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Ferrihydrite coating reduces microplastic induced soil water repellency†

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Addition of microplastics (MP) to soil has the potential to increase soil water repellency. However, coating of MP with soil abundant substances e.g., iron compounds, can reduce this effect. Here, we tested if pre-coating or *in situ* coating of MP with ferrihydrite (Fh) reduces soil water repellency. We applied hotspots of pristine and coated MP (20–75 μm, PS and PET) to sand and imaged capillary rise *via* neutron radiography. Capillary rise experiments in wetting–drying cycles were conducted using water and Fh suspension. Pristine MP hotspots were not wettable. Capillary rise of water into coated MP hotspots differed in wettability depending on polymer type. While coated PS was still non-wettable, water imbibed into the coated PET hotspot. Capillary rise of Fh suspensions in wetting and drying cycles also showed varying results depending on polymer type. MP hotspots were still non-wettable and local water content increased only marginally. Our results indicate that Fh coating of MP changes MP surface wettability depending on polymer type and therefore counteracts the hydrophobic properties of pristine MP. However, MP coating is likely to be slowed down by the initial hydrophobicity of pristine MP. Dynamics of MP coating and increasing wettability are key factors for biotic and abiotic degradation processes.

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Environmental significance

The contamination of soil with microplastics (MP) is ubiquitous and occurs locally by deposition of larger plastic fragments with hotspots of high MP content leading to a loss of soil wettability. Consequently, locally reduced soil water content and air entrapment may occur, affecting soil functions such as capillary water flow as well as MP degradation. Here, we demonstrated that, depending on polymer type, coating of MP with iron (oxy)hydroxides reduces hydrophobicity of MP and allows for capillary infiltration of water. However, the initial low wettability of MP is likely to slow down the coating of MP. We conclude that the interaction of MP with surface-active environmental particles is important for the fate of MP and its impact on soil wettability.

1. Introduction

Terrestrial ecosystems are confronted with ever increasing amounts of microplastic (MP) particles of different origin, type, shape, size, and state of degradation.¹ The extent of MP contamination depends on anthropogenic activity levels.² In

floodplains in Swiss nature reserves, concentrations up to 593 particles per kg of MP are reported.³ Zhou *et al.* (2018) estimated 1.3–14 713 particles per kg in a coastal soil,⁴ Vollertsen & Hansen (2017) reported 53 000–528 000 particles per kg in agricultural soils⁵ and in industrial soils 0.3–67.5 g kg⁻¹ of MP were reported.⁶ However, also in remote locations like Antarctica, MP has been found in soils.⁷ The global plastic production of 2021 is estimated to 367 Mt.⁸ The total accumulated plastic waste generation is estimated to reach over 25 000 Mt by the year 2050 with maximum half of it being either correctly discarded, incinerated or recycled.⁹ Polyethylene terephthalate (PET), one of the main polymers used for packaging materials, constitutes a big part of plastic litter.^{1,8,10,11} Soil analysis for polymers mainly revealed groups of polyethylene (PE), polypropylene (PP), polyamide (PA) and polystyrene (PS).^{1,2}

Hitherto, most studies are focusing on the effect of MP on biological impacts while investigations on soil properties and functions are scarce. Since MP particles are typically hydrophobic, once they are deposited at the soil surface or incorporated into the bulk soil, they are likely to locally increase soil water repellency.¹² Previous experiments showed that pristine

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MP increases soil water repellency with the effect of inhibiting capillary rise.¹² Water bypasses areas of high MP contents, leading to locally reduced soil water contents by increasing tortuosity and entrapping air.¹² These observations imply that MP in soil is not easily wetted by soil water. Due to the importance of water on soil biotic and abiotic processes, this is expected to impact transport and fate of MP.

