A Review of Dissolved Organic Matter Transport Processes Affecting Soil and Environmental Quality

S. K. Deb and M. K. Shukla*

Department of Plant and Environmental Sciences, New Mexico State University, MSC 3Q, P.O. Box 30003, Las Cruces, NM 88003

Abstract

Dissolved organic matter (DOM) affects several processes in soil and water including nutrient cycling, soil and water pollution and CO₂ flux between the soil and atmosphere. The aim of this review is to collate and synthesize the literature on the transport processes of DOM in soil. The DOM normally comprises of only a small fraction of soil organic matter (SOM) and originates mainly from the decomposition and solubilization of SOM, which is accumulated on soil surface or soil profile from plant residues and additions of organic amendments such as animal and poultry manures and other biosolids. The DOM is one of the most reactive and mobile SOM fractions and has a major influence on biogeochemical processes in both terrestrial and aquatic environments. Terrestrially borne DOM is subjected to microbial decomposition, photodegradation and adsorption on soil mineral surfaces. It is sorbed on mineral surfaces and high adsorption capacities of clay minerals and oxides for DOM sorption are demonstrated in laboratory studies. However, these high sorption values are not reproduced in limited field studies. Similarly, a few data available on the transport of DOM through macropores also demonstrate the limited control of sorption on DOM retention in soil profile. Thus, there is a need to further investigate the physical and chemical protection mechanisms, as well as the biodegradability of DOM shown in laboratory studies. There is an increasing need to clearly understand the formation, fate and transport of DOM at field scales. The environmental factors such as precipitation and temperature, land use change, land management, and biological factors have profound and discrete influences on DOM dynamics in soil profile. Future research efforts must focus on the assessment of the influences of these factors by conducting field studies in different climatic zones, soils, and land use and management systems.

Keywords: Dissolved organic matter; Dissolved organic carbon; Sorption; Biogeochemical processes; Nutrient cycling; Anion exclusion; Mineralogy; Climate change; Water quality

Introduction

The importance of soil organic matter (SOM) in supplying nutrients, contributing to cation exchange capacity (CEC), and improving soil structure is well recognized. When in soil solution and pass a filter pore size of 0.4–0.6 µm, SOM is known as dissolved organic matter (DOM) [1,2]. The organic matter in terrestrial and aquatic environments primarily comprises of two operationally defined phases: (i) particulate and (ii) DOM; and both are complex mixtures of many organic compounds and humic substances. Almost all ecosystems contain DOM and the size of its pool can vary at multiple scales ranging from amino acids to complex high molecular weight SOM. The DOM has an important influence on soil forming processes [3,4], mineral weathering [5], and pollutant transport [6,7]. When present in soil solution, DOM plays a vital role in many biogeochemical processes, and transport and fate of several contaminants including heavy metals in soil, surface water and groundwater [7,8]. Recycling of elements (i.e., C, N, P and S) in the soil is strongly influenced by DOM. Equally affected is the export of nutrients to surface and groundwater resources. The DOM also influences the transport behavior of pesticides and herbicides through the vadose zone. Several natural systems behave as a three-phase system where contaminants present in the vadose zone are sorbed to soil matrix, sorbed to DOM, or present as a freely dissolved substance. The relative proportion of contaminant in each of the three phases is important and determines the transport behavior, fate and characteristics of the solute with respect to migration and surface and groundwater contamination. For example, according to Carter and Suffet [9], Chiu et al. [10], and Hassett and Milicic [11], DOM increases the solubility and decreases the volatility of certain organic contaminants, and thus increases their transport through the soil profile. Hydrophobic contaminants in natural waters exist in two states, i.e., one that is truly dissolved and one that is bound to DOM [12]. While this binding decreases a hydrophobe's apparent volatility, bioavailability, and attachment to particles and increases its apparent solubility, binding also brings a hydrophobic contaminant into a chemical environment that is distinct from water and thus can affect its reactivity.

Estimates of the impact of DOM on terrestrial C balance mostly rely on the fluxes measured in river waters. The flux of DOM in river water ranges from 1 to 10 G C m⁻² y⁻¹ for many ecosystems [13]. Therefore, these fluxes are considered a small or insignificant component of the ecosystem C budget as compared to the C fluxes associated with primary productivity or heterotrophic respirations in terrestrial ecosystems [14]. However, there are several reasons to believe that DOM fluxes are more important than that suggested by usually small river fluxes. Over time scales ranging from few months to several years, a consistent loss of DOM from the soil profile can reduce the capacity of ecosystems to sustain primary productivity [15]. The DOM fluxes through soil profile can be much larger than the fluxes associated with runoff that eventually ends up in stream water (Table 1 and Table 2). Within a soil profile, DOM dynamics is an important mechanism involved in soil formation. Therefore, it is of great importance to

