in releasing adsorbed ligands.

A presumed link between particular bonding mechanisms and biotic mineralization of ligands has not been systematically addressed. Laboratory biodegradation studies suggest OM adsorption, particularly to high-surface area oxides in low-pH systems <sup>108-111</sup>, can hinder decomposition. It is not clear, however, how such findings translate to natural soils and sediments, where dynamic weathering and solution chemistry will alter surface functionality, and where exoenzymes from fungi and bacteria can be abundant in solution. Indeed, whole-soil isotope tracer experiments suggest that metabolism of adsorbed ligands is continuous <sup>112</sup>. Besides microbial degradation and consumption after direct desorption, displacement by biotic exudates can contribute to the decomposition of previously-adsorbed molecules <sup>113</sup>, supported by observations that a substantial proportion of mineral-adsorbed organic matter is microbial in origin <sup>114-116</sup>. Such evidence for the multifaceted role of biotic consumption, exchange, and deposition of organic ligands highlights the uncertainty in underlying controls and their quantitative effect on mineralization rates.

### [H1] Catalysis

The potential of a mineral surface to act as a catalyst [G] and enhance reaction rates results from the combined action of multiple factors. For instance, minerals can concentrate organic compounds to several orders of magnitude higher than the bulk solution <sup>117,118</sup>, thus increasing the pre-exponential frequency factor in the Arrhenius equation. Association with a mineral surface can reduce the degree of orientational freedom [G] of an adsorbate by aligning monomers along two dimensions at planar surfaces <sup>119</sup>, thereby facilitating recurring reaction patterns. Minerals also create centers of reactivity by providing coordinatively unsaturated sites and steric enhancement [G] of reactions at surfaces with substantial topography <sup>38</sup>. Furthermore,

mineral surfaces can transfer protons to a sorbate as well as offer empty electron orbitals for the reception of bonding electron pairs (Broensted- and Lewis-acid/base functionalities<sup>120</sup>). Lastly, minerals entirely made of transition metals (such as Fe and Mn oxides) or acting as solid supports for adsorbed transition metals at their surfaces and in interlayers (phyllosilicates) can contribute transition metal specific catalytic functionality <sup>121,122</sup>.

Laboratory observations made on the catalytic reactivity of different minerals can be used to recognize three mineral types of chemical reactivity (**Fig. 6**), although the intensity of the organomineral interactions within each class varies with changes in mineralogy, OM concentration, and geochemistry of the environment.

Class I minerals are dioctahedral 2:1 clays that adsorb large organic molecules through cation-bridging and through hydrophobic expulsion and aromatic ring  $\pi$ -interactions with surfaces or Lifshitz - van der Waals forces. Direct complexation of OM can occur at edge sites. Because hydrophobic expulsion is the primary driver for adsorption, kinetics of adsorption are fast, macromolecular structures are rarely disturbed, and limited electron transfer is expected. However, heterogeneous oxidation of OM can occur on clay surfaces in the presence of adsorbed  $O_2$  and transition metals  $O_2$  and transition metals  $O_2$  are catalytic activity of Class I minerals is expected to be low with slow kinetics  $O_2$  and  $O_3$  are catalytic activity of Class I minerals is expected to be

Class II minerals are metal oxides, silicates, and 1:1 clays, and exhibit high adsorption capacity (**Fig. 6**). These minerals tend to catalyze moderate changes to the structure and electronic state of adsorbed molecules, primarily through direct bonding, hydrolytic breakdown from reactions with surface OH, and heterogeneous oxidation with adsorbed  $O_2$  <sup>123,126,127</sup>.

Finally, Class III minerals are highly reactive in terms of magnitude of sorption, and in terms of hydrolytic and electrolytic breakdown and heterogeneous oxidation (**Fig. 6**). Reduced clays (those with substituted transition metals), Mn oxides, sulfides and magnetite belong to this category <sup>123,127-129</sup>. Mineral carbonates and sulfides, through major changes of surface acidity at the interface, play an important role in the hydrolytic breakdown of macromolecular organic molecules. Breakdown of large molecules into small molecules is possible, as is the subsequent transformation into larger molecules through nucleophilic addition, radical mediated polymerization and large molecule formation reactions (**Fig. 6** <sup>129-131</sup>). Because of these changes, substantial alterations to mineral surface structure, including even

eventual dissolution, are anticipated in the case of Class III minerals (**Fig. 6**). Strongest proof for all of these observations comes from solution studies, rather than from spectrometric work directed at the interface <sup>123,132,133</sup>. The solution studies reveal the progress of catalytic reactions and their rates; however, not the reaction mechanisms at the interface.

The impact of mineral catalytic behavior on OM stability is expected to be greater where there is direct organomineral interaction, or where there are monolayer coverages of OM (**Fig.** 6). As a consequence, the intensity of organomineral interactions should be strongest during monolayer OM adsorption. In multilayer adsorption, conversely, the overall role of minerals as catalysts should be comparatively diminished as OM-OM interactions increasingly predominate over mineral-OM interactions (**Fig.** 6). Although a majority of molecular studies conducted on

soils or sediments belong to the latter category <sup>74,134,135</sup>, the impact of layer thickness on OM behavior remains relatively poorly understood because of the analytical difficulties in exploring these systems.

