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Mobility of Solid and Porous Hollow SiO₂ Nanoparticles in Saturated Porous Media: Impacts of Surface and Particle Structure

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Graphical Abstract:



HIGHLIGHTS:

- The porous shell in porous hollow silica nanoparticles (PHSN) resulted in significantly different particle physicochemical properties compared to solid silica nanoparticles (SSN) of similar size.
- Higher surface roughness in PHSN than that of SSN was responsible for the deviations in standard DLVO theory.
- High ionic strength (IS) and low pH decreased mobility and increased retention for both particle populations.
- Retention was 3-fold higher for PHSN compared to SSN at the highest IS and lowest pH.

ABSTRACT

Silica nanoparticles (SiO₂ NPs) are of increasing interest in nano-enabled agriculture, particularly as nanocarriers for the targeted delivery of agrochemicals. The direct application of these in agricultural soils may lead to the release of SiO₂ NPs in the environment. Although some studies have investigated transport of solid SiO₂ NPs in porous media, there is a knowledge gap on how different SiO₂ NP structures incorporating significant porosities can affect the mobility of such particles under different conditions. Herein, we investigated the effect of pH and ionic strength (IS) on the transport of two distinct structures of SiO₂ NPs, namely solid SiO₂ NPs (SSNs) and porous hollow SiO₂ NPs (PHSN), of comparable sizes (~200 nm). Decreasing pH and increasing ionic strength reduced the mobility of PHSNs in sand-packed columns more significantly than for SSNs. The deposition of PHSNs was approximately 3 times greater than that of SSNs when pH was 4.5 and IS 100 mM. The results are non-intuitive given that PHSN has a lower density and the same chemical composition of SSN but can be explained by the greater surface roughness and ten-fold greater specific surface area of PHSN, and their impacts on van der Waals and electrostatic interaction energies.

Introduction

Rapid advances in nano-enabled agriculture have been focused on making significant improvements in enhancing crop yields, mitigating energy and water footprints, and reducing unintended pollution from the use of fertilizers and pesticides [1-3]. The latter is achieved through precise delivery of pesticides and fertilizers using nanocarriers, which can dose these agents to plant tissues in a targeted manner and with a higher efficiency than traditional pesticide and fertilizer formulations [3-8]. Porous nanosilica is a promising candidate for pesticide and fertilizer nanocarriers because SiO₂ is an earth-abundant, biocompatible material [9, 10] that promotes plant growth, and provides resistance towards pathogens and unfavorable environmental conditions [10-12]. SiO₂ nanocarriers may be introduced to agricultural soils either through direct application or through indirect releases such as discharges to soils following foliar application [13-17]. It is important to investigate the mobility of silica nanocarriers to account for their environmental transport and that of their cargo. Although several studies have investigated the transport of SSNs in geologic deposits [18-21], it is unclear how the differences in particle structure, in particular the high porosity of silica nanocarriers, fundamentally influence their mobility in porous media.

Among studies on SSN transport, Wang, et al. [20] reported that particle size and concentration influenced the transport and retention profiles of SSN in porous media. Small SSN of 8 nm diameter caused higher retention and, thus, less mobility when compared to SSN with mean diameter of 52 nm. Moreover, that study showed that increased ionic strength reduced the overall SSN mobility in the porous medium. Zhang, et al. [22] reported that humic acid improved mobility of SiO₂ nanoparticles in saturated porous media because of enhanced electrostatic forces. HonetschlÄgerová, et al. [23] showed that coating nanoscale zerovalent iron (nZVI) particles with a SiO₂ shell enhanced their colloidal stability and mobility in porous media compared to bare nZVI, which have high aggregation tendency and thus are very colloidally unstable.

Common parameters that influence the mobility of nano- and micro-colloids in porous media are (i) particle surface charge and coating [23], (ii) particle size [20], (iii) ionic strength [24], (iv) pH [25], and (v) temperature [26]. However, few studies have analyzed the effects of the structure of nanoparticles. For SiO₂ nanoparticles, while most studies investigate the transport of SSNs, most applications in agriculture focus on the use of mesoporous SiO₂, such as MCM-41 [27-31], and SiO₂ nanoshells, such as PHSN [32-36], which are structurally different from SSNs and, therefore, may lead to different transport profiles. Given the increasing use of silica nanocarriers in agriculture, and the increased potential for release into soils, there is a critical knowledge gap on the transport behavior of nanocarriers in natural porous media, and whether their transport differ from those of SSN which have been studied before.

