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Effect of rhamnolipid biosurfactant on transport and retention of iron oxide nanoparticles in watersaturated quartz sand†

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Although prior studies have investigated the effects of solution constituents, including dissolved organic matter and synthetic polymers, on nanoparticle mobility in porous media, far less attention has been directed toward evaluating the impacts of biosurfactants secreted by microorganisms on the transport and retention behavior of nanomaterials. The objective of this study was to explore the influence of rhamnolipid, a biosurfactant associated with biofilms, on the transport and retention of iron oxide nanoparticles (IONPs) in a water-saturated guartz sand. Column experiments were conducted using aerobic medium (ionic strength = 50.4 mM) or 10 mM NaCl as background electrolyte at a pore velocity of 0.43 m per day and pH 6.8 \pm 0.2. In aerobic medium columns, nearly all introduced nanoparticles were retained when IONPs were injected alone, whereas the presence of 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid resulted in ~25% and ~50% breakthrough of the injected IONP mass, respectively. Moreover, preflushing media with 50 mg L^{-1} rhamnolipid further increased IONP mass breakthrough by $\sim 30\%$. Similar enhancement of nanoparticle mobility by 50 mg L⁻¹ rhamnolipid was also measured in lower ionic strength (10 mM NaCl) columns. Mathematical models that incorporated nanoparticle filter ripening and biosurfactant competitive adsorption successfully reproduced experimental observations. Modeling results predicted an order-of-magnitude decrease in IONP filter ripening rate coefficient and a three-fold drop in average IONP retention capacity in the presence of rhamnolipid, consistent with a stabilizing effect and competition for surface sites. These findings demonstrate that rhamnolipid biosurfactant can potentially enhance nanomaterial stability and mobility in subsurface environments and that these effects should be considered when evaluating the impact of biological process on nanoparticle fate and transport in porous

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Environmental significance

The mobility and fate of engineered nanoparticles (ENPs) in the subsurface can be strongly influenced by biological process, including the presence of biosurfactants in the aqueous phase and adsorbed on solid surfaces. In this work, a combination of column transport experiments and mathematical modeling was undertaken to assess the impacts of a representative biosurfactant, rhamnolipid, on the transport behavior of oleic-acid coated iron oxide nanoparticles (IONPs) in water-saturated quartz sand. Co-injection of rhamnolipid increased IONP transport by up to 50%, while preflushing rhamnolipid further increased IONP transport by an additional 30%, demonstrating that both aqueous and adsorbed phase rhamnolipid contribute to enhanced IONP mobility. A multi-constituent mathematical model that accounts for competition at surface sites and filter ripening was able to accurately capture the measured IONP effluent breakthrough curves and retention profiles, and revealed that rhamnolipid can decrease the rate of filter ripening by up to an order of magnitude and decrease IONP attachment by 3-fold. These findings demonstrate that in contrast to biofilms, biologically generated surfactants have the potential to dramatically enhance ENP mobility in water-saturated porous media and should be accounted for when describing ENP transport behavior in biologically active systems.

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1. Introduction

In response to public and scientific community concerns related to engineered nanoparticle (ENP) toxicity and fate in the environment, 1,2 substantial research efforts have been made to improve our understanding of the transport, transformation and retention of ENPs in porous media.3-5 Due to their ability to effectively retain particles, surface soils and other natural porous media systems (e.g., aquifers) will likely serve as important sinks, and may ultimately entrap most ENPs entering the environment⁶ from unintentional releases (e.g., landfill leachate) and intentional applications (e.g., subsurface characterization and enhanced oil recovery). The transport behaviour of natural and synthetic micro-size particles, such as bacteria and viruses, 7-12 clay minerals, 13,14 latex particles, 15-17 and silica microspheres, 18,19 and various nanoparticles^{1,20,21} in porous media has been extensively studied for more than 30 years. 22,23 Previous experimental studies of particle transport and retention in packed columns have demonstrated that particle attachment can vary spatially and temporally, leading to non-exponential particle retention distributions.²⁴⁻²⁶ For example, Wang et al. reported nearly constant retention of nanoscale fullerene aggregates (nC_{60} , mean diameter = 95 nm) along the length of water-saturated columns packed with 40-50 mesh Ottawa sand ($d_{50} = 0.36$ mm), indicating a limiting or maximum retention capacity.²⁷ Liang et al. co-injected 10 mg L⁻¹ stabilized silver nanoparticle with a mixture of 10 mg L⁻¹ and polyoxyethylene glycerol trioleate polyoxyethylene (20) sorbitan monolaurate (Tween 20), respectively, and observed retention that was lowest near the column inlet and increased to a maximum at the column midpoint.²⁸ Similarly, several studies by Johnson et al. have demonstrated non-exponential retention, including hyperexponential and nonmonotonic profiles, of microspheres and colloids (from 0.1 to 6.8 µm) under unfavourable (energy barrier) deposition conditions.^{26,29–33}

These studies demonstrated that transport and retention of colloids and ENP in porous media is complex and can involve a number of processes (aggregation, attachment, detachment, physical straining, filter ripening and particle near-surface retention/mobilization^{32,34,35}) that are strongly influenced by interactions between particles and the porous media. Such interactions are not only influenced by the physicochemical conditions³⁶ of the transport system, such as ionic strength (IS) and pH, but can also be impacted by nanoscale surface heterogeneity^{33,35} and co-constituents present in the aqueous phase and/or adsorbed on solid phases. In regard to co-constituents, the effects of natural $(NOM)^{37-40}$ organic matter and polymer/surfactant stabilizers^{41–46} on particle transport have received considerable attention. The majority of these studies observed enhancements in nanoparticle mobility in the presence of NOM/polymers under environmentally relevant conditions. Increased effective transport, corresponding reduction in attachment, has been attributed

to greater electrostatic repulsion and/or steric hindrance between nanoparticles and the solid phase due to surface adsorption or coating by NOM/polymers. 36,47-49 Additionally, several studies have demonstrated that competition between ENP and NOM/polymers for attachment sites can also limit nanoparticle deposition. 28,50,51 For example, Wang et al. 44 reported reduced retention of quantum dots (QDs) near the column inlet and an increasing solid-phase OD concentration with distance, observations that contradict classical colloid filtration theory. This behaviour was attributed to the presence of residual polymer coatings in the QD suspension, which preferentially adsorbed and competed for attachment

Similarly, extracellular biological constituents, such as biofilms and biosurfactants, have the potential to impact interactions between nanoparticles and porous media surfaces.⁵² A number of studies have demonstrated that biofilms enhance nanoparticle retention^{53–55} in porous media through reduced repulsive force(s), hydrogen bonding, hydrophobic or hydrophilic bridging interactions, 56,57 and strengthened filtration and physical straining processes due to biofilm induced roughness⁵⁸ and smaller pore size.⁵⁹⁻⁶¹ In contrast, far less attention has been directed to the potential impact of biosurfactants such as rhamnolipids, which are produced by some species of Pseudomonas (e.g., Pseudomonas aeruginosa), on nanoparticle mobility. Rhamnolipids are anionic surfactants that consist of rhamnose linked to one or two molecules of β-hydroxydecanoic acid. 62 Due to their surface activity, rhamnolipids have been considered as potential replacements for synthetic surfactants and have been shown to exhibit similar levels of detergency compared to sodium dodecyl sulfate (SDS), a widely used anionic surfactant. 63 Basnet and coworkers investigated the effects of rhamnolipid biosurfactant on the aggregation and transport behavior of palladium-doped zerovalent iron nanoparticles (Pd-NVZI) in a water-saturated quartz sand⁶⁴ and a loamy sand.65 Rhamnolipid-coated Pd-NVZI exhibited good colloidal stability and enhanced transport in both porous media even at the highest IS considered (300 mM NaHCO₃), suggesting that rhamnolipid acted to stabilize the Pd-NZVI suspension. They also found that deposition rate of rhamnolipid coated Pd-NZVI increased with time (exhibiting ripening-type behavior), distinctive from the decreasing or constant deposition rate observed for carboxymethyl cellulose coated and soybean flour coated nanoparticles. 65 However, in these studies, rhamnolipid was only used as a nanoparticle surface modifier and the potential effects of rhamnolipid in the aqueous and solid phases on nanoparticle mobility were not considered.