Metal atoms of mineral surfaces in the proximity of adsorbed molecules also tend to experience changes to the adsorbed water and surface hydroxyls, and to their electronic state. These can be three-fold: changes in the metal-ligand coordination (and hydration and hydrolysis) environment; complete e-transfer and oxidation or reduction of surface metal atoms; and perturbations to the energy levels of valence electrons <sup>133,136,137</sup>. Although changes are less notable during the formation of weak H-bonded complexes, the formation of direct ionic and covalent interactions impact the binding of metal atoms of mineral surface strongly, with some additional impact to the buried atom layers next to the binding site <sup>38</sup> (**Fig. 6**). As a result, adsorption of OM to mineral surfaces often modifies the characteristics of both OM and mineral surfaces, and only in rare occasions are the chemical characteristics of each preserved at the interface during bonding <sup>38,85,130</sup>. These considerations highlight the difficulty to distinguish between two fundamental functions (catalysis and redox reactions) of mineral surfaces when considering OM transformations at solid earth interfaces. At this time, it seems there is considerable conceptual uncertainty regarding these functions, with the available evidence favoring a role as reactant.

#### [H1] Redox reactions

Electron transfer reactions between OM and minerals, directly or mediated by microbes, are a primary pathway contributing to the oxidation, transformation, and mineralization of carbon within soils and sediments (**Fig. 7a**). Minerals and OM can serve as either electron donors or acceptors for a vast array of heterotrophic or mixotrophic [G] microbial metabolisms spanning a broad ecologic and taxonomic diversity <sup>138</sup>. Various mechanisms are employed in the microbial coupling of organic carbon oxidation to the reduction of Fe and Mn oxides, including direct electron transfer via outer membrane enzymes <sup>139</sup> or conductive nanowires [G] <sup>140,141</sup> and indirect transfer via endogenously produced electron-shuttling molecules that are secreted by an organism after which they adsorb to an oxide surface and transfer the bacterial-derived electron to the mineral <sup>142</sup>. The latter reaction pathway is a coupled biotic-abiotic pathway, whereby the second step is a direct electron transfer reaction between the organic molecule and mineral surface.

A variety of organic molecules can be transformed via direct electron transfer at the surface of minerals, including Fe and Mn oxides, sulfides, and phyllosilicates <sup>123</sup>. Within natural systems, metal oxides have garnered particular attention for their ability to serve as effective oxidants of a wide range of organic compounds <sup>143</sup>, including humic acids, phenols, anilines, low molecular weight organic acids (such as oxalate, pyruvate, citrate) <sup>7,144,145</sup>, and more recently proteins <sup>128,146</sup>. The overall reaction sequence involves adsorption of the organic reactant to the

(hydr)oxide surface, transfer of electron(s) to the mineral-hosted metal, and subsequent detachment of the reduced metal and oxidized organic molecule <sup>123</sup>.

The adsorption, fractionation, and oxidative transformation upon reaction of dissolved organic matter with minerals are dependent on mineral surface chemistry, including the composition and abundance of terminal bonds <sup>123,147</sup>. Despite typically lower adsorptive capacities, Mn oxides are stronger oxidants than Fe oxides, in large part owing to differences in redox potential <sup>123,148,149</sup>. Indeed, Mn oxides are considered the strongest naturally occurring oxidants, and thus considerable attention has been placed in investigating the role of Mn oxides in organic carbon transformations <sup>143</sup>. In general, organic compound oxidation rates increase with average oxidation state, redox potential, and specific surface area of the Mn oxide and decrease with apparent activation energy of the overall reaction and the pH<sub>pzc</sub> of the oxide <sup>143</sup>. Furthermore, OM oxidation rates increase with decreasing pH, which has been primarily attributed to the higher redox potential at lower pH.

As mineral-OM redox reactions rely on surface complex formation <sup>150</sup>, the efficacy of mineral-induced organic matter oxidation and mineralization is ultimately determined by factors controlling initial adsorption. Thus, the extent and rates of organic carbon transformation and mineralization at mineral surfaces decline if surfaces become passivated [G], for instance by high organic matter loadings <sup>123</sup> and/or reaction products blocking reactive sites on the surface <sup>143</sup>. Further, mineral ripening [G] and recrystallization to less reactive phases during reaction can constrain mineral surface reactivity over time. Extrapolation of the rates and products of mineral-mediated OM transformations within natural systems from these reactions involving pure phases is therefore limited and convoluted.

Nevertheless, correlations between oxidized Mn and oxidized OM point to a causal link between Mn oxidation and carbon transformations <sup>113,149,151</sup>. In fact, the intimate and diverse association of OM with Mn oxides <sup>149,152</sup> leads to a continuum of reactions, including mineralization and polymerization within layers coating the oxide surface <sup>129</sup>. Yet beyond correlations, there is limited data directly linking natural Mn oxides (or any mineral) to OM oxidation and transformation, pointing to a clear need for future investigations targeting mineral-mediated OM redox mechanisms.

## [H1] Reactive Oxygen Species

Mineral-derived ROS<sup>153</sup>, particularly hydrogen peroxide ( $H_2O_2$ ), superoxide ( $O_2^{\bullet,-}/HO_2$ ), and hydroxyl radical ( $HO^{\bullet}$ ), represent likely important but currently underappreciated players in the transformation of soil and sediment organic matter <sup>154</sup>. Among various biological and (photo)chemical processes <sup>155,156</sup>, common soil and sediment minerals have also demonstrated the ability to produce ROS, including  $H_2O_2$  and  $HO^{\bullet,157-161}$ . These ROS-producing minerals include primary silicates (such as plagioclase), oxides, sulfides, and clays, with the titanium dioxide phases rutile and anatase ( $\alpha$ - and  $\beta$ -TiO<sub>2</sub>), iron oxide hematite ( $\alpha$ -Fe<sub>2</sub>O<sub>3</sub>), and Fe(II)-bearing clays such as biotite and chlorite having the highest ROS yields observed to date <sup>157,161</sup>.