The objective of this study was to evaluate how the porous structure of the SiO₂ NPs influences their mobility. Two SiO₂ NPs with very distinct structure and similar size were synthesized, namely (i) SSNs which are solid spheres and (ii) PHSNs which are porous and hollow and represent nanocarriers with high porosity. The synthesis methods ensured that the particles were composed only of SiO₂. Experiments were conducted to assess the colloidal stability and mobility of these two particles in saturated, acid-washed, sand-packed columns, over a range of pH and ionic strength (IS). Furthermore, theoretical Derjaguin-Landau-Verwey-Overbeek (DLVO) interaction energies and single collector contact efficiency calculations were performed to investigate how porosity and density differences in the two types of particles influenced colloidal stability and deposition on collector surfaces. A fundamental understanding of the impacts of particle structure on its mobility is essential before investigation of the effects of various environmental conditions and complex particle compositions.

Materials and Methods

Porous Media. White quartz sand with 50-70 mesh particle size (Sigma-Aldrich) was used as the porous medium in this study. Scanning electron microscopy was used to confirm the sand grain morphology and size (Figure S1). The average grain diameter of the quartz sand was 250 μ m, with diameters ranging from 210 to 297 μ m. Prior to use, the sand was treated with HNO₃ (70% v/v) for 16 hours to remove metal oxides and other impurities as reported elsewhere [37-40]. The acid-washed sand was thoroughly rinsed with DI water (ASTM Type 1, Thermo Fisher) followed by three 20-minute cycles of sonication in water bath. The cloudy

DI water was replaced at the end of each cycle. The electrical potential on the surface of the sand particles was quantified by measuring the zeta potential under varying conditions of pH and IS, remaining negatively charged across the board (Table S1).

SiO₂ Nanoparticles. SSNs were synthesized following a protocol based on the Stöber method [41]. A solution of anhydrous ethanol (200 mL, 100%, Commercial Alcohols, Canada) and NH₄OH (15 mL, 28% NH₃ basis, Sigma-Aldrich) was stirred under 400 rpm for 30 minutes to ensure complete mixing. Then, 10 mL of tetraethyl orthosilicate (TEOS 98%, Sigma-Aldrich) was added dropwise at a rate of 1 mL/min. After 5 hours, the reaction was stopped by drying the cloudy suspension overnight at 80°C. Finally, the nanoparticles were calcined for 5h under 500°C.

PHSNs were synthesized based on a protocol previously described [36]. In summary, a solution of DI water (125 mL), anhydrous ethanol (75 mL), NH₄OH (7.5 mL), hexadecyltrimethylammonium bromide (CTAB, 300 mg, Sigma-Aldrich) and Pluronic P123 (850 mg, Sigma-Aldrich) was stirred under 1000 rpm for 1 hour until reagents were completely mixed. Then, 10 mL of TEOS was added dropwise at a rate of 0.75 mL/min. After 5 hours, the reaction was stopped by drying the cloudy suspension overnight at 80°C. Finally, the nanoparticles were calcined for 5h under 500°C.

The morphology of the nanoparticles was characterized by TEM using a Philips model CM200 TEM at an acceleration voltage of 200 kV. The particle size distribution was characterized through DLS using Malvern Zetasizer Nano ZS (Malvern Instruments, UK). DLS analysis was conducted using SSN and PHSN suspended in DI water with a concentration of 100 ppm. The specific surface area was determined using the BET method through nitrogen sorption/desorption experiments using a Quantachrome Autosorb-1 (Quantachrome GmbH & Co., Netherlands). FTIR spectra were obtained for both SSN and PHSN with a Spectrum II (Perkin Elmer) Spectrometer with a single bounce diamond crystal. Spectra were recorded in the range from 4000 to 400 cm⁻¹ at a resolution of 1 cm⁻¹.

Solution Chemistry. 100 ppm of SSNs and PHSNs were prepared by suspending the dried SiO_2 powder obtained from the synthesis in DI water, followed by sonication in water bath for 30 minutes. IS was adjusted in the nanoparticle suspensions and background solutions

using NaCl (ACS grade, Sigma-Aldrich). The pH was adjusted dropwise with 0.1 M HCl and 0.1 M NaOH, and the counter-ions (Na⁺ and Cl⁻) were taken into account for the final IS of the system.