In the environment, MP particles may undergo various processes changing their surface properties *e.g.*, weathering by UV-light, hydrolysis, microbial activity and interactions with environmental particles and substances.^{13–17} Once incorporated into soil, MP is exposed to a variety of abundant soil and soil water constituents being able to alter MP surfaces¹⁸ and potentially increase their wettability. Such binding agents include inorganic substances such as minerals, metal hydroxides or organic matter.^{18–20} Recent studies demonstrated the formation of a surface coating and heteroaggregation of PS particles with ferrihydrite (a ferric iron (oxy)hydroxide) in the aqueous phase.²¹ Depending on the pH value, the surface properties of the MP were controlled by the ferric particles.²¹ Such iron compounds are not only highly abundant components in aquatic systems but also in soil and soil water (up to 20% (w/w)).²² MP can get in contact with ferrihydrite (Fh) suspension in unsaturated sand, a situation comparable to and commonly found at the capillary fringe, where anoxic Fe(II) bearing groundwater may come in contact with oxygen with subsequent formation of Fe(III) to Fh.²³ Thus, they are posing important coating agents of MP surfaces in terrestrial environments. First studies showed that MP particles and iron oxides interact in porous media and the relative smaller particles adsorbed onto the surface of the larger ones.²⁴ The resulting changes in surface properties influence the (co)transport of MP particles.²⁴ However, studies on the interactions between MP and soil components and the resulting changes of MP surface properties are scarce as most studies focus on the interactions of MP and environmental particles in aquatic systems.^{21,25,26}

Here, we tested if Fh coating influences the wettability of MP in soil. To this end, we investigated whether hotspots of pristine and coated MP (20–75 μm , PS and PET) embedded in a porous medium consisting of quartz sand will show differences in wettability during capillary rise (Fig. 1). Such regions of high MP content in soil are relevant since contamination usually occurs locally by deposition of larger plastic fragments, for example, by mulch film residues and their subsequent incorporation into the soil followed by fragmentation.^{27–29} The resulting wettability of MP hotspots was visualized and quantified by means of water saturation using neutron imaging and by determination of static contact angles (CA) by use of the sessile drop method (SDM). Neutron imaging is non-destructive to the soil structure and highly sensitive to hydrous material rendering this method optimal for imaging water in porous media.^{30–32} The common process of extracting and determining MP in soil, *e.g.*, density separation, is labor intensive and destructive to the soil structure. Neutron imaging allows to determine MP particle properties *in situ* without destroying the soil structure. Furthermore, hydrogen atoms are abundant in polymers allowing neutron imaging to distinguish PET and PS from dry soil material.^{12,33}



Fig. 1 Schematic setup of a sample container filled with sand and MP hotspot.

Differing material neutron attenuation coefficients can be used to quantify sample constituents in space and time.³²

We hypothesize that (i) the coating of MP particles with Fh increases their wettability; and (ii) the addition of Fh to water leads to a partial coating of pristine MP particles during capillary rise and hence increases the wettability of MP.

2. Experimental

2.1 Investigated media

We applied MP hotspots to porous media in varying combinations to estimate the effect of pristine MP and coated MP on soil wettability, respectively. As polymers we used PET and PS provided by Veolia Deutschland GmbH. It has been milled by a centrifugal mill (ZM200, Retsch GmbH) and sieved into a 20–75 μm fraction. Particle size analysis (Microtrac, Retsch GmbH) revealed irregularly shaped particles, ranging from near spherical to fibrous, resembling an environmentally relevant mixture of MP shapes. Size distributions and shapes are reported in the ESI (Fig. S1†).

As model soil particles, quartz sand (Raneem, Sand-Schulz GmbH, Berlin) in the size range of 700–1200 μm was used. To exclude effects of organic matter on wettability, the sand was treated with hydrogen peroxide (30%) at 95 $^{\circ}\text{C}$ for 8 h and subsequently washed with deionized water. Additionally, magnetic particles were removed to not obstruct the signals from neutron imaging.