Minerals contribute to ROS production via water and O<sub>2</sub> reaction at defect sites or with mineral-hosted or adsorbed Fe(II), and via photochemical electron transfer <sup>158,161,162</sup> (**Fig. 7b-c**). Alternatively, the reductive or oxidative dissolution of minerals leads to the release of metals (particularly iron) or ligands (particularly sulfide) that can create ROS upon reaction with oxygen <sup>158</sup> (**Fig. 7b**). Thus, albeit limited in number, these investigations of mineral-derived ROS indicate that the mechanisms and pathways of ROS formation vary as a function of the mineral, light, and aqueous environment.

Mineral-derived ROS within natural soils and aquifers have been measured in a limited number of investigations. In subsurface soils and groundwaters, fluctuating redox conditions and oxygenation of Fe(II)-bearing minerals are the primary contributors to ROS production <sup>161,163</sup>. Formation of ROS including O<sub>2</sub>\* and HO\* has been linked to the oxidation of aqueous Fe(II) emanating from subsurface marine 164 and lake sediments 165. Indeed, the oxygenation of reduced soils, sediments, and waters results in a cascade of ROS pathways, including initial formation of both  $O_2^{\bullet -}$  and  $H_2O_2$ , ensuing dismutation of  $O_2^{\bullet -}$  to  $H_2O_2$ , and ultimately formation of  $HO^{\bullet}$  via reaction of aqueous Fe(II) and H<sub>2</sub>O<sub>2</sub> (Fenton reaction)<sup>166</sup>. In sunlit surface environments, photochemical reactions at oxide surfaces have also been found to play an important role in ROS (particularly  $O_2^{\bullet}$  and  $H_2O_2$ ) formation, including in both desiccated and wet desert soils <sup>159</sup>. Production of ROS in soils pre-washed to remove photosensitizers such as nitrate point to direct mineral-induced ROS generation upon irradiation <sup>159</sup>. Superoxide and peroxides can be stabilized within desiccated soils via complexation to metal oxide surfaces <sup>159</sup>. These ROS are rapidly released when soils are (re)saturated, leading to pulses of oxidants within pore waters. Thus, several oxidants are formed upon oxygenation of soils and sediments, the species and flux controlled by mineral composition and mechanism of ROS generation.

ROS vary in their selectivity and reactivity toward carbon functional groups, and react with a wide range of organic compounds, including carbohydrates, fatty acids, and biomolecules (such as DNA and proteins  $^{167}$ ). The reaction progression upon interaction of DOM and ROS ranges from partial oxidation of organic carbon compounds and formation of low molecular weight organic acids to complete oxidation to  $CO_2$ , depending on the reactants and conditions (such as pH). Aromatic moieties, such as quinones or humics, have been identified as primary sinks for  $O_2^{\bullet -}$  in marine DOM, the reaction of which could lead to a catalytic cycle regenerating the original reactant and forming  $H_2O_2^{-168}$ . Hydroxyl radical is a particularly unselective and strong oxidant  $^{169}$ , having high reaction rates with DOM ( $10^8$  M  $C^{-1}$  s<sup>-1</sup>) $^{170}$  and the ability to oxidize carbon compounds within the DOM pool that are otherwise difficult to photo- or biodegrade  $^{171,172}$ .

Although indirect evidence points to probable contributions of mineral-derived ROS in carbon processing, few studies to date have directly interrogated the role of ROS in carbon transformation and degradation. Still, the potential for ROS to be quantitatively relevant in carbon cycling was exemplified by a previous investigation where rates of  $CO_2$  produced from  $^{\circ}OH$  (109 µmol  $CO_2$  m $^{-2}$  d $^{-1}$ ) in Arctic soils were found to be on the same order of magnitude as bacterial mineralization of DOM in surface waters  $^{165}$ . Similarly, DOM (including coloured

DOM) is a predominant sink of  $O_2^{\bullet}$  within several marine systems  $^{173,174}$ , and photodecomposition of DOM in lake waters has been linked to ROS formation <sup>175</sup>. One of the first studies to directly link ROS to remineralization of carbon within soils showed that 'OH derived from Fe(II) oxidation was responsible for DOM oxidation to CO<sub>2</sub> in Arctic soils <sup>154</sup>. Specifically, production of CO<sub>2</sub> in Arctic soil incubations scaled with 'OH production stimulated via H<sub>2</sub>O<sub>2</sub> addition. Similarly, stimulation of carbon mineralization within aerobic incubations of humid and tropical soils amended with Fe(II) pointed to contributions from Fenton based reactions and ROS <sup>57,176</sup>. In support of this, carbon mineralization decreased (8% less CO<sub>2</sub>) compared to controls) within these incubations upon the addition of 'OH scavengers, further implicating 'OH as a terminal oxidant of OM <sup>57</sup>. Similarly, addition of superoxide alone or in combination with Fe to soil incubations triggered 'OH formation and concomitant degradation of soil organic matter <sup>177</sup>. Oxidation of tetracycline to CO<sub>2</sub> was also recently linked to 'OH formation upon oxidation of sediments in lab incubations <sup>161</sup>. Lastly, soil organic matter oxidation and decomposition in the absence of enzymatic activity (inactivation via autoclaving) within soils implicated mineral-derived ROS <sup>56,178</sup>; yet, the mechanism at play was not elucidated.