*Corresponding author: Manoj K. Shukla, Department of Plant and Environmental Sciences, New Mexico State University, MSC 3Q, P.O. Box 30003, Las Cruces, NM 88003, Tel: 575-646-2324, Fax: 575-646-6041, E-mail: shuklamk@nmsu.edu

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The DOM dynamics has been extensively studied in numerous laboratory experiments using disturbed soil samples, but few studies have been conducted under field conditions [17]. Since 1970s, several attempts have been made to understand the behavior and transport dynamics of DOM through the soil profile. Therefore, the objectives of this review are to: (1) collate and synthesize the available information on DOM processes and dynamics, (2) review the biogeochemistry of DOM in the vadose zone, and (3) identify and prioritize future research needs.

**Constituents of Dissolved Organic Matter**

The DOM is a complex mixture of numerous organic compounds and humic materials in different proportions. The DOM found in streams, rivers or in soil solution is from plant biomass, litter, humus, roots, amendments applied on land surface, etc. Chemical identification of the constituents of DOM is possible but limited to a small fraction of low molecular weight substances such as organic acids, sugars and amino acids [1]. Most common compounds in DOM are: amino acids, carbohydrates, hydrocarbons, and aliphatic and aromatic acids (Figure 1). A large portion of DOM in soil solution consists of humic substances with average molecular weight of 1,000 [18]. Organic colloidal material with average molecular weights ranging from 2,000 to 100,000 may also be present as humic and lipid aggregates. Operationally, DOM is defined as the organic molecules of different sizes that pass through a filter of 0.2 µm or 0.45 µm [2], which can pass through 0.45 µm filter. A 0.2 µm filter is used when some organic or inorganic colloidal materials is present (i.e., organic particulates), which can pass through 0.45 µm filter. Major structural components in freshwater DOM include carboxyl–rich alicyclic molecules (CRAM), heteropolysaccharides, and aromatic compounds [1]. As suggested by Lam et al. [20], the bulk of freshwater DOM is aliphatic in nature, with CRAM derived from cyclic terpenoids as well as material derived from linear terpenoids.
Origin of Dissolved Organic Matter

Despite several studies undertaken since 1970s, origin of DOM is still not very clear and contradictory accounts exist in the literature. McDowell and Likens [22] and Zsolnay [2] propose that humified organic matter or leaching and microbial decay of humus is largely responsible for the DOM production in soil. This argument is mainly due to the presence of relatively high proportion of humus than litter in the soil. In contrast, Qualls and Haines [23] have reported high amounts of simple sugars and nonhumic–bound polysaccharides in DOM solutions and thus proposed a high contribution of recent litterfall in DOM production. The Qualls and Haines's [23] argument was supported by Qualls et al. [24] for a deciduous forest ecosystem where highest increases in DOM fluxes were reported in the upper part of forest floor or litter layer. As reviewed by Kalbitz et al. [17], there remains a considerable discussion and uncertainty in the literature regarding the importance of fresh versus older SOM as sources of DOC in forest floor soil solution. While studies on the chemical composition of DOM (e.g., [25]) suggest that most DOM is an end product of the microbial metabolism, the fresh litter also contributes significantly to the production of DOC [26].

Ultimately, as stated by McDowell [27], the source of virtually all DOM in soils is photosynthesis, which includes both recent photosynthetic (throughfall, leaf litter, root exudates, decaying fine roots) and the leaching or decomposition of older, microbially processed SOM. As DOM ranges in age from hours to days, up to decades to thousands of years, the range of sources can be equally varied [27–29].

Measurement of Dissolved Organic Matter

The DOM fluxes have been measured in rivers, streams, ponds, lakes and in the soil profile [13]. Most studies are carried out on discrete catchments ranging in size from 1 to 50 km². The concentrations of DOM and other nutrients and chemicals are measured both at upstream and downstream levels and effects due to land use and management are assessed by the comparisons of paired catchments [30]. Some other approaches to measuring DOM include: (i) assessment of total DOM or nutrient export by analyzing water samples collected at the lower reaches of a river or catchment [31,32], and (ii) collecting and analyzing samples from representative sections of an entire river system [33–35]. Most of the samples collected from rivers can be termed as instantaneous grab samples. The frequency of water sampling mostly depends on the size of catchment and varies from twice daily, daily, bi-weekly, weekly, bi-monthly, monthly to event basis (i.e., during high or low flows). Number of samples can also increase with increase in river flow [22,36].