Mathematical models are valuable tools for analyzing nanoparticle transport behavior in porous media and for identifying associated attachment parameters. Several studies have demonstrated that simultaneous model fitting to both nanoparticle effluent concentration data and retention profiles can provide important insight into retention processes. For example, Li et al.24 employed a mathematical

model, based upon a modification of classical colloid filtration theory, to fit experimental observations of fullerene (C₆₀) breakthrough and retention in packed columns. This simultaneous fitting process revealed that nanoparticle attachment on mineral surfaces exhibited a maximum capacity (S_{max}) that was far smaller than monolayer coverage, and that the attachment rate coefficient and S_{max} were dependent upon flow velocity and grain size. This work was extended by Becker et al.25 who developed a multiconstituent site blocking (MCB) model based upon column observations of nanoparticle transport in the presence of free polymer. In that study, residual polymer present in quantum dot suspensions was shown to compete with nanoparticles for attachment sites, resulting in nanoparticle retention that increased with distance from the column inlet, counter to predictions from classical colloid filtration theory. In a subsequent study, the developed modeling framework was implemented in a multi-dimensional simulator to successfully predict the transport of magnetite nanoparticles in a heterogeneous flow cell.⁶⁶

Building on this prior work, the goal of this research was to investigate the effects of aqueous phase and adsorbed rhamnolipid biosurfactant on the transport and retention behaviour of ENPs in a water-saturated porous medium through a combination of experimental and mathematical modelling studies. We hypothesize that rhamnolipid adsorbed on the solid phase will inhibit nanoparticle attachment, while rhamnolipid present in the aqueous phase will act to stabilize nanoparticles as well as compete for attachment sites, and that the combined effects of these processes will result in enhanced nanoparticle mobility compared to rhamnolipid-free control experiments. Oleic acid coated iron oxide nanoparticles (IONPs) were chosen as model ENPs due to their stability, low toxicity, broad range of applications, and reproducibility. 67-71 Batch studies were conducted to determine the effect of rhamnolipid, as a function of concentration, on the aggregation kinetics and zeta potential of IONP. A series of column experiments was then performed to investigate the impacts of rhamnolipid on IONP mobility and deposition. Aqueous and solid phase concentrations of IONP were measured to construct IONP breakthrough curves (BTCs), retention profiles, and to calculate mass balance. Based on these experimental data, a two-site, multi-constituent transport model, was developed that incorporates both filter ripening (particle-particle interactions) and competitive adsorption effects on attachment. This model, an extension of the MCB model, 25 was then used to estimate nanoparticle attachment rate and capacity parameters in the presence and absence of rhamnolipid.

2. Materials and methods

2.1 Materials

Iron oxide nanoparticles (magnetite, 8 nm core diameter) were synthesized using the method described by Li et al. 70 Briefly, 0.178 g FeO(OH) fine powder, 2.26 g oleic acid and 5.0 g 1-octadecene were mixed in a three-neck flask equipped with a heating mantle and a temperature controller. The system was maintained at 120 °C for 1 h to remove residual water and then heated to 320 °C for 1 h under constant argon environment. The resulting brown-black colloid was purified using acetone and hexane with 4-5 washes by centrifugation (Sorvall WX Ultra 80, Thermo Scientific, MA) at 8000 rpm for 10 min to remove unreacted iron salts. After the final wash, nanoparticles in the hexane phase were transferred to water by adding a second layer of oleic acid (as a stabilizer) using a probe sonicator.⁶⁷ After sonication, the suspension was stirred for 24 h to evaporate residual hexane. The aqueous phase was collected and washed with Milli-Q water (18.2 M Ω cm) 5-6 times to separate out the nanoparticles. Under the conditions and time frames considered in this study, the oleic acid coating is stable and not considered to be a sensitive variable.72

Rhamnolipid biosurfactant (90% purity) was purchased from AGAE Technologies (Corvallis, OR). According to the manufacturer, the ratio of mono-rhamnolipid to dirhamnolipid is approximately 3:2, which yields an average molecular weight of 563 g mol⁻¹ with a reported critical micelle concentration (CMC) of approximately 100 mg $L^{-1.73}$ Aerobic medium and 10 mM NaCl solution were prepared to represent biotic and environmental-relevant background electrolytes, respectively. Aerobic medium was prepared with $1.50 \ \ g \ \ L^{-1} \ \ KH_2PO_4, \ \ 1.00 \ \ g \ \ L^{-1} \ \ (NH_4)_2SO_4, \ \ 0.10 \ \ g \ \ L^{-1}$ $MgSO_4\cdot7H_2O$, 0.05 g L⁻¹ CaCl₂·2H₂O and 5.00 g L⁻¹ glycerol (IS = 50.4 mM).⁷⁴ The aerobic medium was sterilized using an autoclave (Amsco Century Medium Steam Sterilizer; Steris, OH) with a liquid cycle maintaining at 120 °C for at least 60 minutes. Prior to use, a sterilized NaOH solution was added to adjust the pH of aerobic medium and 10 mM NaCl solution to 6.8 \pm 0.2. All chemicals used as background electrolytes were purchased from Fisher Scientific (Fair Lawn, NJ). Ottawa sand (80–100 mesh) with a mean diameter (d_{50}) of 165 µm was obtained from US Silica (Ottawa, IL) as F65 and separated using a programmable sieve shaker (Tyler Ro-Tap, W.S. Tyler Industrial Group, OH). Prior to use, the 80-100 mesh size fraction was cleaned using a sequential acid wash, water rinse, ultrasonication, and oven-drying procedure.²⁷

2.2 Column experiments

To minimize potential microbial contamination, column equipment and materials were sterilized using either an autoclave or disinfectant (70% ethanol) prior to use. A total of 15 columns were packed with quartz sand and completely saturated with water following procedures described previously.44 Briefly, borosilicate glass columns (5 cm diameter × 15 cm length) were packed with washed 80-100 mesh Ottawa sand in 1 cm increments under gentle vibration. The column endplates were fitted with two 40 mesh nylon screens and a 40 µm nylon filter to support the

solid phase and promote uniform flow. The packed columns were then flushed with CO2 gas for at least 60 min to facilitate subsequent dissolution of entrapped gas during the water imbibition process. To fully saturate the columns, at least 10 pore volumes (PVs) of background solution were introduced into the column in an up-flow mode using a highliquid chromatography performance (HPLC) pump (ChromTech, MN). After water saturation, a non-reactive tracer test was conducted using sterilized NaBr solution (50 mM or 10 mM) to characterize water flow and hydrodynamic dispersion. The effluent bromide ion concentrations were measured using ion-selective electrode(Cole-Parmer, IL) and fit to a one-dimensional form of the advective-dispersivereactive (ADR) transport equation using CXTFIT ver. 2.1 (ref. 75) to obtain the hydrodynamic dispersion coefficient $(D_{\rm H})$ and the retardation factor $(R_{\rm F})$.