2.2 Characterization of MP

2.2.1 Static contact angle. To evaluate static contact angles (CA) of MP, the sessile drop method (SDM) was applied.³⁴ MP, pristine and coated, was fixed to a microscopy glass slide by attaching a hydrophobic, double-sided adhesive tape.^{34,35} 1 μL droplets of deionized water ($n = 11$) were placed on the MP surface with a syringe. Simultaneously, the drop shapes were



recorded with a camera (device: Drop Shape Analyzer DSA30S, Krüss GmbH).

Each droplet geometry was analyzed with an elliptical shape recognition algorithm and measured at the three-phase contact line of liquid, solid and vapor 500 ms after the droplets detached from the syringe (software: Krüss Advance, Krüss GmbH). CA are the result of the combination of surface tensions of the aforementioned three phases according to eqn (1),^{36,37} where CA_e^Y is the equilibrium contact angle, γ is surface tension at the respective interfaces between solid (S), liquid (L) and vapor (V):

$$\cos CA_e^Y = \frac{\gamma_{SV} - \gamma_{SL}}{\gamma_{LV}} \quad (1)$$

Statistical analysis was conducted using the software MATLAB (R2021a). A parametric one-way ANOVA test was applied to the CA data of the MP to disclose a potential significant difference and the post hoc test of Bonferroni was employed to identify significant differences between the variants ($p < 0.001$). The letters A, B and C indicate significant differences.

2.2.2 Streaming potential. The streaming potential measurements have been performed with the electrokinetic analyzer SurPASS 3 (Anton Paar GmbH) and the cylindrical cell. The cell was filled to its maximum capacity with 128.4 ± 7.1 mg PS and 154.3 ± 6.3 mg PET, respectively. The particles were immobilized by polyamide-6,6 filter mats (Pieper Filter GmbH) with nominal pore diameters of 10 μm . The solutions were composed of analytic grade KCl (BioUltra, Merck), HCl, and KOH (Titrisol®, Merck). Titrations were performed from pH 4 to pH 10 at a constant overall ionic strength of 10 mM to neglect influences of surface conductivity. The reported zeta potentials have been calculated by means of the Helmholtz–Smoluchowski approximation.

2.2.3 Coating of MP with ferrihydrite. As shown in previous work, Fh colloids adsorb onto the surface of MP.²¹ The coating of MP was achieved by shaking 2 g MP per liter Fh suspension. Fh colloids were synthesized after Cornell & Schwertmann (2003).³⁸ For the coating process, the stock solution of Fh was diluted with deionized water to 100 mg L⁻¹. Additionally the ionic strength was set to 10 mM by addition of NaCl. After shaking the mixture for 2 h in an overhead shaker (Reax 20/4, Heidolph Instruments GmbH & Co. KG) at 16 rpm, the suspension was filtered through a 10 μm filter (nuclepore polycarbonate hydrophilic membranes, Whatman, Cytiva) to separate the coated MP particles from the remaining Fh colloids. The filter was then dried overnight at 40 °C.

Scanning Electron Microscopy (SEM) was conducted to confirm the coating of MP with Fh. The dried material was mounted onto a standard sample holder and coated with a thin layer of platinum (Cressington 208 HR sputter coater). SEM images were recorded with a Zeiss ULTRA PLUS (Carl Zeiss Microscopy GmbH, Germany). To calculate the amount of Fe adsorbed to the MP surface, the specific surface area of PS and PET was determined by the Brunauer–Emmett–Teller (BET) method. This method measures the surface area by a multi-

point surface N₂-adsorption isotherm at a temperature of 77 K (NOVA 2000e, Quantachrome Instruments).

Furthermore, the amount of iron adsorbed to the MP surfaces was determined. Approximately 0.03 g of coated MP were added to 5 mL 1 M HCl. The mixture was placed on an overhead shaker for 24 h to dissolve all ferric particles from the MP surface and afterwards filtered through 10 μm . The iron concentration of the filtrate was determined spectrophotometrically (DR 3800 VIS, Hach) at 512 nm using acetate buffer solution and 1,10-phenanthroline. Prior to the measurement, all ferric iron was reduced to ferrous iron by adding ascorbic acid to the sample. Triplicate samples were prepared.