DOM solutions in soils are collected using a number of devices. For laboratory studies, DOM is extracted using centrifuges in batch experiment studies or effluent solution is collected from the bottom of soil columns. Some laboratory column studies also employ suction cups for DOM collection. McDowell et al. [37] evaluated the comparability and reproducibility of batch, kinetic and bioreactor methods to measure biodegradable fraction of dissolved organic C (BDOC) for solutions from a variety of terrestrial sources such as throughfall, soil solution and soil extracts. Their results suggested that the precision of various BDOC methods was similar (5-15%) across a broad range of BDOC (12–56% of DOC) [37]. For fields, which are not tilled, several devices can be installed to collect leachate from vadose zone. Some of these devices are: suction cups, piezometers, sampling wells, and tension or zero tension lysimetry [22,38,39]. In tile drained fields, samples can be collected on a flow proportional basis or volume average basis. In general, quality of collected solution sample depends upon the device used and, therefore, results also often technique dependent. A comparison of DOC concentrations in soil solutions using a tension–free collector, a tension Prenart collector and a tension Rhizon collector sampling methods was made over a range of soil types [40]. The DOC concentrations in soil solutions were significantly higher with tension–free collectors than with tension devices. The lower concentrations obtained with the tension devices were more likely due to artifacts associated with sorptive removal of DOC by the collector or with localized soil compaction induced by the applied vacuum, as well as the influence of tension on the source of soil water from different pores [40]. Tension–free collectors were more representative of free–flowing water through the organic horizon, which is the key to representing DOC fluxes through topsoils.

Chantigny et al. [41] presented the general procedures for collection and analysis of soil DOM, including selected procedures for analyzing C and N concentration, key spectroscopic and chemical properties, and biodegradability. Different extraction procedures can result in different DOM samples. As suggested by Zsolnay [42], even minor alterations in extraction protocols can have a serious impact on DOM yield. Understanding the phase distribution of different contaminants in aquatic systems is necessary to determine which factors regulate binding of these contaminants to DOM. The feasibility of solid phase extraction (SPE) and stir bar sorptive extraction (SBSE) for the determination of eight polycyclic aromatic hydrocarbons (PAHs) in run–off water samples was evaluated by García–Falcón et al. [43]. Both the SPE and SBSE were appropriate to determine PAHs in run-off waters. While the SPE was suitable for the determination of overall PAHs, the SBSE was used for the determination of free PAHs. The combination of both extraction techniques could be used to estimate PAHs bound to DOM. Once the fraction percentage the free concentration represented was estimated, the only use of SBSE...

Figure 1: Size range of various carbon compounds in natural waters. F: Fulvic acid, H: Humic acid, FA: Fatty acid, CH: Carbohydrate, AA: Amino acids, HA: Hydrophilic acids, HC: Hydrocarbons (modified from Thurman [18]; Hope et al. [13])
was proposed for the estimation of all PAHs fractions (free, bound to DOM, and overall). The determination of PAHs in peat samples is complicated due to the high content of organic matter in peat, which affects both extraction efficiency and analytical quality [44]. Rey-Salgueiro et al. [44] reviewed and addressed the most relevant analytical methods for determining PAHs in peat, as well as evaluated three different extraction procedures, such as ultrasound-assisted solvent extraction (UASE), shaking and pressurized liquid extraction (PLE). The UASE extraction method using hexane:dichloromethane (80:20) as extractant was robust enough to determine the selected PAHs in peat samples. As suggested by Salgueiro et al. [44], the UASE did not demand sophisticated equipment and long extraction times. In contrast, the PLE involved sophisticated equipment and showed important variations in the results.

Processes Affecting Dissolved Organic Matter Fluxes

The processes associated with DOM fluxes can be subdivided into several categories: (i) atmospheric deposition and production of C, (ii) seasonal effects, (iii) climate and geology, (iv) transport and export to streams or groundwater, and (v) adsorption and storage in the vadose zone [13].

Seasonal effects and geology

Atmospheric inputs of DOM consisting of both dissolved organic C and dissolved inorganic C are low. The concentration of DOM in precipitation ranges between 1 and 10 ppm [18,45] and is usually small compared to internal fluxes. However, some studies have reported concentrations in atmospheric deposition being equal to or more than those in the exported effluent [22,46]. Concentrations of dissolved inorganic C (DIC) in precipitation are generally very small [47,48].

Temperature and precipitation affect the production of allochthonous and autochthonous organic C. Several studies have reported high maxima for DOM concentrations in soil solution and CO₂ levels in soil air during summer [16,49–52]. Climate and geology have major influence on DOM export through a catchment. Climate impacts primary production and decomposition rates of DOM. In arctic, alpine and arid regions of the world, DOM productivity and export are lower than those in tropical and temperate regions. The terrestrial export of DOM varies with land use under the influence of climatic variability [53]. Climate and vegetation have important role on surface runoff and soil erosion whereas geology has major influence on geomorphology and rates of physical and chemical weathering [34,54,55].