### [H1] Definition of microbial habitats

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Mineral-organic matter associations play a critical role in structuring and compartmentalizing the biological reaction space of soils and sediments into microsites [G], and, in combination with the water potential, determine the extent to which the individual microsites are connected <sup>179</sup> (Fig. 8). Pore connectivity, morphology, and size distribution at the microsite scale either enhance or restrict microbial access to OM <sup>180</sup>. They also determine the conditions that microbial decomposers are exposed to, impacting the range and magnitude of the microbial activities that occur. For instance, the compartmentalisation of space results in the juxtaposition of microsites with different environmental conditions that can be mutually exclusive (for example, oxic and anoxic) at very fine scales such as within a single aggregate <sup>171,172</sup>. This juxtaposition leads to the emergence of gradients, allowing a range of processes to proceed. It also facilitates the development and persistence of extensive microbial diversity, because interactions that tend to reduce diversity, such as competition for resources or antagonism, are attenuated in compartmentalised space <sup>173,174</sup> (**Fig 8**). Understanding the link between microbes and their environment is fundamental to understanding soil and sediment microbial ecology, microbial activity and the involvement of microbial decomposers in OM dynamics <sup>16,199,200</sup>. As many of the functional properties of soils and sediments (nutrient cycling and pollution degradation, for example) emerge from the diversity of their microbial inhabitants 175, it is not unreasonable to suggest that this functioning is ultimately dependent on the minerals' role in the structuring of space.

Indeed, we posit that the overriding function of mineral matrices (Tier 1 in **Fig. 1b**) should be seen in their role in compartmentalizing the system. Microbially driven decomposition

requires decomposers and organic substrates to encounter one another <sup>201</sup>. However, OM is heterogeneously distributed at scales that are relevant for microbial decomposition <sup>202</sup>, as are microbial communities themselves <sup>203</sup>. This distribution results in a patchwork of cold and hotspots of microbial activity <sup>204</sup>, which is a fundamental characteristic of soils. Furthermore, there can be non-linear, concave relationships between the concentration of available organic substrate and microbial activities. These can be particularly apparent in activity hotspots and are possibly related to the saturation of cellular transport systems or local O<sub>2</sub> limitations <sup>132</sup>. A major consequence of non-linear, concave relationships between the concentration of available substrate and decomposition result is the lowering of overall organic matter decomposition relative to when substrate and decomposers are homogeneously distributed<sup>132</sup>.

The variable connectivity of the microsites (**Fig. 8**) means that microbial communities function and evolve more or less independently of communities in other microsites and under different environmental conditions <sup>176,177</sup>. Taxa that are better adapted to the prevalent microsite conditions, such as through the capacity to use electron acceptors other than O<sub>2</sub> in anoxic conditions <sup>178</sup>, can competitively exclude other, less adapted taxa, with the potential loss of metabolic capacity from the community. Metabolic capacity can also be lost when mutations cause the inactivation of genes that are not maintained by selective pressure (in essence, the gene does not provide any benefit to the microbe in the microsite) or genetic drift <sup>179</sup>. Such losses of capacity from microsites could have consequences for subsequent organic matter dynamics, as they would allow organic molecules to persist over time, regardless of the intrinsic properties of the molecules or the microsite conditions.

Just as plant roots affect and select microbial communities within their sphere of influence, the rhizosphere, minerals also exert influence over the microbial communities within their vicinity, and therefore the metabolic pathways and organic matter dynamics that are present in microsites <sup>185</sup>. The sphere of influence of minerals has been termed the "mineralosphere" <sup>186</sup>. For example, different types of minerals select different microbial communities <sup>187-189</sup>, and the selective pressures exerted by minerals can even be greater than those associated with different environmental conditions or in the presence of different organic matter sources <sup>188,190</sup>. Minerals, therefore, should not be viewed simply as supporting surfaces onto which microbial communities arrive from the surrounding environment in a random and passive fashion. The selective pressure exerted by minerals is likely derived from differences in resource availability at the surface or in the vicinity of the minerals induced by their function as a mobilisation filter, from the mineral acting as an electron acceptor or donor <sup>191</sup>, from differences in surface charge, area, or topography<sup>192</sup>. Microbial attachment to mineral surfaces can also trigger wholesale changes in microbial functioning, with metabolic activity being either stimulated or inhibited, or growth and biofilm formation being promoted or suppressed, depending on the mineral type <sup>193,194</sup> and depending on the microorganism <sup>195,196</sup>. The types of microbial metabolic pathways present at mineral surfaces is likely to have a major impact on mineral-organic matter associations and, indeed, a number of studies have suggested that a large part of mineral associated organic matter has been processed by microbial communities <sup>109,197</sup>, <sup>181</sup> particularly in agroecosystems or in

contexts where microbial communities flourish <sup>182,183</sup>. Furthermore, the extent to which microbially processed organic matter associates with minerals could even depend upon the composition of the microbial communities <sup>198</sup>. The mechanisms underlying the production of mineral associated organic matter via microbial processing remain as yet un-elucidated however. Nevertheless, the overriding message that emerges from these studies is that microbial communities not only mineralise organic matter but are also agents in the generation of mineral-organic matter associations.

Ultimately therefore, the properties, including the biotic composition, of the compartmentalized space are intimately linked to microbial-mineral associations. These properties affect the local environmental conditions, which, in turn, modulate subsequent biogeochemical processes <sup>184</sup>. This kind of functioning, in which feedback controls are prominent and in which the overall performance of the system is not reflected in the properties of the individual components, but emerges from the component interdependencies, is typical of complex systems <sup>184,185</sup>. The feedback controls impose a number of static and dynamic constraints on the system (Tier 2 in **Fig. 1b**). These constraints then determine the intensities and rates at which molecular scale biogeochemistry might be able to proceed (Tier 3 in **Fig. 1b**). Depending on the extent to which higher level (Tier 1 and 2) constraints operate in different parts of the system, different lower tier processes could occur simultaneously within the same system. How the structuring of space into a network of more or less connected microsites affects the assembly of microbial communities and determines local environmental conditions that modulate the expression of microbial functions and, ultimately, ecosystem performance has not yet been fully explored.