2.3 Neutron imaging

Time-series neutron radiography of capillary rise into aluminum containers (inner dimensions: 6 × 16 × 60 mm) filled with sand and MP was conducted in wetting–drying cycles at the NEUTRA beamline at the Paul Scherrer Institute in Villigen, Switzerland.³⁹ The containers were pre-filled half with sand before MP was added *via* a funnel with subsequent addition of the other half sand (Fig. 1). The applied MP hotspots were pristine or coated with Fh. The mass of the MP hotspots was 0.02 g and range in size from height of *ca.* 1.5–2.5 mm and width of *ca.* 5.0–7.0 mm covering the whole depth of the sample container (6 mm). Capillary rise was imaged for a duration of 150 s to eventually estimate the water saturation of the capillary fringe. Differences in water saturation allow conclusions about the wettability of MP hotspots. An open container with in and out flow and a permeable bridge as stand for the sample containers was providing a constant water table for capillary rise. The respective liquid was delivered by a peristaltic pump at constant flow rate. Imbibition of different liquids was recorded for ten and six replicates of different MP polymer types (Table 1) with an acquisition time of 2 s and a pixel size of 53 μm . Three different liquids for capillary rise were used: (i) deionized water with 10 mM NaCl (water), (ii) 10 mg per L Fh suspension with 10 mM NaCl (Fh10) and (iii) 100 mg per L Fh suspension with 10 mM NaCl (Fh100). We conducted one wetting–drying cycle with water and two cycles with Fh10 and Fh100. Before the second wetting, the samples were dried at room temperature until constant weight.

Differing material neutron attenuation coefficients can be used to quantify the distribution of sample constituents.³² The attenuation coefficients for water and Fh suspensions were derived from step-wedge samples with defined thickness filled with the respective liquid. The coefficients were used to calculate water saturation based on porosity and dimensions of the

Table 1 Capillary rise experiment setups for neutron imaging

MP	Liquid	Repetitions
Pristine PET	Water, Fh10, Fh100	10
Coated PET	Water	6
Pristine PS	Water, Fh10, Fh100	10
Coated PS	Water	6



sample.⁴⁰ According to eqn (2), where, in each pixel of the images, Θ is the saturation of the respective liquid, $\text{liquid}_{\text{signal}}$ is the gray value signal after subtracting the dry soil signal, $\text{AC}_{\text{liquid}}$ is the respective attenuation coefficient of a liquid, s is the thickness of the sample material in direction of the neutron beam and n is the porosity of the sample material.⁴⁰

$$\Theta = \frac{\text{liquid}_{\text{signal}}}{\text{AC}_{\text{liquid}}} / (s \times n) \quad (2)$$

The gray value data of the images were normalized to the open beam signal, spot cleaned, and noise reduced.^{31,41} Subtracting gray values of the dry sample references from their wetted counterparts provides the signal of imbibed water.⁴⁰ The analyzed subsection of sample was set to 21 (width) \times 35 (height) pixel resulting in a region of interest (ROI) of merely the MP hotspot of 0.11 mm (width) and 0.2 mm (height). Statistical analysis was conducted using the software MATLAB (R2021a). The non-parametric Kruskal–Wallis test was applied to the water saturation data to disclose a potential significant difference and the post hoc test of Bonferroni was employed to identify significant differences between the treatments ($p < 0.001$). The letters A, B and C indicate significant differences where the combination of two letters shows similarity between those groups.