Transport to streams or groundwater

As precipitation falls through the canopies, on to the forest floor and soil surface, concentration of DOM progressively increases [16,38]. The part of DOM runs off as overland flow towards a ditch or a stream and remaining water infiltrates into soil. During the overland flow, part of DOM is lost to depressional storage, adsorption to soil particles, and interception due to vegetation (Figure 2). All these processes reduce the concentration of DOM in overland flow. The concentration of DOM increases as the infiltrating water moves through the soil profile. DOM is leached in the soil profile and is either stored in the deeper soil profile, adsorbed in the soil or joined to the shallow groundwater. A part of infiltrated DOM can also move laterally and join the open ditch or stream. The role of surface water–groundwater interfaces in the retention and transport of DOM appears highly variable and remains challenging at different scales (i.e., from stream reach scale to the regional and global scales) [56].

Mobility of agrochemicals particularly pesticides in soils plays an important role in the fate and transport of contaminants. A comprehensive review about the mobility and degradation of pesticides in soils and the pollution of groundwater resources was provided by Arias–Estévez [57]. The influence of organic matter and clay contents on the sorption/desorption and degradation of pesticides and their access to groundwater and surface waters were discussed. As described by Arias–Estévez [57], the processes, which directly control the transport of pesticides within the soil and their transfer from the soil to water, include a variety of complex dynamic physical, chemical and biological processes including sorption–desorption, volatilization, chemical and biological degradation, uptake by plants, run-off, and leaching. While the relative importance of these governing processes varies with the chemical nature of the pesticides and the properties of the soil, the degradation and sorption processes are crucial.

Physical and chemical processes in the soil

The origin and production of terrestrial C is derived from vegetation and soil. Inputs from vegetation are from biomass, which consists of shoot, fruit, leaf and roots. It can also be from the leaching of C from live vegetation and litter. Inputs from soil are from SOM, root exudation, microbial metabolism, and erosion and leaching of SOM [18]. DOM concentrations measured in the vadose zone are the net result of processes that remove and/or release DOM. The process that remove DOM from soil solution are adsorption or decomposition and the processes that release DOM are the leaching from litter, organic wastes or desorption from solid phase (Figure 3). This physical and chemical alteration of SOM and the formation of soluble humic substances are generally assumed to be dominated by biota. However, several reports have indicated that the DOM concentrations in soil solutions are mainly controlled by abiotic processes such as desorption, and dissolution from the pool of potential DOM [58,59].

Transport through the soil profile

A number of studies have been conducted on the fate of DOM during transport through soil profile. Several studies have reported a
decrease in DOM concentrations and fluxes as water flows through the soil profile largely due to the sorption of organic acids in the B horizon, which occurs as part of podzolization process [16,18,60]. Sorption is often the primary cause for declining DOM concentration in soil profile of forested watersheds [3,16,22,38,60–62]. Similar observations for a grass land system have been reported by Hornung et al. [63].

Forest soils receive high inputs of PAHs (and other organic contaminants) because of the large intercepting surface constituted by foliage that eventually falls and contributes to the accumulation of PAHs in the soil organic layer [64,65]. García–Falcón et al. [66] studied the levels of eight representative PAHs in the 1–5 cm layer of a peruvian woodland soil, which had undergone wildfire and compared with those measured in nearby and distant unburnt perurban woodland soils and in a distant unburnt rural woodland soil. García–Falcón et al. [66] reported that PAH levels at the burnt site fell along the months as the result of rainfall and the prevention of further input from the atmosphere by the overlying layer of wood ash with a very high PAH adsorption capacity, which did not itself appear to act as a source of PAHs. García–Falcón et al. [66] concluded that PAH transport may have been assisted by increased mobilization of PAHs associated with DOM due to an increase in soil pH caused by the leaching of alkaline ash components from ash.

The primary controls for potential and actual DOM concentration are biotic and abiotic in nature. The DOM concentrations in soil profile are primarily biotically controlled with favorable temperature changes increasing the mineralization in soil profile. However, biotic influences can be masked by the variations in pore water velocity through soil profile and may become more important with increasing time scales [17]. The DOM pool can be further divided into mobile and immobile fractions. Such a classification is mainly based on the pore size distribution of soil matrix [2,67]. The portion of DOM present in micropores is immobile and can interact with the mobile fraction by diffusion. On the other hand, DOM associated with macro and mesopores is mobile and is subjected to convective as well as dispersive transport through soil profile [17].