## [H1] Summary and future perspectives

The mechanisms underlying mineral-organic interactions have a long tradition of being explored through adsorption experiments conducted with defined phases in tightly controlled laboratory environments. However, we strongly urge that this research take a more holistic, multidimensional view (**Fig. 1b**). We advise the scientific community to resist the temptation to attribute given manifestations of carbon dynamics to a single predictor value, such as the abundance of a certain mineral species, phyllosilicate clay content, or operationally determined specific surface area values, as has been frequent practice in the past. Rather, soils and sediments should be investigated as multidimensional entities, whose overall functional performance depends on the extent to which the structure of the matrix supports molecular scale reactions. Translating this insight into novel modeling approaches will be key to improved predictions of global biogeochemical cycles. Quantitative information about pore size, pore morphology and pore connectivity can be obtained from modern multidimensional imaging techniques such as computed tomography, rendering an explicit consideration of architectural features fundamentally possible. In this final section, we examine some of the outstanding questions and pressing research needs in organomineral interaction research.

## [H2] Structure and prokaryotic performance

Interactions within the mineral-organic matter complex not only have immediate and direct effects on organic matter dynamics as described above but might also have indirect and longer term consequences. Metabolic dependency [G], when microbial cells lose the ability to produce essential metabolites and instead use those released into the environment by other cells, is common in natural microbial communities <sup>179,180</sup>. Such metabolic dependency develops only if communities remain spatially co-located long enough for coevolution to occur <sup>181,182</sup>. Metabolic dependency has been shown to evolve in experimental populations in 1000 to 2000 generations <sup>186</sup>. In soils and sediments, the probability of such co-evolutionary interactions occurring is greater if the physical structure of the microsite persists over time (Fig. 8). Microsite persistence is related to the organic matter content <sup>183</sup>, but could also be related to the mineralogy. For example, there is evidence to suggest that 1:1 clays and oxides produce more persistent microsite structures <sup>183</sup>, probably owing to electrostatic interactions between the variable charged minerals <sup>184</sup>. In view of the suggestion that the persistence of organic matter in soil is related to the heterogeneous distributions of organic substrate and metabolic capacities <sup>187,188</sup>, it would be interesting to test the hypothesis that the mineral compartmentalizing the environment affects the distribution of metabolic capacity at the microsite scale. This could be achieved using shotgun metagenomics <sup>189</sup>, at appropriate scales, in soils with different mineralogies.

## [H2] Adsorption, desorption and protection

Decomposition of adsorbed OM is typically substantially slower than decomposition of the same type of OM in a freely suspended or dissolved state <sup>190</sup>, leading to the notion of 'sorptive protection' of OM. However, desorption can be facilitated by changes in pH, electron availability, or by modifications to sorbent surfaces <sup>105</sup>. Indeed, evidence is mounting that plants are equipped with tools to achieve this exact purpose, such as the exudation of organic compounds designed to release bound organic matter from mineral surfaces <sup>191</sup> and to adjust patterns of microbial community composition for optimal functionality <sup>192</sup>. With these developments in mind, we suggest that sorptive protection should be re-evaluated as an explanation for organic matter persistence, especially in systems inhabited by plant roots.

More broadly, the idea that there is reduced bioavailability of adsorbed substrates must be further scrutinized. Potentially, it is not that the adsorption process exerts overwhelming bonding forces, preventing breakdown of sorbates. Instead, adsorption could temporarily fix the location of an adsorbed substrate within a given pore network. Such localization would force the decomposer community to overcome the resulting spatial complexity of substrate locations, slowing the rate of decomposition <sup>187,193</sup>. To understand the relationship between OM persistence, bioavailability, and adsorption, enzyme activity experiments should be moved away from traditional batch experiments conducted with phases dispersed in a slurry. Instead, investigations must consider and preserve the three-dimensional microstructure of natural mineral-organic interfaces <sup>46,194</sup>.

Although the importance of various physicochemical variables in determining the magnitude of OM adsorption to select phyllosilicates, metal oxides and carbonates are well understood, the

influence of these variables on the rates of adsorption and desorption is not. In addition, most OM sorption and desorption kinetic studies have been conducted on clean minerals exposed to aqueous phase OM. The kinetics of sorption onto realistic soils and sediments, where the newly offered adsorbate could compete with and actually induce the release of previously adsorbed organic compounds, have been studied on rare occasions <sup>86</sup>, presumably because of the difficulty in unequivocally attributing resulting observations to certain mechanisms when the sorbent is complex <sup>195</sup>.

## [H2] Adsorption as a framework-stabilizing process

In addition to localizing the adsorbed substrate within a pore network, adsorption has the potential to significantly influence the stability of this network, thus influencing the microbial dynamics outlined above <sup>196</sup>. For example, investigations of sediment dynamics demonstrate that fine-grained minerals and mineral-OM assemblages have a controlling influence on the cohesive nature of sediment. Detailed understanding of the relevant relationships, unfortunately, remains elusive due to complexity of interparticle interactions in assemblages of fine-grained minerals. Even in the absence of OM, colloidal interactions [G] involve a variety of interaction mechanisms with different sensitivities to distance and particle shape and charge, such that different interactions to predominate in different systems <sup>197-199</sup>. In particular, for layered minerals, the combination of a long-range osmotic repulsion and a long-ranged attraction due to configurational entropy enables the formation of extensive aggregates with relatively dilute yet cohesive gel-like structures <sup>200,201 202,203</sup>.