Five sets of column experiments were performed to evaluate the effects of rhamnolipid on nanoparticle transport and retention in water-saturated 80-100 mesh Ottawa sand. The initial set of column experiments was conducted to obtain baseline IONP transport and retention behaviour in water-saturated porous medium in the absence of rhamnolipid. Here, 3 PVs of approximately 5 mg L⁻¹ IONP (as iron) either in aerobic medium or 10 mM NaCl were injected to the saturated columns, followed by 3 PVs of IONP-free background electrolytes to displace unattached nanoparticles from the column. To characterize the transport of rhamnolipid alone, a second set of column experiments was conducted by injecting solutions containing only rhamnolipid. Here, 5 PVs of 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid in aerobic medium or 10 mM NaCl were introduced to water-saturated columns, followed by 3 PVs rhamnolipid-free background solution. The third set of column experiments was performed to explore the influence of rhamnolipid pre-treatment on IONP transport. Here, the water-saturated column was first pre-flushed with approximately 5 PVs 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid solution, followed by 3 PVs of background electrolyte solution to remove any free rhamnolipid from the column. A 3 PV pulse of approximately 5 mg L⁻¹ IONP solution (as iron) was then introduced into the column, followed by 3 PVs IONPfree background solution. In the fourth set of column experiments, IONP transport and retention behaviours were evaluated in the presence of free rhamnolipid (i.e., coinjection). Here, 3 PVs aerobic medium solution containing 5 mg L⁻¹ IONP (as iron) and either 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid or 3PVs 10 mM NaCl solution containing 5 mg L⁻¹ IONP (as iron) and 50 mg L⁻¹ rhamnolipid were introduced to the water saturated columns, followed by 3PVs background electrolyte solution (aerobic medium or 10 mM NaCl solution). In the fifth and final set of column experiments, IONP transport was evaluated under the more realistic condition where rhamnolipid was present in both solid and aqueous phases (i.e., pre-treatment + co-injection). Here, the water-saturated columns were first flushed with 5 PVs of 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid solution and 3 PVs rhamnolipid-free background solution (aerobic medium or 10 mM NaCl), followed by the injection of 3 PVs of suspension containing 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid and approximately 5 mg L⁻¹ IONP (as iron), and finally, 3 PVs background electrolyte solution.

All the column experiments described above were conducted at a constant flow rate of 0.06 ± 0.001 mL min⁻¹ corresponding to a pore-water velocity of approximately 0.43 m per day. All influent solutions were prepared by coinjecting the stock nanoparticle solution (~150 mg L⁻¹) and background solution (with or without rhamnolipid) at a flow rate of 0.002 mL min⁻¹ and 0.058 mL min⁻¹, respectively, and the two solutions were mixed immediately before introduction to the column (Fig. S2†). The nanoparticle size was monitored at the start and end of the injection and no significant changes were observed. Effluent samples from the columns were collected continuously using a fraction collector (CF-2, Spectrum Laboratories, CA). To construct IONP BTCs, effluent concentrations expressed as the relative concentration (C/C_0) , where C_0 is the applied influent concentration, were plotted versus the number of dimensionless PVs of solution delivered. To obtain IONP retention profiles at the conclusion of each experiment, the columns were sectioned into ten increments and the solidphase concentrations were measured and plotted versus the distance from the column inlet. Duplicate experiments were conducted for 50% of the columns to confirm reproducibility. Experimental conditions for IONP column studies are summarized in Table 1, while the rhamnolipid only column results are provided in Table S2.†

2.3 Analytical method

The hydrodynamic diameter and electrophoretic mobility of IONP in suspension were characterized with a Malvern ZetaSizer Nano ZS (Malvern, Cambridge, United Kingdom) using dynamic light scattering (DLS) and laser Doppler velocimetry, respectively. These measurements were conducted at room temperature (i.e., approximately 22 °C) and a scattering angle of 90 degrees. The Smoluchowski approximation was used to relate electrophoretic mobility to zeta potential. An ultra-high-resolution field emitter scanning electron microscope (SEM) (Zeiss LEO 1530 VP, MN) equipped with an energy dispersive spectrometer (EDS) was used to image sand grains and to characterize surface chemistry. Prior to SEM imaging and EDS analysis, all samples were secured on carbon conductive tape and coated with a layer of gold and palladium. IONP core size and morphology were characterized by transmission electron microscopy (TEM, FEI Tecnai G2 Spirit, OR) operated at 120 kV (Fig. S1a†). TEM sample was prepared by placing a small drop (10 µL) of diluted IONP suspension on a carbon coated copper grid (Electron Microscopy Sciences, PA) and left to dry at room temperature (22 \pm 0.5 °C). The IONP core size distribution was obtained by counting more than 500 randomly chosen nanoparticles from the TEM images using

Table 1 Summary of IONP column experimental parameters and mathematical modelling results

	Experimental parameters				Mathematically fitted parameters					
Column identifier ^a	$ \begin{array}{c} C_0^b \\ (\text{mg L}^{-1}) \end{array} $	PW ^c (PV)	BT^d (%)	MB ^e (%)	$k_{\text{att}}^f(\mathbf{h}^{-1})$	$S_{1, \max}^{p} (\mu g g^{-1})$	$k_{\mathrm{rip}}^{}h}\left(\mathrm{h}^{-1} ight)$	$k_{ m ads}^{i}({ m h}^{\scriptscriptstyle -1})$	$lpha^j$	R^2
AE-IONP-1	5.0	3.0	3.2	94.8	0.52 (0.39, 0.65)	5.60 (1.38, 9.82)	0.64 (0.23, 1.05)	NA	0.051	0.99
AE-IONP-2	5.6	3.0	1.1	96.1	0.49 (0.37, 0.61)	6.11 (1.21, 11.01)	0.57 (0.20, 0.94)	NA	0.046	0.99
AE-R10_IONP	4.9	3.1	1.0	93.9	0.55 (0.43, 0.67)	4.03 (1.44, 6.62)	0.49 (0.22, 0.76)	NA	0.046	0.99
AE-R50_IONP	4.6	3.2	12.9	105.1	0.19(0.16, 0.22)	2.24 (0.81, 3.67)	0.21 (0.10, 0.31)	NA	0.017	0.97
AE-R10&IONP	5.1	3.1	24.9	107.4	0.32 (0.25, 0.39)	6.85(5.72, 7.98)	0.06 (0.05, 0.07)	2.73 (1.75, 3.71)	0.017	0.96
AE-R50&IONP-1	5.0	3.1	56.6	106.6	0.43 (0.05, 0.81)	5.98(0,41.78)	NA	0.08(0, 0.87)	0.021	0.92
AE-R50&IONP-2	4.8	3.0	43.2	99.7	0.38 (0.26, 0.50)	7.67(0,42.54)	NA	0.09(0, 0.74)	0.027	0.87
AE-R10_R10&IONP	4.9	3.1	28.4	97.9	0.25 (0.23, 0.27)	6.74 (5.89, 7.59)	0.07 (0.06, 0.08)	1.70(1.19, 2.21)	0.014	0.92
AE-R50_R50&IONP-1	5.4	3.1	78.7	99.0	0.09(0.06, 0.12)	2.64(0,53.55)	NA	0.03(0, 0.96)	0.007	0.95
AE-R50_R50&IONP-2	4.8	3.0	83.0	97.6	0.16 (0.07, 0.25)	1.94 (0, 34.86)	NA	0.05(0, 1.34)	0.010	0.95
NaCl-IONP-1	5.3	3.0	14.9	104.5	$0.70 \ (0.55, 0.85)$	3.75 (3.57, 3.93)	NA	NA	0.030	0.97
NaCl-IONP-2	5.1	3.0	12.3	102.9	0.55 (0.44, 0.66)	4.04 (3.74, 4.34)	NA	NA	0.024	0.96
NaCl-R50&IONP-1	5.8	3.0	67.3	92.2	0.84 (0.60, 1.08)	8.56 (7.95, 9.17)	NA	2.01 (0.68, 3.34)	0.037	0.98
NaCl-R50&IONP-2	5.2	3.0	66.9	99.7	0.58 (0.46, 0.70)	9.08 (8.50, 9.68)	NA	1.68(0.77, 2.59)	0.025	0.99
NaCl-R50_R50&IONP	5.8	3.0	95.3	100.4	0.18 (0.09, 0.27)	2.37 (1.36, 3.36)	NA	183.8 (161.5, 206.1)	0.008	0.99