**Im mobilization**

Limited research has been carried out to quantify the impact of soil properties in DOM immobilization. The DOM consists of both hydrophilic and hydrophobic substances, but the potential for sorption in the vadose zone differs. The hydrophobic DOM generally has higher affinity for soils than hydrophilic [68]. The DOM can be immobilized in soil through complex interactions with the mineral surfaces [69,70]. Immobilized C may be stored within the B horizon for several hundred years [22,38]. The capacity of soil to adsorb DOM is related to the soil pH, mineralogy, antecedent SOM content and amount of Fe and Al oxides in the soil. The DOM sorption on mineral surfaces is strongly pH dependent, with higher sorption observed at lower pH [71]. Smaller reductions in DOM with depth are observed in peat soils than mineral soils because of the high antecedent soil organic carbon (SOC) concentration [50,51,72]. It is mostly assumed that adsorption of DOM on mineral surfaces is far more pronounced than reduction of DOM due to the decomposition [17]. Approximately 40–370 kg DOM ha⁻¹ y⁻¹ is retained in mineral subsoil with sorption being the dominant process of stabilization [73,74].

Gibbsite, especially near pH 5, has the potential to absorb large amounts of fulvic and humic acids from lake sediments [75]. Some other studies have shown that appreciable amounts of DOM can be absorbed on a variety of synthetic Fe oxides and hydroxides [70,76]. Phyllosilicates can also absorb a significant amount of DOM [77,78]. Tipping [79] reported greater adsorption potential for higher than lower molecular weight humic substances in some studies with lab synthesized goethite, hematite, and amorphous Fe gels.

**Mechanisms of Sorption**

Mechanisms of DOM immobilization with mineral surfaces are not clearly understood and mostly hypothetical in nature [71]. Anion sorption, ligand exchange, protonation, hydrogen bonding, cation bridging, and physical adsorption are some of the mechanisms responsible for the DOM sorption on mineral surfaces [80].

**Physical Adsorption**

The adsorption of DOM on mineral surfaces is related to the antecedent SOM content and mineralogy of soil. Jardine et al. [71] reported that large amount of DOM (50 to 70%) is retained by crystalline and noncrystalline Fe oxides and hydroxides from the soil with phyllosilicates, in the less than 2 µm clay fraction, accounting for the remaining DOM. Jardine et al. [71] observed that kaolinite, which had lower CEC than illite, exhibited a higher adsorption capacity than illite. With removal of Fe oxides and hydroxides from a soil, a significant reduction in the amount of DOM sorption is observed. The Fe minerals in the soil are basically hematite (α-Fe₂O₃) and maghemite (γ-Fe₂O₃) and their abundance increases with depth which is in accord with the increasing DOM sorption with depth. Adsorption of humic acids by hematite was also reported by Tipping [79] and Ho and Miller [81].

The sorptive protection of DOM in soil is primarily indicated by the positive relationships between clay content and SOM [82]. Further evidence is given by the close relationships between DOM and BET surface areas in subsoil horizons [83]. The adsorption hypothesis was further supported by the positive correlations that exist between surface area and SOC contents for sediment samples [84,85]. Soil samples, collected from the A horizons of sixteen states across US, were analyzed for BET specific surface area and SOC content. A linear relationship exists between specific surface area and SOC across a range of soil types (Figure 4) [83].

The contribution of clay minerals on the adsorption of DOM can be assessed by either using pure minerals such as kaolinite and gibbsite or treating the soil with citrate–bicarbonate buffer and sodium dithionite to remove the iron oxides. As much as 250 to 600 mg kg⁻¹ of DOM can...
be adsorbed on a fine loamy soil with Fe oxide removed, 700 to 2100 mg kg\(^{-1}\) on kaolinite and 700 to 1900 mg kg\(^{-1}\) in gibbsite [71]. Thus, clay minerals play an important role in the adsorption of DOM. Significant adsorption of DOM on gibbsite and kaolinite was also reported by Farfitt et al. [86] and Davis [77]. There can also be other potential adsorbents such as high activity (2:1) clay minerals [87]. In contrast, Jardine et al. [71] reported that 2:1 phyllosilicate illite adsorbed 85% less DOM than kaolinite.

Degradation of sorbed DOM on surfaces already occupied by SOM is generally more and the sorptive stabilization mainly limited to juvenile mineral surface [74]. In general, DOM sorption on mineral soil results in stabilization [88]. The main mechanisms for stabilization include the selective sorption of intrinsically stable compounds and strong chemical bonds to the mineral surface. Stabilization is also the result of physical protection and inaccessibility of SOM to microorganisms. However, stability of DOM is largest for molecules that adsorb on juvenile mineral surfaces [89]. Surface OH in gibbsite, goethite, and imogolite is replaced by carboxyl groups of fulvic and humic acid [86]. Humic substances are responsible for ligand exchange of water and surface OH\(^{-}\) from Fe oxides [79]. Kodama and Schnitzer [90] proposed that surface water in sepiolite is displaced through a nonionic hydrophobic adsorption mechanism.