Column Experiments. A 10-cm long acrylic column (Chromaflex, Fisher) with internal diameter of 1 cm was packed with acid-washed sand using a wet packing technique described in Oliviera, et al. [42]. Prior to the addition of the acid-washed sand, the column was filled with DI water. Then, saturated acid-washed sand was deposited in increments of 1 cm layer at a time while vibrating the column for 20 seconds. This procedure was repeated until the saturated acid-washed sand occupied the whole space within the 10 cm long column, yielding a porosity of 0.391 from fitting the advection dispersion equation to the tracer breakthrough curve (BTC).

At the start of each run, 3 pore volumes (PVs) of DI water were introduced to the system with the aid of a peristaltic pump at a flow rate of 1.25 mL/min, which corresponds to a Darcy velocity of $2.64 \times 10^{-4} \text{ m s}^{-1}$ within our setting. The flow rate remained constant throughout the run and for all the different conditions. 10 PVs of background solution amended with NaCl (25 mM) followed to produce the breakthrough curve (BTC) for the tracer to ensure consistency and reproducibility of the column wet packing procedure. Then, 5 PVs of particle-free electrolyte solution at desired IS and pH, 10 PVs of the particle suspension at 100 ppm at the respective IS and pH. Then 5 PVs of the particle-free electrolyte solution and, finally, 3 PVs of DI water were injected. Ten different experimental conditions were carried out in triplicates as detailed in Table 1. The SiO₂ NP concentration of 100 ppm was selected based on concentrations employed in previous column transport studies for SSN [20] and concentrations relevant to foliar application of SiO₂ NPs [13].

NaCl and particle (SSN and PHSN) concentrations exiting the column were quantified in real-time by UV-Vis spectroscopy (Agilent Technologies, Model 8453) in a quartz flow-through cell (Hellma Analytics, GE, 10 mm path length, 300 μ L volume) at wavelengths of 196 nm, and 350 nm, respectively. Particle size distribution and zeta potential at different experimental conditions were measured using a Zetasizer Nano ZS (Malvern Instruments). Size distribution was determined using the dynamic light scattering (DLS) mode and electrophoretic mobility mode [43].

Table 1. Particle characterization data and calculated DLVO interaction energies by nanoparticle type, pH, IS (mM, electrolyte NaCl), and measurements of zeta potential (mV), Z-average diameter (nm), Polydispersity Index (PdI), depths of primary minima (Φ_{min}), heights

3 of energy barriers (Φ_{max}) and depths of secondary wells (Φ_{sec}) for the DLVO energy profiles. Errors denote standard deviations

4 corresponding to measurement of 3 samples.

Experiment	Structure	pН	IS	Zeta Potential	Z-avg dia.	PdI	Particle-Particle (k _b T)			Particle-Collector (k _b T)		
			(mM)	(mV)	(nm)		Φ_{\min}	Φ_{max}	Φ_{sec}	Φ_{\min}	Φ_{max}	Φ_{sec}
Exp 1	SSN	4.5	1	-34.6 ± 0.9	238 ± 1.9	0.14 + 0.01	-160.4	120.2	/	-789.2	178.7	/
Exp 2	SSN	6.5	1	-52.5 ± 0.8	229 ± 5.4	0.12 ± 0.02	-4.5	255.7	/	-555.9	355.4	/
Exp 3	SSN	9.5	1	-71.3 ± 1.3	201 ± 4.9	0.05 ± 0.02	152.4	373.2	/	-289.6	495.5	/
Exp 4	SSN	9.5	10	-57.3 ± 1.1	214 ± 4.8	0.07 ± 0.03	33.2	238.0	/	-381.7	358.5	/
Exp 5	SSN	9.5	100	-37.4 ± 3.0	220 ± 3.6	0.07 ± 0.02	-132.5	82.8	-0.6	-587.9	171.2	-2.0
Exp 6	PHSN	4.5	1	-10.7 ± 0.7	240 ± 1.1	0.20 ± 0.02	-142.0	10	/	-791.3	49.1	/
Exp 7	PHSN	6.5	1	-29.2 ± 1.1	239 ± 1.8	0.19 ± 0.02	-52.6	90.4	/	-601.3	210.9	/
Exp 8	PHSN	9.5	1	-42.5 ± 2.0	221 ± 1.9	0.20 ± 0.01	47.0	173.8	/	-387.0	349.3	/
Exp 9	PHSN	9.5	10	-42.1 ± 1.5	220 ± 3.1	0.27 ± 0.03	42.3	155.8	/	-391.6	296.7	/
Exp 10	PHSN	9.5	100	-31.3 ± 0.5	239 ± 2.0	0.18 ± 0.02	-41.6	72.3	-0.3	-536.6	158.2	-1.9