^a "AE": aerobic medium as background electrolyte, "NaCl": 10 mM NaCl as background electrolyte, "R10": 10 mg L⁻¹ rhamnolipid solution, "R50": 50 mg L⁻¹ rhamnolipid solution, "_": preflood, "&": co-injection, e.g. "AE-R10&IONP": IONP and 10 mg L⁻¹ rhamnolipid were co-injected into the column using aerobic medium as background electrolyte; "AE-R50_IONP": column preflushed with 50 mg L-1 rhamnolipid solution and then IONPs were injected using aerobic medium as background electrolyte. "NaCl-R50_R50&IONP": column preflushed with 50 mg L⁻¹ rhamnolipid and then IONP with 50 mg L⁻¹ rhamnolipid were injected using 10 mM NaCl as background electrolyte. "-1" and "-2" indicate duplicate experiments. b Input IONP concentration (as iron). Pulse width. Percent of IONP breakthrough mass to total applied mass. Mass balance. f IONP attachment rate coefficient. Maximum retention capacity of porous media surface (type 1 site capacity for two-site model). ^h IONP filter ripening attachment rate coefficient. ⁱ Rhamnolipid adsorption rate coefficient. The 95% confidence limits of fitted parameters are also presented after fitted parameters in parenthesis. The relatively large 95% confidence intervals (CI) for S_{\max}^p and k_{ads} in the columns "AE-R50&IONP" and "AE-R50_R50&IONP" indicate that the model predictions are not sensitive to maximum retention capacity and first order adsorption rate under the experimental conditions. Additionally, zero-truncation for the lower bound of CI was used, given that S_{\max}^p and k_{ads} are positive values. Analysis of the CI obtained for these column experiments are documented in the ESI. † Collision efficiency factor. Collision efficiency factors were estimated by classical filtration theory 97 and empirical correlation equations developed by Tufenkji and Elimelech. 98 The sum of attachment rate and ripening attachment rate coefficient was used in the calculation.

ImageJ software (National Institutes of Health, MD), as shown in Fig. S1b.†

Quantification of rhamnolipid concentrations in aerobic medium was performed with a Jasco V-730 UV-vis spectrophotometer (Easton, MD) using a modified anionic surfactant-methylene blue method.^{76,77} containing 10 mM NaCl as the background electrolyte, rhamnolipid concentrations were determined using a total organic carbon analyzer model TOC-L series with Autosampler ASI-L (Shimadzu Scientific Instruments, Columbia, MD). For both methods, seven-point calibration curves were constructed using rhamnolipid solutions (0-100 mg L⁻¹), which yielded linear fits with regression coefficients (R^2) greater than 0.98.

Iron concentrations in aqueous and solid samples were determined using a Jasco V-730 UV-vis spectrophotometer based on a modified ferrozine method.78 The method detection limit for total iron was 10.7 µg L⁻¹, obtained using the method of Armbruster and Pry. 79 Prior to UV-vis measurement, the IONPs in aqueous and solid samples were digested by 10% hydrochloric acid (HCl) under sonication (ultrasonic batch, Fisher Scientific, MA) for 2 h at room temperature. The average background iron concentration of the cleaned 80–100 mesh Ottawa sand was $4.7 \pm 0.3 \mu g$ Fe per g sand for this digestion method. Accuracy of the UV-vis

method was confirmed using an Optima 7300 DV inductively coupled plasma optical emission spectrometer (ICP-OES) (PerkinElmer, Waltham, MA).

2.4 Mathematical modelling

A multi-constituent nanoparticle transport model developed in prior work²⁵ serves as the foundation for the mathematical model employed herein. Consistent with the experimental observations described below, two types of sites were modelled. The first type of site (type 1 site) accounts for nanoparticle attachment and biosurfactant adsorption on porous media surfaces, assuming a Langmuir-type limiting or maximum surface capacity. The second type of site (type 2 site) accounts for filter ripening, where previously deposited nanoparticles provide additional sites for nanoparticle deposition. A coupled system of one-dimensional aqueous phase transport equations is used to describe constituent transport and retention in a homogeneously packed column:

$$\frac{\partial C^{i}}{\partial t} + \frac{\rho_{b}}{\theta_{w}} \frac{\partial S^{i}}{\partial t} = D_{h} \frac{\partial^{2} C^{i}}{\partial x^{2}} - \nu_{p} \frac{\partial C^{i}}{\partial x}$$
 (1)

where, C^{i} is the concentration of constituent i, nanoparticle (i = p) or rhamnolipid (i = R), in the aqueous suspension (M L^{-3}); S^{i} is the solid-phase concentration of constituent i (M

 M^{-1}); t is time (t), x is distance from the column inlet (L); $\nu_{\rm p}$ is the average pore-water velocity (L t^{-1}), D_h is the hydrodynamic dispersion coefficient for the column (L² t⁻¹), $\rho_{\rm b}$ is the bulk density of the porous medium (M L⁻³); and $\theta_{\rm w}$ is the volumetric water content. For the nanoparticle transport equation, the solid phase accumulation term is expanded, consistent with a two-site deposition model, as:

$$\frac{\rho_{b}}{\theta_{w}} \frac{\partial S^{p}}{\partial t} = \frac{\rho_{b}}{\theta_{w}} \frac{\partial S_{1}^{p}}{\partial t} + \frac{\rho_{b}}{\theta_{w}} \frac{\partial S_{2}^{p}}{\partial t}$$
(2)

The first term in (2) accounts for competitive deposition/ sorption between nanoparticles and rhamnolipid at type 1 sites (i.e., sand surfaces):

$$\frac{\rho_{\mathbf{b}}}{\theta_{\mathbf{w}}} \frac{\partial S_{1}^{i}}{\partial t} = k_{\mathbf{a}}^{i} \Psi C^{i} - k_{\mathbf{d}}^{i} \frac{\rho_{\mathbf{b}}}{\theta_{\mathbf{w}}} S_{1}^{i} \tag{3}$$

while the second term accounts for nanoparticle attachment due to filter ripening at type 2 sites:

$$\frac{\rho_{\rm b}}{\theta_{\rm w}} \frac{\partial S_2^p}{\partial t} = k_{\rm rip} \Phi C^p \tag{4}$$

Here, S_1^p and S_2^p represent nanoparticle retention concentrations (M M⁻¹) at type 1 and 2 sites, respectively (note $S^p = S_1^p + S_2^p$). S_1^i is the solid-phase concentration of constituent i (M M⁻¹) at type 1 sites (i.e., sand surface), k_a^i is the first-order attachment or adsorption rate coefficient for constituent i due to physical attachment or preferential sorption (t^{-1}) , k_d^i is a first-order detachment or desorption rate coefficient (t^{-1}), and k_{rip} is a first-order ripening rate coefficient describing the nanoparticle filter ripening process (t⁻¹). Based on the absence of tailing in IONP effluent BTCs, the IONP detachment rate coefficient was assumed negligible. Deposition of nanoparticles and rhamnolipid at type 1 sites are coupled through a surface site depletion function Ψ (M M⁻¹), which decreases from 1 to 0 as nanoparticles and/or rhamnolipid deposit on the surface of the porous medium (type 1 sites):

$$\Psi = 1 - \frac{S_1^p}{S_{1,\text{max}}^p} - \frac{S_1^R}{S_{1,\text{max}}^R} \tag{5}$$

Here, S_1^p and S_1^R represent nanoparticle and rhamnolipid retention concentrations (M M⁻¹) at sand surfaces (type 1 sites), $S_{1,\text{max}}^p$ and $S_{1,\text{max}}^R$ are the maximum retention capacities of nanoparticle and rhamnolipid at type 1 sites (M M⁻¹). The individual $S_{1,\text{max}}^p$ and $S_{1,\text{max}}^R$, for nanoparticle and rhamnolipid, are scaled by molecular size, which influences their effective surface coverage on the porous medium (detailed calculations provided in the ESI†).