The adsorption and desorption of agrochemicals in crop soils govern their fate. Although organic matter is the universal soil colloid that binds most herbicides, metallic hydrous oxides might also have some influence [91] studied the adsorption–desorption of three quaternary ammonium herbicides, i.e., paraquat (PQ), diquat (DQ) and difenzoquat (DFQ), on soils with different chemical–physical characteristics using a batch equilibration method before and after the following sequential selective dissolution procedures: removal of organic matter, and removal of organic matter plus free iron oxides. The adsorption on soils (based on the herbicide soil/solution distribution coefficients) was in the following order: PQ > DQ ≥ DFQ [91] concluded that the removal of organic matter from soils seems to leave free new adsorption sites for quats on the clay surface, which is no longer occluded by organic matter. As reported by [91], the amount and nature of the clay surface that remains available after the removal of single soil constituents is a critical parameter in determining the sorptive behavior of cationic contaminants.

**Anion Exchange**

Anion exchange reaction involving adsorption of inorganic anions by hydrous oxides is also known as non–specific adsorption [92]. Terms specific and nonspecific are also used to distinguish adsorption due to polar and nonpolar van der Waals forces. Below a pH of 8, aluminium and iron hydrous oxides generally have positive sites. Therefore, coulombic attraction causes organic anions to be attached to oxide surfaces (Figure 5). However, the adsorbed organic anion can also be exchanged with chloride or nitrate [69,93].

Anion exchange mechanism is operative during the adsorption of DOM on allophane as well as on mineral surfaces [71,80,87]. Anions such as sulfate and phosphate present in soil solutions complete with DOM for adsorption [79,80]. Contrary accounts are available about the affinity of DOM for soil as compared with sulfate. Kaiser and Zech [94] reported that DOM exhibits greater affinity for soil than sulfate while Jardine et al. [71] reported greater affinity of DOM for sulfate than soil. In contrast, Vance and David [95] and David and Zech [96] concluded that it is not clear whether sulfate and DOM compete for the same sites. Sulfate anions have strong electrostatic interactions with positive sites associated with soil and do not induce ligand exchange reactions where surface OH groups are replaced. In a study on sulfate adsorption on a kaolinite, it has been shown that the ligand exchange is not operative [97]. The contribution of anion exchange during DOM adsorption on a fine loamy (Paleudult) soil was also reported by Jardine et al. [71] who observed that as the concentration of sulfate is increased DOM adsorption in the soil is decreased. This is possible because sulfate eliminates all anion exchange reactions of DOM. In contrast, Vance and David [80] observed that DOM has greater affinity for soil than sulfate as presence of DOM in solution reduces the amount of sulfate adsorbed especially in spodosols soil horizons. Gobran and Nielson [98] also reported competition between DOM and sulfate for exchange sites and reported no sulfate sorption until sulfate concentration exceeded 15.6 mmol L\(^{-1}\).

**Ligand Exchange**

In addition to the anion exchange reactions, specific adsorption of anion by hydrous iron and aluminium oxides and kaolinite takes place (Figure 6). This reaction takes place when the anion penetrates the coordination shell of an aluminium or iron atom in hydroxide surface and gets incorporated with the surface hydroxyl layer [92].

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**Figure 4:** Relationship between BET specific surface area (SSA) and soil organic carbon concentration (SOC) for soil samples collected from 16 states across the United States for 12 different soil types (drawn according to data reported by Mayer [83]).

**Figure 5:** Anion exchange mechanism for interaction of humic acids with hydrous aluminium oxide surfaces. Note: R is the remainder of humic molecules which can also be associated with the surface. O is oxygen ion, H is hydrogen ion and C is clay. (redrawn from Greenland [69]).
anion exchange reactions, the incorporated anion in a ligand exchange reaction cannot be displaced by leaching with a non–specifically adsorbed anion [92].

**Protonation**

The organic molecules of basic amino groups can accept a proton from clay surface and may become cationic after adsorption at the clay mineral surfaces. Such a phenomenon is known as protonation [99]. Some of the factors influencing the ability of clay surface to donate protons are: nature of minerals and exchangeable cations, and water content of mineral. The hydrolysis of water associated with exchangeable metal cations (M) is given by the following reaction:

$$M(H_2O)_x^{n+} = [M(H_2O)_{x-1}]OH + H^+$$

where $n$ is positive charge of the hydrated cation. A hydrated Al$^{3+}$ is a better proton donor than hydrated Na$^{+}$. The reaction goes to the right as charge to radius ratios increases or water content decreases [99,100].