7 Interaction Energies and Collector Efficiency. The van der Waals forces and repulsive 8 electrostatics forces for SSN and PHSN for the different experimental conditions were calculated to determine the particle-particle and particle-collector interaction energies 9 10 according to the classical DLVO model (Equation 1) using the expressions proposed by 11 Gregory [44] (full list of equations and theoretical considerations in Supporting Information). 12 Interaction energies due to Born were not accounted for because of their negligible 13 magnitudes [45-47]. The Hamaker constant was calculated using the expressions proposed 14 by Lipkin, et al. [48], and accounted for the contributions of water core and silica shell for 15 PHSNs The full list of physical properties used to compute the DLVO energy profiles for 16 SSNs and PHSNs can be found in Table S2. For PHSNs, the interior and exterior fluid 17 compositions were considered to be the same, thus the van der Waals energy profile was 18 calculated by subtracting the resulting forces generated by a particle with diameter 19 corresponding to the inner diameter of the PHSN from the resulting forces calculated for a 20 particle with diameter corresponding to the outer diameter of the PHSN [49, 50] as per 21 Equation 2.

$$V_{tot} = V_{vdW} + V_{edl} \tag{1}$$

22

$$V_{vdW} = V_{vdW}^{out} - V_{vdW}^{in}$$
(2)

The depths of primary minima (Φ_{min}), heights of energy barriers (Φ_{max}) and depths of secondary wells (Φ_{sec}) for the DLVO energy profiles are summarized in Table 1.

The single collector contact efficiency was calculated based on the empirical expressions derived from the following studies: Yao, et al. [51], Rajagopalan and Tien [52], Tufenkji and Elimelech [53], Ma, et al. [54], Nelson and Ginn [55], Ma, et al. [56], and Messina and Sethi [57]. The comprehensive list of expressions used in this step is presented in Table S3.

29

30

32 **Results and Discussion**

33 SiO₂ NPs Characterization. The TEM images in Figure 1 show that while the SSN 34 synthesized based on the Stöber method (Figure 1a and b) are solid with an average primary particle diameter of 184 nm, the PHSN (Figure 1c and d) are hollow with an average diameter 35 36 of 205 nm. Figure 1b and 1d show a more magnified TEM image of singular particles of 37 SSN and PHSN. The shell thickness of the PHSN ranged from 22 to 38 nm. An important 38 difference in the surface characteristics of the two particle types is the significantly higher 39 surface roughness of PHSN created in part by the highly porous shell surface. The calculated 40 mass of the SSNs and PHSNs were very similar. Thus, the particle number concentration of 41 SSNs and PHSNs were very similar in all experiments.

42 The primary particle size distributions (N = 100) for SSN and PHSN represented in Figure 43 S2a and Figure S2b, respectively, show that both particles have a comparable size 44 distribution. The hydrodynamic diameter size distribution, obtained by DLS for SSNs 45 (Figure 2a) is slightly narrower than for PHSNs (Figure 2b) as reported in Table 1 represented 46 by the polydispersity index (PdI). The DLS measurements were obtained at pH 9.5 and IS 1 47 mM because optimal colloidal stability was obtained at these conditions. Overall, both 48 particle populations were successfully synthesized to yield comparable size and shape, to 49 enable direct comparison of their mobility in porous media.

50 The surface characteristics were investigated using BET and FTIR analyses (Figure S3). 51 Although, SSN and PHSN have relatively similar sizes, the specific surface area of PHSN 52 $(287 \pm 30 \text{ m}^2 \text{ g}^{-1})$ is approximately 10 times greater than the specific surface area of SSN (29 53 $\pm 8 \text{ m}^2 \text{ g}^{-1}$). This is due to the presence of micro-scale pores of around 2.5 nm in the PHSN 54 surface as characterized in previous work [36].

The chemical composition of the SSN and PHSN were identical with no chemical bonds present other than those attributable to silica, as determined by FTIR analyses (Figure S3c). Although both populations of SiO₂ nanoparticles have the same chemical composition, the differences in their structure result in different Hamaker constants (6.59×10^{-21} J for SSNs and 5.79×10^{-21} J for PHSNs) and zeta potentials (Table 1). Given the higher surface area of 60 PHSN, there are more counter-ions around the PHSNs compared to SSNs, resulting in a61 thinner solvation layer and zeta potential closer to zero [58].