3. Results and discussion

3.1 Characterization of pristine and coated MP particles

Contact angles are a measure of polarity and hence allow conclusion about the wettability of particles.^{34,35} Typically, $CA < 90^\circ$ are regarded to allow for good wettability, while $>90^\circ$ can be considered hydrophobic.^{34,35} In our study, all particles exceed this threshold (Table 2). The analyzed MP surfaces partly indicate significant differences between pristine and coated MP as well as in polymer type (Fig. 2a, Tables 2 and S1†). The lower initial static CA of pristine PET (123.81°) compared to PS (143.71°) is significantly different and indicates a higher surface polarity of pristine PET⁴² and therefore, a higher wettability.

Coated PET has an average static CA of 101.67° and is significantly different from the other variants. Coating of PS (127.06°) resulted in static CA merely in the range of pristine PET (123.81°) indicating no significant difference between them and no wettability for both polymers. The coating of both polymer types with Fh significantly decreased the contact angle indicating a change in polarity. Nychka & Gentleman (2010)

mention that a moderate CA, besides other important surface properties, facilitates microbial colonization.⁴³ They differentiate between cell-substrate adhesion, where high wettability ($CA = 24^\circ$) is favorable and cell–cell cohesion where hydrophobicity ($CA = 84^\circ$) is of importance. Neither of those CA are observed in our experiment and thus, the used substrates were highly water repellent. Of all MP materials, coated PET showed the most favorable wettability for colonization exhibiting a CA of 101.67° .

In line with CA measurements, the coating of PET with Fh was more effective compared to PS. On the PET surface, $87.4 \mu\text{mol Fe}$ adsorbed per m^2 surface area of MP. Hence, surface coverage with Fh is more than four times higher compared to PS with $20.31 \mu\text{mol Fe}$ per m^2 surface area MP (Tables 2 and S2†). This is confirmed by SEM images showing a denser coating of the PET surface with small Fh particles compared to PS (Fig. S2†). The zeta potentials of PS and PET are negative in the analyzed pH range between 4 and 10 and become more negative with increasing pH value. The pH value of the Fh suspension used in the experiments was approximately 5.7. At this pH value, the zeta potential values of both polymer types were similar (Fig. S3† and Table 2) and thus cannot be the reason for the different adsorption behavior of Fh. Therefore, we assume that the presence of polar functional groups of the PET structure enhances the adsorption of Fh compared to PS. This is consistent with previous studies that investigated the adsorption of heavy metals or ciprofloxacin on MP.^{44–46} They observed that the adsorption capacity of MP particles increased with increasing UV-weathering. The UV-treatment increased the number of functional groups on the surface of MP particles and therefore provided more adsorption sites.¹⁸ For PET, the polymer backbone contains such functional groups even without weathering and therefore provides more adsorption sites for Fh compared to PS.

3.2 Neutron imaging

3.2.1 Capillary rise of water. In line with results from static contact angle measurements, the analysis of water saturation from neutron imaging shows similar qualitative patterns (Fig. 2b). After imbibition of water, mean water saturation values of pristine PET (0.10) exhibit non-significantly higher water saturation compared to pristine PS (0.04) (Table S3†). Both pristine polymers induce low water saturation and need to be considered non-wettable as water bypassed these hotspots (Fig. 3).

Table 2 Characterization of pristine and coated MP. The letters A, B and C indicate significant differences in contact angles of polymer types

MP type	Mean contact angle ($^\circ$), $n = 11$	Significance contact angle	$\mu\text{mol Fe}$ per m^2 MP, $n = 3$	Zeta potential (mV) (pH 5.7 ± 0.1), $n = 4$
Pristine PS	143.71 ± 3.29	A	—	-32.7 ± 0.5
Pristine PET	123.81 ± 3.87	B	—	-28.1 ± 0.1
Coated PS	127.06 ± 3.74	B	20.3	—
Coated PET	101.67 ± 4.37	C	87.4	—