**Hydrogen Bonding**

Hydrogen bonding is an important mechanism of clay mineral and organic interaction. It can happen in two ways. A polar organic molecule forms a hydrogen bond directly with water because it cannot displace a water molecule solving a metal exchange cation [99]:

$$\begin{align*}
\text{M} & \quad \text{O} & \quad \text{H} & \quad \ldots & \quad \text{O} = \text{C} \\
\text{H} & \quad \ldots & \quad \text{O} & \quad \text{C} \\
\text{R} & \quad \ldots & \quad \text{R}
\end{align*}$$

The other type of H-bonding takes place between an organic cation on an exchange site and another organic molecule:

The proton may also be shared equally between the two amines and the H–bond will be symmetrical. Such type of bond is found on clay surfaces for certain amines, amides, and urea [101].

$$\begin{align*}
\text{NH}_2 & \quad \text{C} = \text{O} & \quad \ldots & \quad \text{H}^+ & \quad \ldots & \quad \text{O} = \text{C} \\
\text{NH}_2 & \quad \ldots & \quad \text{O} & \quad \text{C} \\
\text{NH}_2 & \quad \ldots & \quad \text{R}
\end{align*}$$

**Precipitation**

Precipitation is another mechanism, which influences the retention or immobilization of DOM in soil horizons. Precipitation of DOM may also contribute to the formation of stable SOM. However, this mechanism has not been investigated in details [88].

**Global Flux**

The oceans contain some 50 times as much carbon dioxide as the atmosphere, and small changes in the ocean carbon cycle can therefore have large atmospheric consequences. Various agencies have tried to estimate the DOM fluxes in rivers and those entering in to the ocean. The RiOMar (River-dominated Ocean Margins) workshop reported that estimates for the global flux of atmospheric C transported by rivers ranged between 500–700 Tg y$^{-1}$, of which approximately 35% is DOC, 20% particulate organic C (POC) and 45% DIC [102]. As reported by RiOMar [102], estimates of the particulate and dissolved fractions of riverine organic C ranged over a factor of 2–3, but global average values at POC/DOC ratios of ~1 (~250 Tg y$^{-1}$ each). According to the estimates of the Committee on Flux of Carbon to the Ocean 1981, about 50% of all global DOM entering in rivers ends up in the ocean. About 15% of total DOM in rivers is oxidized and 25% is stored in as particulate organic matter (POM) and never reaches the oceans. The total DOM export to the oceans by rivers can be estimated by extrapolating the annual C flux data from each river around the globe and are presented in Table 3 [13]. The total global C flux in the world is reported to vary from 0.3 × 10$^{11}$ kg C y$^{-1}$ to 10.0 × 10$^{11}$ kg C y$^{-1}$ [103–105]. The estimates of global C fluxes are 3 to 10 times smaller than the rate of increase in atmospheric CO$_2$, which is reported to range from 1.8 to 4.7 × 10$^{11}$ kg C y$^{-1}$ [106].

The role of continental erosion in the global C cycle and its influence on river C fluxes may change with changing climate particularly between glacial and interglacial cycles [107]. Future global warming may cause significant shifts in rainfall patterns and storm intensities [108], which in turn will directly affect the export of C from terrestrial environments.

**Dissolved Organic Matter and Colloidal Transport**

The fate and transport of colloids and associated anions can be strongly influenced by DOM [109]. DOM in soil is composed of humic and fulvic acids and is pH–dependent. Numerous studies have demonstrated that even small amounts of DOM increase the mobility of colloid–associated contaminants (e.g., phosphorus, hydrophobic organic compounds, radionuclide plutonium, agrochemicals, carbon nanotubes, lead, etc.) and colloid–sized pathogens through the hydrologic pathways [109–116]. Laboratory batch kinetic and isotherm experiments have demonstrated that DOM can increase the stability of colloid and nanoparticle suspensions and can facilitate the colloid transport.
and nanoparticle transport [117–119]. The impact of DOM on colloid mobility in saturated porous media was systematically examined in terms of pore water velocity and deposition kinetics [110,117]. Evaluations included the effects of monovalent versus divalent cation concentrations, ionic strength on attachment efficiency, and charge reversal by organic matter adsorption [110,120,121].