In the presence of OM, the stability of the mineral framework is further stabilized <sup>196</sup>. One conceptual model that could shed light into this stabilization is that the interaction of OM with mineral surfaces, at the OM abundances that exist in most soils and sediments, should be somewhat analogous to a wetting phenomenon. This analogy to wetting, in particular, is qualitatively consistent with the existence of either patchy or uniform distributions of OM in different systems <sup>16,204-206</sup> (although patchy distributions can partly reflect microbial lifestyles <sup>207,208</sup>, they are also observed in studies examining the uptake of dissolved organic matter on pristine mineral surfaces in abiotic conditions as well as in soils <sup>135,209</sup>), with observations of preferential uptake of OM on rough surfaces or in colloidal aggregates <sup>79,207</sup>, and with the tendency of OM to induce lasting mineral aggregation <sup>94,96,210</sup>. It also is consistent with the hydrophobic character of the siloxane surface of phyllosilicate minerals in the absence of surface charge <sup>211,212</sup> or even, in the case of smectite, in uncharged surface patches resulting from the non-uniform distribution of isomorphic substitutions <sup>62,67</sup>. In the case of oxides, studies indicating OM fractionation during uptake by Fe oxides suggest that the uptake of an initial 'contact layer' plays an important role in enabling mineral wetting by OM <sup>76</sup>. A number of other concepts developed in studies of wetting phenomena could prove useful in studies of mineral-OM interactions, including theoretical representations of contact angles and thin films <sup>213-215</sup> and descriptions of the impacts of biosurfactants on microorganism distribution, carbon cycling, and bioremediation 45,216,217.

### [H2] Mineral-catalyzed polymerization

There is substantial interest in the ability of minerals to both induce molecular growth (such as polymerization and large molecule formation) and contribute to molecular fragmentation, as it has implications for the global carbon cycle <sup>218,219</sup> and in prebiotic chemistry <sup>220</sup>. The fundamental ability of minerals to modify organic compounds has been confirmed in laboratory settings <sup>220-222</sup>, but the catalytic synthesis of novel organic compounds has not yet been observed in the environment. To investigate this phenomenon, observations need to occur simultaneously on extremely small spatial scales but on very long time scales, invoking serious experimental challenges. In the environment, mineral-catalyzed synthesis could occur but could be counterbalanced by mineral catalysed disassembly as soon as the newly made product diffuses away from its compartment of origin. Moreover, the same type of mineral can assume opposite functions in the microbial conversion of adsorbed substrates <sup>223</sup>. Directing greater research effort at the quantitative relevance of these phenomena in natural systems would be particularly useful to settle the ongoing debate regarding the importance of mineral-induced catalysis in the environment.

## [H2] Redox and ROS

In direct opposition to the expectation that minerals act as stable repositories for OM, the close association of mineral and organic compounds also enables electron transfer and coupled redox reactions. It is now well established that a plethora of microbial taxa and biochemical pathways couple the decomposition of organic matter to the oxidative or reductive transformation of minerals. Yet, despite tremendous progress in understanding the biogeochemical underpinnings of mineral-dependent life, the taxonomy of, enzymatic machinery employed by, and ecological controls on mineral-transforming microbes remain active and essential areas of inquiry. Importantly, the thermodynamic and kinetic constraints on operative metabolisms within natural soils and sediments remain poorly constrained, and yet are essential components for the efficacy of qualitative and quantitative biogeochemical models.

Further, despite some direct and indirect evidence of abiotic reactions mediating carbon transformation and mineralization at mineral surfaces, investigations of these interactions are sparse and, for those that have been conducted, conclusions are often based on correlations and/or anecdotal evidence. Moving forward, systematic and targeted investigations specifically linking OM cycling with direct and indirect reactions at mineral surfaces is needed. These investigations will undoubtedly require new methods and approaches to specifically probe the OM-mineral interface and track reaction products, such as incorporation of isotope labels and/or utilization of sophisticated imaging and spectroscopic techniques.

Acquisition of a clearer understanding of ROS-mediated carbon decomposition in soils and sediments has been hindered by several obstacles, including the difficulty in measuring short-lived ROS and identifying the fate of ROS among the numerous potential decay pathways within complex matrices. In fact, OM competes with numerous other ROS sinks, such as metals,

chloride, bromide, or carbonates <sup>165,170,224</sup>, and thus the contribution of ROS to OM cycling will undoubtedly vary with local geochemical conditions, and between freshwater and marine systems. Further the promiscuity of ROS such as O<sub>2</sub>\* and H<sub>2</sub>O<sub>2</sub> to act as both reductants and oxidants depending on the reactants and aqueous conditions leads to complex reaction networks even within relatively simple systems. Investigations of dityrosine formation within marine waters recently highlighted the potential for ROS-induced polymerization as a means to generate less bioavailable "humic"-like fluorescent DOM in the ocean <sup>225</sup>, indicating that ROS can mediate polymerization in addition to carbon mineralization. Thus, given the emerging recognition of minerals as sources of high ROS fluxes within the subsurface and the well-known role of ROS in organic carbon transformations, mineral-based ROS serve as a new frontier in understanding the carbon cycle.