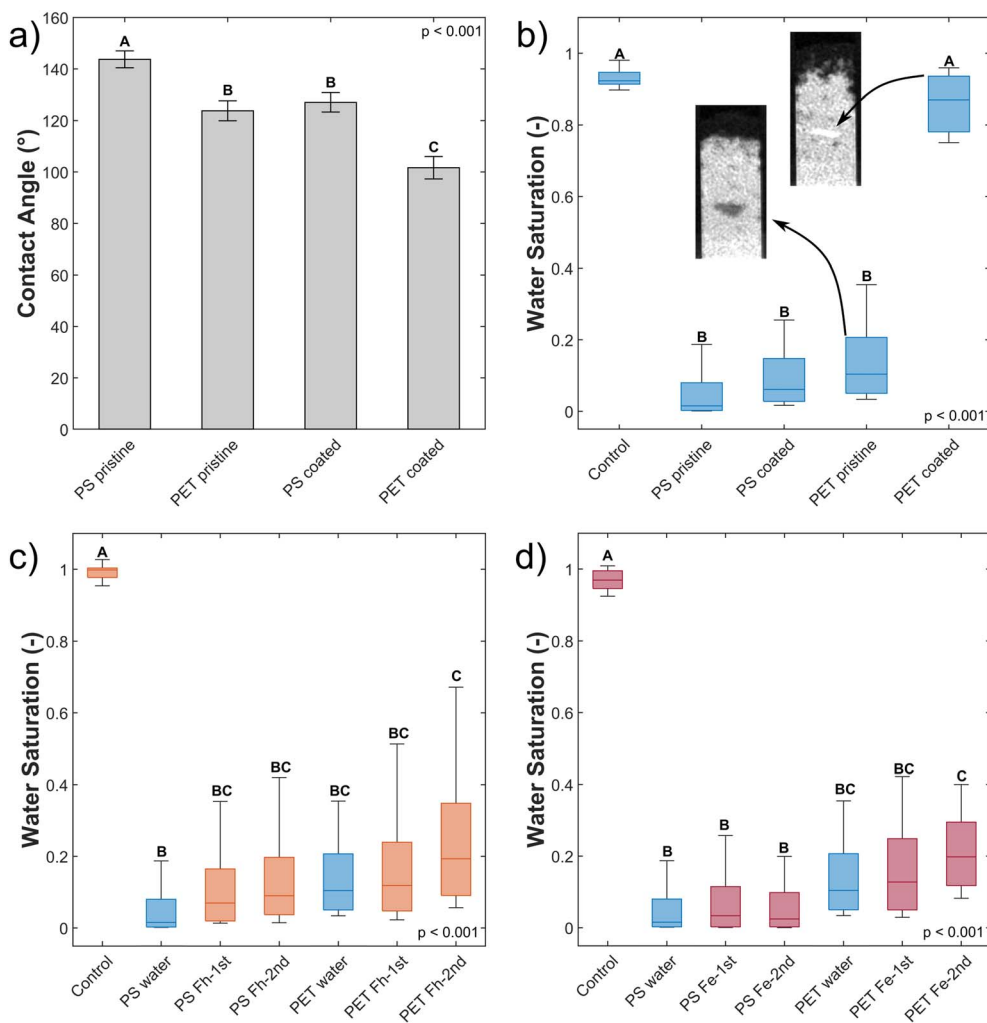


Fig. 2 (a) Contact angle of pristine and coated PS and PET. (b) Water saturation of MP hotspots after capillary rise with water. One wetting cycle for control samples without MP hotspot ($n = 10$), pristine ($n = 10$) and coated MP variants ($n = 6$) are shown. For PET, neutron images after capillary rise are shown. (c) Water saturation of MP hotspots after capillary rise with Fh10 (orange). One wetting cycle for control without MP ($n = 6$) and two wetting–drying cycles for pristine PS and PET variants ($n = 6$) are shown. For comparison, the results of the wetting of pristine PS and PET with water (blue) are shown. (d) Water saturation of MP hotspots after capillary rise with Fh100 (red). One wetting cycle for control samples ($n = 10$) and two wetting cycles for pristine PS and pristine PET variants are shown ($n = 10$). For comparison, the results of the wetting of pristine PS and PET with water (blue) are shown. The letters A, B and C in each graph indicate significant differences whereas the combination of two letters (e.g., AB) shows similarity between those groups.