Arias–Estévez et al. [122] evaluated the sorption of PAHs to colloid dispersion of humic substances (HSs) in water. Notably, HSs can complicate many environmental pollutants such as PAHs, and the distribution of PAHs in the aqueous environment (free and bound to HSs) is important since it affects their stability and persistence. As reported by Arias–Estévez et al. [122], both steric hindrance and molecular interactions between the PAHs and the HSs’ aromatic structures might play an important role in their binding reactions. The distribution of PAHs in colloidal dispersions of HSs in water (i.e., their stability–persistence and possible bioavailability in the natural aqueous environments) was controlled by the hydrophobicity and the water solubility of PAHs. Arias–Estévez et al. [122] also emphasized that adsorption of the different PAHs to HSs could be modeled in terms of both of these driving factors. In a river basin partially devoted to vineyards that are regularly treated with copper–based grapevine fungicides, Fernández–Calviño et al. [115] examined the copper (Cu) levels in soils and river sediments. Soils differed relatively little regarding the distribution of Cu among various fractions; for instance, about 50% of total Cu content was generally bound to soil organic matter. Fernández–Calviño et al. [115] observed that the distribution of Cu in sediments was more heterogeneous. Sandy sediments had Cu distributions similar to those observed in soils; whereas in silty sediments with large inorganic colloids fractions, it was these amorphous colloids that bound the largest proportion of Cu.

Systematic interaction and transport of DOM and colloids in unsaturated soils have not yet been studied in detail. Only limited studies have examined the effect of DOM and pH on colloid transport in unsaturated porous media and the effect of colloid transport in alkaline DOM rich conditions [111,123]. In general, interfaces with air in unsaturated porous media promote colloid retention and organic matter of hydrophobic character preferentially fractionates to the air–water interface [124]. Morales et al. [124] examined the effect of DOM on colloid characteristics through changes in surface potential, adsorbed layer thickness, and mass of adsorbed organic matter. They conducted transport experiments with tandem internal process visualization for key constituents of DOM, humic and fulvic acids at acidic, neutral and basic pH and two CaCl₂ concentrations. Humic acid improved colloid transport significantly, while fulvic acid marginally affected transport [125]. Experiments with fluvic or no DOM promoted colloid retention at solid–water interfaces, while experiments with humic acid enhanced colloid retention at air–water interfaces due to partitioning of humic acid at the air–water interface and/or increased hydrophobic characteristics of humic–colloid complexes.

Summary

Dissolved organic matter plays an important role in many biogeochemical processes in soils and waters. It also plays an important role in elemental cycling and influences the cation leaching processes, mineral weathering and soil formation. The mobility of DOM in soils is controlled by its sorption to mineral surfaces. Several mechanisms i.e., anion sorption, ligand exchange, cation bridging, and physical adsorption, are postulated for describing DOM sorption on mineral surfaces. However, these mechanisms are not yet clearly understood and more research is needed to properly understand the processes associated with physical and chemical stabilization in soil. More efforts are needed to understand the steric effects of the silicate structure on orientation of organic absorbates. Equally important is to understand the effects due to structural and charge differences in clay, which may lead to different kinds of association with organic molecules. The degree of DOM sorption on mineral surfaces is dependent on pH, average molecular weight of DOM, antecedent C content of soil, and surface soil characteristics. Little information is available on the relationship between mineralogy and the chemistry of adsorbed organic matter; particularly, the quantification and evaluation of the different interactions with mineral surfaces (e.g., impact of ligand exchange, cation bridges, hydrophobic interactions, complexation reactions) need to be addressed. It is generally assumed that adsorption diminishes the biodegradability of DOM. However, this has not yet been quantitatively established. Most of the research on sorptive protection is based on the results from laboratory experiments, which generally disregard changing hydrological conditions. More efforts are needed to investigate the biotic and abiotic controls on DOM concentration and fluxes in soil horizons especially by conducting field experiments. Such an effort should be made in conjunction with proper accounting of flow through the soil profile. The temperature dependence of these controls is also important and needs to be further investigated. Further research is necessary to bridge the information gap between the physicochemical changes of DOM–colloid complexes and soil interfaces and the effect that these systematic changes have on the transport and fate in unsaturated soils (i.e., the vadose zone). While most studies over the past several decades focus on DOM dynamics in forests and grasslands or stream ecosystems, little attention has been paid to quantifying DOM fluxes in human–dominated systems such as agricultural soils or soils of heavily populated basins.

The interactive effects of climate change and dissolved organic matter is another important area needing further efforts. Climate change may reduce DOM concentrations in aquatic ecosystems, thereby exacerbating UV effects by changing the amount and flowpaths of DOM from upland and wetland ecosystems. Therefore, there is a need to conduct experiments through a combination of landscape and hydrological analyses, biogeochemistry, photodegradation, and biodegradation of DOM. Such research will greatly enhance our knowledge of the interactive effects of climate change, landscape watershed attributes, and DOM on aquatic ecosystems.

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