The deposition of nanoparticles on type 2 sites is related to type 1 site attachment through use of a ripening Φ (M M⁻¹), which increases evaluation function proportionally with the total deposition nanoparticles(S^p):^{80,81}

$$\Phi = \frac{S^p}{S_{1,\text{max}}^p} \tag{6}$$

Here, S^p is the total nanoparticle retention concentration (M M^{-1}) at both two types of sites, i.e. $S^p = S_1^p + S_2^p$. As more nanoparticles occupy the surface sites (type 1) and more nanoparticles are deposited on previously attached nanoparticles (type 2 sites), the number of type 2 sites grows, creating an increase in the effective ripening rate $(k_{rip}\Phi)$. S_1^p max is used here to normalize the influence of total nanoparticle deposition S^p in the ripening attachment term and to maintain a similar functional form for the k_{rip} and k_a^i

In addition to the two-site multi-constituent model described above, a two-site single constituent model (TSC), which only accounts for filter ripening (no competition for type 1 sites), was used to model experimental results of IONP transport in columns where there was no secondary constituent. For NaCl columns, a modified filtration theory model (MFT)24 was employed in conjunction with a one-site multi-constituent model (MCB)²⁵ to simulate experimental data when there was no indication of filter ripening. Governing equations and comparisons of these models to other commonly used mathematical models (e.g. clean-bed filtration model) are summarized in the ESI.†

The mathematical expressions were implemented using a central-in-space and fully implicit-in-time finite difference scheme in MATLAB R2020a (The MathWorks, Natick, MA), and the transport and solid-phase interaction equations for each constituent were solved sequentially. Model parameters, including attachment/adsorption rate coefficients, filter ripening rate coefficient and maximum retention capacity of the porous medium surface, were estimated by fitting the to experimental breakthrough curves and corresponding retention profiles using a least-squares optimization procedure based on the following objective function:

$$F(x) = \beta \cdot \frac{\sum_{t} (\text{ExpBTC} - \text{ModelBTC})^{2}}{\text{TotalMassBTC}} + \gamma \cdot \frac{\sum_{t} (\text{ExpRET} - \text{ModelRET})^{2}}{\text{TotalMassRET}}$$
(7)

where β and γ are weighting factors, which allow the user to fit breakthrough (ExpBTC) and retention (ExpRET) simultaneously with different degree of importance. In this study, weighting factors were optimized for each column to provide best fits to experimental results. The corresponding weighting factors for each column experimental data are listed in Table S7.†

3. Results and discussion

3.1 IONP transport in aerobic medium

To simulate conditions conducive to microbial growth that would likely yield relatively large quantities of rhamnolipid, ten column experiments were conducted with aerobic growth medium (IS = 50.4 mM, pH = 6.8 \pm 0.2), as described above, as the background electrolyte. The transport and retention of oleic acid coated IONPs was evaluated in the absence of rhamnolipid, with rhamnolipid pre-treatment, co-injection,

and pre-treatment plus co-injection at two rhamnolipid concentrations (10 and 50 mg L⁻¹). The experimental conditions for these IONP column experiments are summarized in Table 1. The nanoparticle transport models were fit to the measured effluent breakthrough curves and retention profiles to obtain the following parameters: IONP attachment rate coefficient (katt), surface (type 1 site if twosite model) retention capacities for IONP $(S_{1,\text{max}}^p)$ and if applicable, IONP filter ripening rate coefficient (k_{rip}) , rhamnolipid retention capacity $(S_{1,\max}^R)$, as well as rhamnolipid adsorption rate coefficient (k_{ads}), (Table 1). All R^2 values obtained by model fitting were greater than 0.87, indicating that the mathematical models were able to capture both the transport and retention characteristics of nanoparticles in water-saturated 80-100 mesh Ottawa sand. Experimental breakthrough curves and retention profiles, and the mathematical simulation results are shown in Fig. 1 and 3.

3.1.1 Baseline IONP transport in aerobic medium. When IONPs were injected alone with a background aqueous phase of aerobic medium (AE-IONP-1 and AE-IONP-2), most were retained, with minimal breakthrough $(C/C_0 < 0.01)$ for the duplicate column experiments (Fig. 1a). Most IONPs were retained near the column inlet (Fig. 1b), exhibiting a hyperexponential shaped retention profile, 82-84 similar to that by others for colloids including latex microparticles, ¹⁷ bacteria, ⁸³ biochar colloids, ³⁷ clay minerals ⁸⁵ and ENPs such as multi-wall carbon nanotubes (MWCNTs), 86,87 graphene, 88 TiO2, 43,89 ZnO (ref. 61) and nanoscale zero valent iron (NZVI). 90,91 As discussed in these prior studies, hyper-exponential retention profiles may result from particle physical straining, 92 colloid population heterogeneities size and/or surface media,94 packing of porous heterogeneous hydrodynamics,95 aggregation/straining⁹⁶ and ripening/ straining.⁵¹ Although physical straining demonstrated as an important mechanism underpinning hyper-exponential retention profiles in many colloid transport studies, 86,87,92 it is an unlikely factor in the present experiments. Based on a critical particle to grain ratio of 0.002, as reported by Bradford et al.,95 a nanoparticle size greater than 330 nm would be required to result in straining in 80-100 mesh Ottawa sand (d_{50} = 165 μ m). However, the IONP influent suspension, which was mixed with background

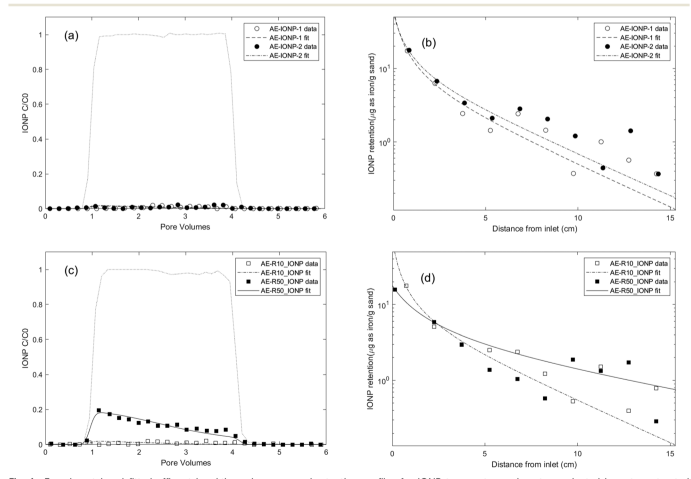


Fig. 1 Experimental and fitted effluent breakthrough curves and retention profiles for IONP transport experiments conducted in water-saturated columns packed with 80-100 mesh Ottawa sand with aerobic medium as background electrolyte at a pore-water velocity of 0.43 m per day using different injection procedures and with different rhamnolipid concentrations. Symbols represent experimental data from each IONP column experiment shown in Table 1. (a) and (b): IONP transport alone column results. (c) and (d): 10 mg L^{-1} and 50 mg L^{-1} rhamnolipid preflood column results.

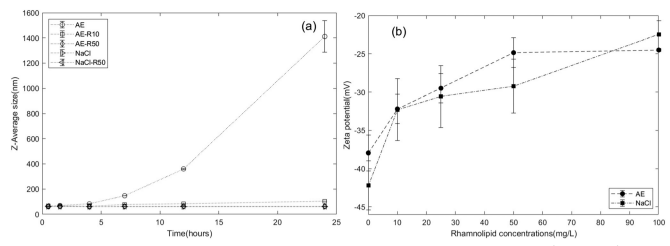


Fig. 2 Changes in (a) IONP diameter over 24 h in aerobic medium(AE) and 10 mM NaCl solution containing 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid and (b) IONP zeta potential as a function of rhamnolipid concentration in background electrolytes.