Coated PET hotspots show a different behavior compared to pristine PET hotspots. The coated PET hotspots are wettable and water flow into the hotspot was observed (Fig. 3). Furthermore, water was even attracted by the hotspot, accelerating towards it and being further distributed from there (Fig. 3). We attribute this observation to a smaller pore geometry of the wettable, coated PET particles resulting in enhanced capillary forces compared to the sand. In the absence of coating, the smaller pore geometry of MP did not affect water flow due to high static CA. The observations are confirmed by water saturation values of the coated PET hotspots comparable to the control but with a broader distribution of values (Fig. 2b).

Other than in the case of PET, only a slight difference in water saturation between pristine and coated PS particles exists. Even though the water saturation of coated PS increased

slightly, the wettability of the hotspot did not change, and the hotspot is still considered non-wettable.

The increase in wettability of coated PET compared to coated PS can be explained by the effectiveness of the Fh coating. The amount of Fh adsorbed onto the PET surface was approximately four times higher compared to PS (Table 2). Therefore, we assume that the higher degree of coating of PET enables the formation of capillary water menisci in the pore space facilitating capillary rise. The wettability of hotspots of coated PET appear to contradict the results from static CA measurements with a CA of coated PET above 90° indicating non-polarity (see Section 3.1). Such discrepancy emerges from the methodological differences. Static CA measurements are derived from a quasi-two-dimensional system while capillary rise adheres to effective, dynamic CA of water in a three-dimensional pore



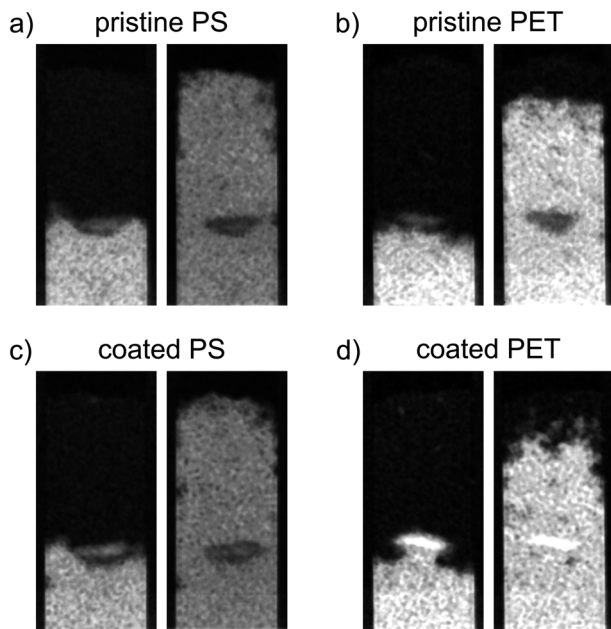


Fig. 3 Neutron images during and at the end of capillary rise of water. The hotspots consist of (a) pristine PS, (b) pristine PET, (c) Fh-coated PS, and (d) Fh-coated PET. For each set, the left image is taken at a time step at which the rising water reaches the hotspot. For coated PET, water flow into the hotspot is observed whereas for the other variants, the water bypasses the hotspot. The right image of each set is taken at the end of capillary rise (150 s).

space. Hence, the static CA derived allow to compare particles and treatments but cannot fully explain the dynamic process of capillary rise into a three-dimensional volume. In case of this study, the solid material consists of irregularly shaped sand grains and MP. The shape of the resultant liquid meniscus, and therefore the CA, is subject to forces based on the geometry of the pore space. In samples mixed with MP, dynamic CA are typically lower compared to static CA.¹² Instead of water not being able to spread beyond low wettability of MP during static CA measurements on a quasi-two-dimensional packing, water, during imbibition into a three-dimensional volume, has the possibility of bypassing into more favorable flow paths for imbibition and enclosing non-wettable locations.¹² Furthermore, forces like inertia of the imbibing liquid can also lead to overcome locations of low wettability. This would explain that the water saturation of coated PET is close to the one of the control but with a wider distribution of values (Fig. 2b).