Disentangling the complex network of mineral-based redox reactions is a necessary next step to understanding the controls on carbon processing and availability within soils and sediments. These reactions will likely be most relevant at redox interfaces and in fluctuating redox environments. Under these conditions, fresh precipitates free of significant surface coatings and prior to ripening will have increased surface reactivity. Further, (re)generation of mineral-bound and aqueous redox reactants will lead to new mineral reactive sites and (re)generation of ROS and other reactive intermediates (such as reactive DOM). Teasing out the relative contributions of light-dependent and –independent reactions on ROS formation and DOM decomposition is also required within sunlit environments. Thus, an improved understanding of OM cycling in soils and sediments also requires quantification of fluxes of solid-phase and aqueous reactants across light and redox gradients, likely requiring the use of spatially resolved techniques and thermodynamic and kinetic modeling to tease out the network of reactions at play.

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## 1316 Author contributions

All authors participated in developing the concept. Figures were developed by E.C. (Figures 1 and 5), M. K. (Figure 2), I.C.B. (Figures 3 and 4), S.M. (Figure 6), C.M.H. (Figure 7) and N.N. (Figure 8). All authors contributed to writing and editing.

## **Competing interests**

The authors declare no competing interests.

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#### **Key points**

- Minerals enable the compartmentalisation of soils and sediments into small yet clearly
  delineated spaces such that different chemical, ecological and evolutionary processes can
  occur concurrently within a larger system context.
- Organic matter (OM) attachment to mineral surfaces is dynamic, sensitive to interfacial energies and topology, and exhibits features reminiscent of a partial wetting phenomenon.
- Mineral-derived reactive oxygen species represent overlooked but undeniably key reactants in the oxidation and transformation of OM within soils and sediments.
  - Correlations between OM and fine-grained minerals, though generally interpreted as reflecting the impacts of minerals on OM, could additionally reflect impacts of OM on mineral nucleation, growth, and transformation.
  - Depending on system logistics and environmental setting, the same type of mineral could act as a sorbent, chemical reactant and catalyst for associated OM, enabling a vast portfolio of potentially opposing outcomes.

 Assessments regarding the fate of OM in the environment should not be derived from correlations with single predictor values, such as abundance of a certain mineral phase or specific surface area.

## Figure Captions

Figure 1 Organic-matter cycling in soils and sediments and mineral-organic matter interactions a Historic representation of organic matter cycling in soils and sediments. Major organic matter (OM) pools and fluxes are represented as black boxes and arrows. Blue arrows represent processes and interactions discussed in this Review. b Multiscale representation of mineral organic interactions. At the largest scale (Tier 1), the main function of the mineral matrix is to delineate compartmentalized space. As a consequence of compartmentalization, a number of static and dynamic constraints are imposed on smaller scales (Tier 2 functions). These constraints then determine the intensities and rates at which molecular scale biogeochemistry could be able to proceed (Tier 3 functions).

**Figure 2. Organic Multifunctionality.** Variation of functional group combinations on organic compounds allows for diverse modes of interaction with mineral surfaces.

**Figure 3. Key properties of fine-grained minerals and related solids**. The material referred to as allophane is a short-range-ordered (SRO) phyllosilicate with imogolite-like local structure and highly variable stoichiometry; its distinguishing features are low crystallinity and transmission electron microscopy (TEM) images suggestive of hollow spheres 3.5 to 5 nm in diameter, though multiple studies have noted that the interpretation of three-dimensional structure from two-dimensional TEM image is ambiguous <sup>95</sup>. Data based on Refs<sup>34,66,94,96,226-229</sup>.

**Figure 4. Size, shape, and global distributions of minerals. a**| Size and shape of minerals discussed here. **b**| Global map showing the most abundant fine-grained mineral in the subsoil (0.3 to 2 m depth) as a function of location, and average relative abundance of different fine-grained minerals in the upper 2 m of soil averaged over the Earth's land surface with the exception of organic-rich soils (such as mollisols) and ice-covered regions<sup>33</sup>. Part X is adapted from ref <sup>33</sup>, CC BY 4.0 (https://creativecommons.org/licenses/by/4.0/).

**Figure 5. Organic ligands at mineral interfaces.** Mineral interfaces occur at a variety of spatial scales and topography in soils and sediments, where a diversity of primarily low-molecular weight, amphiphilic organic ligands may reach the solid-solution interface. Once proximal,

ligands can bind to mineral surfaces through one or multiple complexation mechanisms based on surface charge distribution and structural reactivity.

Figure 6. Molecular mechanisms of OM reactions at mineral-water interfaces. Cartoon at the top shows a mineral surface showing monolayer and multilayer (3-dimensional) sorption of small and large OM. Molecular structures of organic molecules at mineral-water interfaces are: it cation bridging, ii: H-bonded or Lifshitz -Van der Waals interactions, and iii: direct covalent interactions. Expanded views of OM interactions and expected trends for different mineral structures in soils and sediments are shown at the bottom. The nature of chemical reactions are (as numbered in the figure): 1: e-transfer and electrolytic breakdown of OM; 2: larger OM formation from nucleophilic addition and radical mediated polymerization; 3: hydrolytic breakdown of molecules from pH changes at the interface; 4: heterogeneous OM oxidation from adsorbed O<sub>2</sub>; and 5: chemical changes to substrate showing electron transfer and reduction of substrate atoms and changes to the coordination environments of neighboring atoms of substrate surface. Based on their differences in chemical reactivity towards OM, we further classify minerals presented above into 3 classes (I-III). Expected trends for OM behavior are included.

**Figure 7**. **Mineral-induced organic carbon redox pathways**. **a**| Mineral induced oxidation, which represents direct coupling. **b**| Mineral-derived aqueous Fe(II), which involved both direct and reactive oxygen species (ROS) mediated reactions. **c**| ROS mediated reactions. In **b** and **c**, the fate of mineral-derived ROS includes reaction of ROS with organic carbon (OC) or transformation of one ROS to another (dismutation of superoxide to hydrogen peroxide, formation of hydroxyl radical from ferrous iron and hydrogen peroxide via the Fenton reaction) – for simplicity, all these fates are not shown here. The illustrations are meant to show general processes, not mechanistic details, and the list of reactions is not exhaustive.