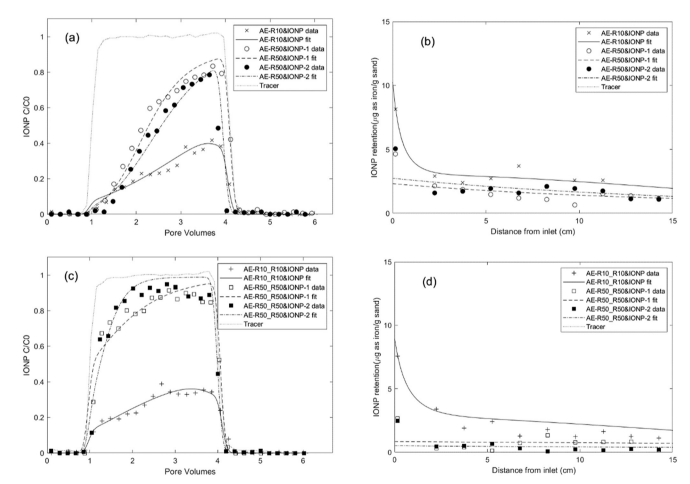


Fig. 3 Experimental and fitted effluent breakthrough curves and retention profiles for IONP transport experiments conducted in water-saturated columns packed with 80-100 mesh Ottawa sand with aerobic medium as background electrolyte at a pore-water velocity of 0.43 m per day using different injection procedures and with different rhamnolipid concentrations. Symbols represent experimental data from each IONP column experiment shown in Table 1. (a) and (b): IONP and rhamnolipid co-injection column results. (c) and (d): rhamnolipid preflood and co-injection with IONP column results.

aerobic medium and measured immediately before introduction to the column, exhibited a mean hydrodynamic diameter of approximately 40 nm with a polydispersity index (PDI) <0.2 (Fig. S4†), which indicates that IONPs were not large enough in the aerobic medium at the time of injection to cause straining. Although the DLS results shown in Fig. 2

indicated that the mean diameter of IONP (z-average size, intensity based) gradually increased from 60 nm to 200-400 nm in the aerobic medium over 12 hours, it is unlikely that IONPs increased to a large enough size during the column residence time (approximately 8.5 h) to cause physical straining at the column inlet. Therefore, the hyperexponential decay retention profile was not attributed to nanoparticle aggregation/straining prior to injection. Meanwhile, the small PDI and narrow surface potential distribution (Fig. S1c†) of nanoparticle in influent suspension suggested the nanoparticles were monodisperse and charged, uniformly thus, the colloid population heterogeneities should not account for the hyper-exponential profile. The employed column packing procedures also resulted in homogeneously packed porous medium along the column,²⁷ as reflected by the symmetrical shape of nonreactive tracer BTCs, and thus, heterogeneity of porous media is also an unlikely explanation. Bradford et al. 95 pointed out the hyper-exponential retention profile could be attributed to inlet hydrodynamics under unfavourable conditions, i.e. the colloid flux near column inlet dominated colloid mass transfer to sand surfaces and consequently resulted in higher deposition at column inlet. With increasing flow velocity, this flux could contribute significantly to hyper-exponential retention profile. However, our column studies were run at a slow pore water velocity (0.43 m per day) and thus inlet hydrodynamics would only play a minor role for the hyperexponential profiles presented here.99

Energy profile calculations based on Derjaguin, Landau, Verwey, and Overbeek (DLVO) theory (Fig. S5a†) revealed a secondary minimum and a small energy barrier (less than 1 kT) between IONPs in the aerobic medium, in agreement with DLS results that IONPs tended to agglomerate in the aerobic medium. The interaction energy between IONPs and the quartz sand surface was also calculated using DLVO theory (Fig. S6†) and resulted in an energy barrier of 4.8 kT. This suggests that IONP attachment might be more favourable to the surface of other IONPs than to that of quartz sands. Thus, we attribute the observed hyperexponential retention profiles to physical straining induced by a filter ripening process, 100 i.e., we assume that IONPs preferentially deposit onto previously attached nanoparticles, creating increasing layers of retained nanoparticles over time near the column inlet96 and induces straining process.101 This ripening/straining phenomenon has been observed previously for other ENPs (e.g., gold nanoparticles, 96 TiO2 (ref. 102 and 103)). For example, Chen et al. 103 observed 10 2 breakthrough curves that displayed a transition from blocking to filter ripening with increasing ionic strength, and hyper-exponential retention profiles occurring at high ionic strength condition. They concluded that net attractive nanoparticle-nanoparticle interactions were responsible for the ripening and subsequent physical straining processes. Mathematical simulations of the experimental data also showed that the two-site mathematical model incorporating a filter ripening process more accurately captured the hyperexponential decay feature of retention profiles compared to one-site models (Fig. S9†). The fitted filter ripening rate coefficients (k_{rip}) were 1.16 and 1.23 times greater than the nanoparticle attachment rate coefficient for the replicate columns, respectively (Table 1). These results imply that filter ripening played a more important role than nanoparticle attachment in the IONP retention process for IONP transport within an aerobic medium background solution. This is also consistent with DLVO calculations, which show that nanoparticle-nanoparticle interactions experience a slightly smaller energy barrier than nanoparticle-sand interactions.

IONP transport with rhamnolipid-preflood. Pretreatment of the packed column with a 10 mg L⁻¹ rhamnolipid solution prior to nanoparticle introduction appeared to have no discernible effect on IONP transport behaviour, as minimal nanoparticle breakthrough and hyperstill exponential retention profiles were (Fig. 1c and d). However, when the concentration of rhamnolipid in the pretreatment solution was increased to 50 mg L⁻¹, IONPs were readily detected in the column effluent (12.9% of total applied mass). In addition, the IONP breakthrough curves obtained following a preflood of 50 mg L⁻¹ rhamnolipid displayed a downward trend with a maximum relative concentration (C/C_0) achieved at breakthrough, with a gradually decreasing concentration over the pulse width, consistent with a filter ripening process. 23,51,104 This observation supported our hypothesis of filter ripening when IONPs were introduced alone with aerobic medium as the background electrolyte. The existence of a filter ripening process was also supported by SEM imaging results (Fig. 4a and b), which show adsorbed agglomerates of IONP on grain surfaces near the column inlet (AE-R10_IONP).

To further understand the effect of the rhamnolipid preflood on IONP mobility, additional batch studies and column experiments were performed, using only rhamnolipid as the solute, to obtain baseline data of rhamnolipid sorption and transport (Fig. S7 and S8†) in 80-100 mesh Ottawa sand. As shown in Fig. S8,† the Langmuir model provided a good fit to the batch adsorption data, yielding a maximum adsorption capacity of 15.38 and 16.36 µg rhamnolipid per g sand in aerobic medium ($R^2 = 0.895$) and 10 mM NaCl ($R^2 =$ 0.963), respectively. These data support the use of a Langmuir-type limiting capacity or maximum surface coverage for type 1 sites in the model. In column experiments, when 10 mg L-1 and 50 mg L-1 rhamnolipid were injected alone in 80-100 mesh Ottawa sand with aerobic medium as the background electrolyte, 12.9% and 17.9% of the injected mass was retained, corresponding to approximately 1.9 µg and approximately 8.0 µg rhamnolipid retention per gram sand, respectively. Using a rate limited sorption model with a desorption term (with the same mathematical form as given by eqn (1) and (3)), the fitted adsorption capacities were determined to be 3.31 \pm 2.51 μg g^{-1} and 11.84 \pm 5.11 μg g^{-1} for rhamnolipid at concentrations of 10 mg L⁻¹ and 50 mg L⁻¹ (Table S2†), respectively, which

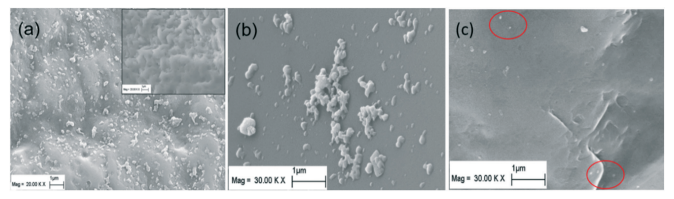


Fig. 4 SEM images of (a) retained IONPs near the column inlet (inset picture shows the clean sand surface before IONP injection), (b) an increased magnification view of deposited IONP aggregates formed due to filter ripening and (c) retained IONPs on sand surface when injected with rhamnolipid.

were also consistent with batch adsorption results at these concentrations (4 μg g⁻¹ and 15 μg g⁻¹). The fitted rhamnolipid adsorption rate coefficient was at least 10 times larger than desorption rate coefficient, which suggested that rhamnolipid desorption from the quartz sand was negligible compared to the adsorption process. Using the fitted parameters, rhamnolipid retention profiles were also simulated (Fig. S7b†), which indicated that the absorbed rhamnolipid was distributed uniformly over the column length, consistent with the limiting sorption capacity assumption of the mathematical model.