3.2.2 Capillary rise of ferrihydrite suspensions. In the first part, we showed that a pre-established Fh coating of MP, depending on polymer type, can increase the wettability substantially. In a second experiment, we investigated to what extent coating of pristine MP hotspots can be established *in situ*. To this end, we used ferrihydrite suspensions instead of water and performed capillary rise experiments with a new set of pristine MP hotspots to mimic a situation commonly found at the capillary fringe.²³ To test if there is a dependency between Fh concentration in the soil suspension and increase in water saturation of the MP hotspot, we used two different Fh concentrations (10 and 100 mg L⁻¹).

Slight, but not significant, increases in water saturation relative to uncoated PS and PET were observed upon imbibition of Fh suspensions independent of concentration (Fig. 2c and d). A second wetting cycle did not lead to a change in wettability of the hotspots. Though, increased water saturation with a broader distribution of values show that Fh coating occurred locally during capillary rise. The explanation is that Fh suspensions have limited contact to the MP hotspots due to MP hydrophobicity. The suspensions bypassed the hotspots and only MP particles at the boundaries had contact with the imbibing Fh suspensions. It means that for coating and reducing the wettability of MP, Fh needs first to wet MP, which is a process that is hindered and slowed down by the initial hydrophobicity of MP. An increased number of wetting and drying cycles would likely allow the Fh suspensions to propagate deeper into the hotspots. The fact that larger increases of water saturation were only observed with PET is in line with differences in pre-coating of pristine MP, where PET coating was more profound than PS due to the presence of more functional groups in the PET polymer backbone.

4. Conclusion

We showed that naturally occurring coating agents like ferric (oxy)hydroxide increase the wettability of MP in porous media. This is of importance since MP has been shown to limit capillary flow in soils leading to air entrapments and reduced local water contents.¹²

Depending on polymer type, capillary driven imbibition of water into MP hotspots was facilitated by Fh pre-coating. While coated PS was still non-wettable, coated PET was wettable. We infer, a coating of particles, sufficiently high to render them wettable, depends on the presence of functional groups on the MP surface. Polymers with more functional groups in the polymer backbone like PET can be coated more efficiently compared to less functionalized polymers like PS.

In contrast to Fh pre-coating of MP, *in situ* coating of pristine MP hotspots with Fh suspension did not establish an effective coating regardless of Fh concentrations. The exposed MP hotspots were still non-wettable. Yet, a slight but not significant increase in water saturation and a wider distribution of values was visible. We summarize, that pristine MP, due to its hydrophobic properties, had limited contact to the imbibing Fh suspensions and only MP particles at the hotspot boundaries were exposed to Fh. We speculate, that adsorption of Fh onto MP in soils is a slow process, which requires multiple wetting-drying cycles in order to effectively alter the surfaces of MP in soil and render it wettable.

Translated into a natural process, our capillary rise experiment with wetting-drying cycles mimics fluctuating moisture conditions in the vadose zone. Over time, this process happens numerous times and therefore might allow MP in soil to get in contact with the soil suspension. However, the initial hydrophobicity of MP delays this process. The time frame necessary to overcome MP's inherent hydrophobicity is unclear and strongly depends on polymer type.



Understanding the coating and wetting kinetics of MP is a key factor to predict MP degradation, as the presence of water accelerates biotic and abiotic degradation processes.

Author contributions

The experimental work was performed by A. Cramer and J. Schmidtman. The manuscript was written by A. Cramer and J. Schmidtman with contributions of all authors. A. Cramer and J. Schmidtman contributed equally. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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