**Figure 8. Compartmentalization and mineral-organic matter-microbe interactions**. Top panel shows structure with high clay content that is compartmentalized and relatively constant over time and lower panel shows structure with low clay content that is relatively dynamic over time. The compartmentalization and constant micro-environmental conditions results in greater niche differentiation, which allows a greater microbial diversity to co-exist through reduced competitive interactions <sup>230,231</sup>. The magnified insets show potential evolutionary consequences in microsites that form constant habitats. The capacity to use organic substrate present is subjected to strong selective pressure and preserved but other pathways could be lost (insets i. and iii.). Less competitive species do not compete for organic substrate present, but exploits metabolic by-product of more competitive species, resulting in a better use of available resources by the microsite community <sup>232</sup> (insets ii. and iv.).

- Adsorption An increase in the concentration of a dissolved substance at the interface of a condensed and a liquid or gaseous phase due to the operation of surface forces.
  - **Catalyst** A substance that increases the rate of a reaction without modifying the overall standard Gibbs energy change in the reaction
    - **Chemotrophic** the ability to use electron donors other than photons for the synthesis of organic compounds containing reduced carbon
  - Coagulation the formation of aggregates from a fluid colloidal system
  - **Colloid** molecules or polymolecular particles dispersed in a medium that have at least in one direction a dimension roughly between 1 nm and 1 µm
  - **Colloidal interactions -** interactions that are enabled when particles become so small (equivalent diameter < 1-2 micron) that surface borne electric forces between particles can effectively control their behavior in a suspension (for instance, prevent them from settling)
  - **Compartmentalization** the division of a system into multiple subsystems with well defined boundaries that provide a certain degree of process autonomy
  - **Coulombic interactions** interactions that result from the electric force between two charged entities
  - **Crystal facet -** a flat plane on a crystal
  - **Crystal growth -** the addition of new atoms into the characteristic arrangement of the crystalline lattice, releasing thermal energy (enthalpy of crystallization)
  - **Depolymerization -** the disassembly of a polymer into its constituent monomers or into a mixture of products
  - **Dispersion** A system in which particles of colloidal size of any nature (solid, liquid or gas) are dispersed in a continuous phase of a different composition (or state).
  - **Fine-grained fraction** mineral grains with an average diameter smaller than 50/63 microns, depending on classification system used
  - Heterotrophic the ability to derive nutritional requirements from complex organic substances
  - Intra-particle region any part of a particle that is not participating in surface reactions
  - **Interfacial energy** excess free energy or work associated with the interface between two phases, per interfacial area
  - **Ligand** any atom or molecule attached to a central atom, usually a metallic element, in a coordination or complex compound; if regarding part of a polyatomic molecular entity as central, then the atoms, groups or molecules bound to that part are called ligands.
  - **Metabolic dependency** a form of adaptation that leads to the absence or loss of the ability to synthesize a certain metabolite essential for the organism, usually in response to an abundance of said compound in the environment
  - **Microsite** a clearly delineated space within an environment with unique conditions or features in which specific microbial processes can occur.

- Mixotrophic deriving carbon and energy from a mix of different sources, typically a combination of inorganic and organic compounds
  - **Nanowire** proteinaceous appendage produced by microbes, particularly bacteria, that is electrically conductive
- Nucleation the process by which nuclei are formed in solution

- **Orientational freedom -** the absence of any physical restrictions to the movement and arrangement of a compound
- **Passivated** a surface that is unreactive owing to alteration or from the formation of a thin inert coating
- **Photochemical lability -** the tendency of a compound to undergo a chemical reaction when exposed to light
- **Phototrophic** ability to capture photons as energy source for the synthesis of organic compounds containing reduced carbon
- **Poorly crystalline** an operational term to distinguish crystalline structures with short range order from others that exhibit order over longer distances
- **Reactive oxygen species** ROS are short-lived oxygen-bearing molecules with half-lives that range from fractions of seconds to days, including hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), superoxide (O<sub>2</sub>·-/HO<sub>2</sub>), hydroxyl radical (HO•), singlet oxygen (¹O<sub>2</sub>), and carbonate radical (CO<sub>3</sub>·-).
- **Ripening** physical and/or structural alteration of a mineral to obtain a lower surface free energy and more energetically favorable state
- **unsaturated soil** a (soil) pore system that is only partially filled with water is unsaturated; a pore system entirely filled with water is considered saturated.
- **Short-range ordered** the regular and predictable arrangement of atoms over a very short distance; in crystals, order does not persist over distances of more than a few nanometers and often extends over the distance of just a few bond lengths; Short range ordered minerals are often also referred to as poorly crystalline minerals.
- Solution a homogeneous phase that results from the mixing of two (or more) phases
- **Steric constraints** factors or effects that either prevent the adoption of a certain spatial orientation that would be required for the reaction to proceed unhindered
- **Steric enhancement -** factors or effects that facilitate the adoption of a certain spatial orientation that would be required for the reaction to proceed unhindered
- **Xenobiotic compound** a substance that is foreign to a given natural environment or ecosystem; usually means that organisms in the system lack adaptations for the metabolic processing of a xenobiotic compound

## Table of contents summary

Minerals and organic matter interact in soils and sediments, impacting biogeochemical cycling and ecosystem functioning. This Review describes the major and emerging environmental mineral-organic interactions observed, and their implications for organic matter persistence.