The results from rhamnolipid-only batch and column studies suggested that the observed increase in IONP mobility following a 50 mg L⁻¹ rhamnolipid preflood was due to blocking of attachment sites on mineral surfaces, thereby reducing nanoparticle deposition. 39,50,105 However, when the rhamnolipid concentration was lowered (i.e., preflood with 10 mg L⁻¹ rhamnolipid), a sufficient number of attachment sites were still available to retain the applied IONP mass. Similar observations of increased nanoparticle mobility after pretreatment with surfactants¹⁰⁵/polymers²⁵/NOMs³⁹ have been reported. For example, Kmetz et al. 105 pretreated a sand column with 1000 mg L-1 HEC-10, followed by the introduction of 2500 mg L⁻¹ magnetite nanoparticles, which resulted in 93% mass recovery of the injected nanoparticles. More recently, Xu et al.39 preflushed soil columns with dissolved organic matter (DOM) and observed a 2- to 5-fold increase in the mobility of hydroxyapatite nanoparticles (nHAP) compared to untreated soil columns with increasing preflood DOM concentrations. The authors concluded that DOM was adsorbed on favourable attachment sites and decreased soil surface potentials by 2-5 mV, as well as provided steric hindrance to nHAP deposition.

The two-site model, which incorporated filter ripening, was able to reproduce the declining plateau of the BTC and the hyper-exponentially decreased retention profiles. Compared to modelling results obtained for the IONP baseline experiment, the fitted type 1 site maximum capacity decreased to 4.03 and 2.24 µg as Fe per g sand, after a preflood of 10 mg L⁻¹ and 50 mg L⁻¹ rhamnolipid solution, respectively, corresponding to a reduction in the number of attachment sites. Additionally, the fitted attachment rate coefficient after a preflood with 50 mg L⁻¹ rhamnolipid was approximately 2.5 times smaller (0.19 h⁻¹) than those obtained from the baseline column experiment (0.49 h⁻¹ and 0.52 h⁻¹), suggesting that adsorption of rhamnolipid on sand surface altered interactions between nanoparticles and the porous medium.

3.1.3 IONP transport with rhamnolipid co-injection. To explore the effect of rhamnolipid in the aqueous phase on IONP transport behaviour, a series of batch experiments was first conducted to evaluate the impact of rhamnolipid addition on IONP stability in the aerobic medium. As shown in Fig. 2b, the zeta potential of nanoparticle decreased (less negative) upon the introduction of 10 mg L⁻¹ rhamnolipid to IONP suspension, and further addition of rhamnolipid resulted in an even lower IONP zeta potential. These findings indicated direct adsorption of rhamnolipid onto IONP surfaces.⁵⁰ Aggregation kinetic studies indicated, without rhamnolipid, IONPs aggregated in aerobic medium and formed large aggregates (Fig. 2a), which was attributed to the high ionic strength (50.4 mM) and complex ion composition (e.g., Ca²⁺, Mg²⁺) of aerobic medium. However, IONPs became stable in the aerobic medium with 10 or 50 mg L^{-1} rhamnolipid (Fig. 2a). This effect was likely due to increased steric repulsion resulting from adsorption of rhamnolipid on the IONP surface. 106 The concentrations of free rhamnolipid in the IONP suspensions after surface adsorption were determined by centrifugation at 4200 rpm for 2 hours. The rhamnolipid concentrations in the supernatant were 9.0 \pm 0.4 mg L⁻¹ and 43.0 \pm 0.7 mg L⁻¹ for a dose addition of 10 and 50 mg L⁻¹, respectively. Although most of the rhamnolipid existed in the aqueous phase, these data indicate a volume fraction of 0.2% and 1.9% of the outer layer is composed of absorbed rhamnolipid (see ESI†). Using these values, DLVO calculations accounting for repulsive interactions (osmotic interaction and elastic repulsion) between rhamnolipid coated nanoparticles showed a much larger energy barrier (>6 kT and >30 kT) for IONP suspensions containing 10 and 50 mg L⁻¹

rhamnolipid compared to IONP suspensions without rhamnolipid in aerobic medium (Fig. S5a†).

Fig. 3a and b showed the effect of co-injecting rhamnolipid with IONPs on the IONP breakthrough curves and retention profiles in 80-100 mesh Ottawa sand, respectively. Compared to the minimal breakthrough measured when IONPs were injected alone (Fig. 1), considerably more breakthrough of IONP was measured in the presence of 10 or 50 mg L⁻¹ rhamnolipid. Specifically, addition of 10 mg L⁻¹ rhamnolipid resulted in 24.9% of the applied mass recovered in the column effluent, while coinjection with 50 mg L⁻¹ rhamnolipid yielded 43.2% and 56.6% IONP breakthrough for duplicate columns. In contrast to IONP baseline columns where near column inlet a large amount of nanoparticle aggregates were observed due to filter ripening, the deposited nanoparticles in IONP and rhamnolipid co-injection columns sparsely distributed and remained monodisperse on the sand surface, as shown in Fig. 4c (nanoparticles are highlighted in red circles). These data indicated that the observed increases in nanoparticle mobility in the presence of rhamnolipid are due to the combined effects of nanoparticle stabilization and the site blocking mechanisms.

All IONP breakthrough curves obtained for the coinjection columns exhibited asymmetrical shapes with the relative concentration (C/C_0) in the column effluent gradually increasing to a maximum and then rapidly declining to 0 after nanoparticle injection ceased (Fig. 3a), consistent with a maximum retention capacity (S_{max}) and irreversible attachment ($k_{\text{det}} = 0$). The retention profiles obtained for IONP on the solid phase displayed a monotonically decreasing trend with the highest IONP deposition at the column inlet (Fig. 3b). For the column co-injected with 10 mg L⁻¹ rhamnolipid, IONP retention near the column inlet may be slightly impacted by filter ripening, as the fitted ripening rate (k_{rip}) decreased to 0.06 h⁻¹, only one fifth of attachment rate (0.32 h⁻¹). For 50 mg L⁻¹ rhamnolipid co-injection columns, model fitting of the data indicated that rhamnolipid adsorption rates were approximately 4-5 times smaller than the IONP attachment rates (e.g., $0.09 \text{ h}^{-1} \text{ vs. } 0.38$ h⁻¹), suggesting that IONP exhibited more favourable deposition than rhamnolipid. This meant that the IONPs were able to effectively occupy more attachment sites near the column inlet compared to the rhamnolipid. As IONP mass in the aqueous phase was depleted with travel distance, rhamnolipid adsorption then became more dominant. As a consequence, the IONP retention profile decreased monotonically, with greatest retention near the column inlet. These results were in contrast to prior studies which showed that a residual coating polymer rapidly adsorbed near the column inlet, resulting in attachment site blocking and an increasing QD retention profile with distance from the inlet. 25,44 The multi-constituent model yielded better fit to the experimental data compared to a single-constituent model (Fig. S10†), demonstrating the importance of accounting for competitive attachment to accurately simulate

IONP transport and retention behaviour in this complex system. Additionally, the fitted type 1 site retention capacities (Table 1) were similar (approximately 5.98–7.67 μg g⁻¹ sand) for co-injection columns at approximately 5 mg L⁻¹ IONP (as iron) co-injected with 10 mg L⁻¹ rhamnolipid (AE-R10&IONP) and with 50 mg L⁻¹ rhamnolipid (AE-R50&IONP, duplicates), further demonstrating the robustness of the (two-site) multiconstituent mathematical model.

3.1.4 IONP transport with rhamnolipid preflood and coinjection. For the final three column experiments, the preflood process was applied and the IONP suspension was co-injected with rhamnolipid to evaluate IONP mobility with rhamnolipid in both aqueous phase and adsorbed onto the solid phase. As shown in Table 1, compared to columns with only pre-flood or co-injection, IONP mobility was further improved to 28.4%, 78.7% and 83.0% breakthrough of the applied mass, respectively. For the column pre-flushed and co-injected with 10 mg L⁻¹ rhamnolipid (Fig. 3c and d), the BTCs and retention profiles had similar features compared to those obtained for 10 mg L⁻¹ rhamnolipid co-injection (Fig. 3a and b), indicating that the preflood with 10 mg L⁻¹ rhamnolipid had little additional effect on IONP transport. For the duplicate columns conducted with a 50 mg L⁻¹ rhamnolipid preflood and co-injection, the relative concentrations (C/C_0) of IONP rapidly increased to a maximum plateau of approximately 0.9 after 2 PVs of IONP were injected, and sharply decreased to 0 once the pulse injection ceased. The retention profiles were relatively flat with a slightly higher amount of IONP (approximately 3 µg iron per g sand) retained near the column inlet. The (twosite) multi-constituent model was able to capture the transport and retention profiles of the IONP in these columns with R^2 greater than 0.87. The corresponding values of nanoparticle attachment rate coefficient, and type 1 site retention capacities $(S_{1,\text{max}}^p)$ were approximately 5-fold and 3-fold smaller than those obtained from IONP only columns, respectively. These findings provided further support for the stabilizing effect and competition for surface sites attributed to rhamnolipid in the aqueous phase and adsorbed on the solid phase, respectively.

3.2 IONP transport in 10 mM NaCl

A second set of column experiments was performed using 10 mM NaCl as the background electrolyte to assess the effect of rhamnolipid on IONP transport and retention under a lower ionic strength condition. Baseline transport and retention data for IONP or rhamnolipid alone in 10 mM NaCl were first obtained, followed by IONP co-injection with 50 mg L⁻¹ rhamnolipid, and finally preflood and co-injection of IONP with 50 mg L⁻¹ rhamnolipid. The resulting IONP BTCs and retention profiles are shown in Fig. 5. For the duplicate IONP-alone column experiments, breakthrough of IONP represented 12.3% and 14.9% of the applied mass, which were higher than the minimal breakthrough obtained for IONP baseline experiments with aerobic medium as the

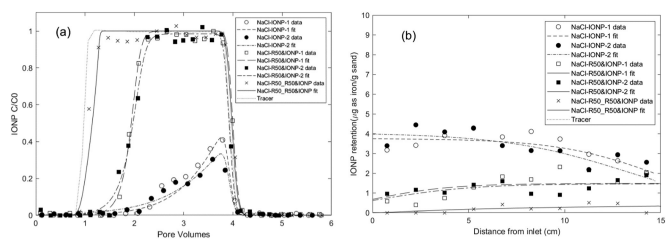


Fig. 5 Experimental and fitted (a) effluent breakthrough curves and (b) retention profiles for IONP column experiments using 10 mM NaCl solution as background electrolyte. First, IONPs were injected alone in duplicate columns (NaCl-IONP-1 and -2), then IONPs were co-injected with 50 mg L⁻¹ rhamnolipid in duplicate columns (NaCl-R50&IONP-1 and -2), and finally the column was pre-flushed and IONPs were co-injected with 50 mg L⁻¹ rhamnolipid (NaCl_R50_R50&IONP). Symbols represent experimental results from IONP column experiments in Table 1.

background solution. This finding is consistent with aggregation kinetic results (Fig. 2a) and DLVO calculations (Fig. S5b†) demonstrating that IONPs exhibit greater stability in 10 mM NaCl compared to the aerobic medium. The improved transport of IONP in 10 mM NaCl is also consistent with prior nanoparticle transport experiments in which mobility increased and retention decreased in lower ionic strength monovalent cation solutions. 103,107,108 In contrast to the hyper-exponentially declining retention profiles observed with aerobic medium (Fig. 1), the retained IONPs in these two columns exhibited a relatively uniform distribution over the length of the columns, consistent with a limiting retention capacity and indicating the absence of filter ripening or straining processes (Fig. 5b). Therefore, a modified filtration theory model²⁴ that incorporates the maximum nanoparticle attachment capacity (S_{max}) was employed to simulate these column data, yielding IONP attachment rate coefficients of 0.5-0.7 h⁻¹ and a maximum retention capacity of approximately 4 µg as iron per g sand. The attachment rates of IONP baseline columns in 10 mM NaCl were similar to the values obtained when using aerobic medium as background, while the retention capacities were slightly smaller than those obtained in the aerobic medium columns, consistent with other nanoparticle transport studies showing that higher IS results in higher retentions. 103,107,109

When IONPs were co-injected with 50 mg L⁻¹ rhamnolipid duplicate columns, enhanced breakthrough nanoparticle was measured (66.9% and 67.3% of the applied mass, Table 1). Additionally, the IONP retention profiles displayed a monotonically increasing trend, with the smallest deposition near the column inlet. These results suggest that the enhanced IONP mobility was related to the competitive effect of adsorbed rhamnolipid which reduced the number of IONP attachment sites. A one-site multi-constituent model developed in our previous work225 successfully captured the delayed IONP effluent breakthrough and the reduction in

solid phase IONP concentrations near the column inlet. The resulting rhamnolipid adsorption rate coefficients were nearly 3-times greater than the nanoparticle attachment rates, which supported the assumption that rhamnolipid outcompeted IONPs for attachment sites. Notably, the fitted maximum retention capacities $(S_{1,\text{max}}^p)$ for IONP were approximately 2-times greater than those obtained without rhamnolipid (4.04 μg g⁻¹ νs . 8.56 μg g⁻¹, respectively), indicating that more sites were available for IONP attachment in the absence of rhamnolipid.

When the column was preflushed and co-injected with 50 mg L⁻¹ rhamnolipid, even greater breakthrough of IONPs was measured (95.3% of injected mass), yielding a BTC that nearly coincided with that measured for non-reactive tracer (Fig. 5a). The corresponding retention profile exhibited a gradual decrease in concentration with distance, with minimal IONP deposition near column inlet, indicating attachment between rhamnolipid competitive nanoparticles. The fitted maximum nanoparticle attachment capacity following the rhamnolipid preflood was determined to be 2.43 µg g⁻¹ using one-site multi-constituent model, which represents a 2- and 4-fold reduction compared to the values obtained from the baseline (no rhamnolipid) and coinjection columns respectively, further demonstrating that adsorbed rhamnolipid enhances nanoparticle mobility by competing for attachment sites.

Conclusions

The effects of rhamnolipid biosurfactant on the transport and retention behaviour of IONP in 80-100 mesh Ottawa sands were investigated using coupled experimental and mathematical modelling studies. When using aerobic medium (IS = 50.4 mM) as the background electrolyte, IONPs were almost completely retained within the column due to the instability of nanoparticles and filter ripening. A preflood

with either 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid resulted in incremental increases in IONP mobility, which were attributed to rhamnolipid adsorption that blocked potential nanoparticle attachment sites. When nanoparticles were coinjected with 10 mg L⁻¹ or 50 mg L⁻¹ rhamnolipid, greater breakthrough of IONP was observed, which was attributed to a combination of improved suspension stability and the site blocking mechanism. The greatest IONP mobility was observed when the columns were preflushed and co-injected with 50 mg L⁻¹ rhamnolipid. These findings were confirmed under a low ionic strength condition (10 mM NaCl), where IONP mobility was considerably greater and the competitive effect of rhamnolipid adsorption on IONP attachment was more apparent. Simulations of the experimental BTCs and retention profiles using mathematical models that accounted for nanoparticle filter ripening processes and/or competitive adsorption were able to capture experimental measurements and provided attachment rate parameters and maximum retention capacities that supported the interpretation of rhamnolipid effects on IONP transport and retention behaviour. These findings demonstrate that biosurfactants can have a substantial effect on the mobility and retention of nanoparticles in porous media and should be taken into account when evaluating the environmental fate of both natural and engineered nanoparticles in the subsurface.

Conflicts of interest

There are no conflicts to declare.

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