62 **pH Effect of the Transport Behavior.** The effect of pH in the transport of SSNs and PHSNs in saturated porous media was evaluated at IS 1 mM for both sets of nanoparticles. The BTCs 63 64 obtained for SSNs (Figure 3a) show that the values for steady-state relative effluent 65 concentration (at 9 PV) decreased as the pH also decreased, from 0.99 at pH 9.5, to 0.95 at 66 pH 6.5 and then 0.91 at pH 4.5, suggesting that transport of SSNs is slightly hindered as the 67 pH becomes more acidic, increasing retention of SSNs in the column. The changes in 68 mobility of SSNs with pH are consistent with another study showing reduced mobility of 69 SiO₂ nanoparticles in carbonate reservoirs at acidic pH [59]. It should be noted that the SSN 70 hydrodynamic diameter as measured by DLS increased slightly with pH decrease from 9.5 71 to 4.5 (from 201 ± 4.9 nm at pH 9.5 to 238 ± 1.9 nm at pH 4.5) suggesting limited aggregation.

72 The DLVO energy profiles for the SSN particle-particle (Figure 4a) and particle-collector 73 interactions (Figure 4c) at IS of 1 mM, show highly unfavorable attachment conditions. 74 Under these conditions, the repulsion forces between SSNs and between SSNs and the sand 75 collector surfaces resulted in elevated energy barriers and absence of secondary minima 76 (Table 1). The DLVO energy profiles for the SSN particle-collector interactions (Figure 4c) 77 support the observations of low retention of SSNs from the BTC trends discussed above 78 (Figure 3a). The calculated particle-particle energy barrier for SSN at pH 4.5 was far greater 79 than 15 k_bT, an approximate threshold for colloidal stability, and thus do not suggest 80 conditions favorable for SSN aggregation. However, the energy barrier was significantly 81 lower at pH 4.5 (120.2 k_bT) compared to pH 9.5 (373.2 k_bT).

82 The values for steady-state relative effluent concentrations in the BTCs for PHSN also 83 decreased with decreasing pH, from 0.94 at pH 9.5 to 0.85 at pH 6.5 and then more 84 substantially to 0.37 at pH 4.5. The extent to which the transport was hindered in PHSNs 85 with decreasing pH was much more significant than that in SSNs. As with SSNs, there was 86 also a small increase in DLS-measured hydrodynamic diameters with decreasing pH (221 \pm 87 1.9 nm at pH 9.5 to 240 ± 1.1 at pH 4.5). Although the DLVO calculations suggest that the 88 primary maximum for particle-particle interactions of PHSN at pH 4.5 was small at 10 kbT, 89 the measured hydrodynamic diameters suggest limited aggregation, comparable to SSNs.

90 At pH 4.5 and IS 1 of mM, the primary energy maximum for PHSN was the lowest (49.1 k_bT) 91 of the 3 pH conditions at the same IS, and thus the DLVO calculations are in qualitative agreement with the experimental observation of significantly low mobility of PHSNs at pH 92 93 4.5. However, there was no secondary minima predicted by DLVO calculations. This 94 suggests that other parameters played a role in the transport of these nanoparticles. Straining 95 is unlikely to be a cause for the significantly higher deposition of PHSN, given that the SSN 96 have similar hydrodynamic diameters at corresponding pH, but had lower deposition. 97 Furthermore, straining is not expected when the colloid diameter (d_c) to sand grain diameter 98 (d_g) ratio (d_c/d_g) is below 0.008 [60]. Here, the d_p/d_g ratio is approximately 0.001 for both 99 SSN and PHSN. According to Xu, et al. [60], straining rates are negligibly small when d_p/d_g 100 < 0.008 and, in this work, such ratio is one order of magnitude lower. There is, however, a 101 gradual increase in C/C_0 with increasing pore volumes, suggesting some blocking or 102 detachment of deposited PHSNs. As a first layer of deposited nanoparticles is formed, the 103 trajectories of the subsequent nanoparticles are significantly impacted by this monolayer of 104 deposited particles, a phenomenon previously referred to as the shadow effect [61]. The 105 extent of the influence of the shadow effect in subsequent particle trajectory is directly related 106 to the hydrodynamic interactions and electrostatic double layer repulsion. Therefore, as the 107 conditions become unfavorable (lower pH and higher IS), repulsion forces decrease, thus 108 increasing both the surface coverage of deposited nanoparticles on the sand grains and the 109 blocking effects in the nanoparticle transport profiles.

IS Effect on the Transport Behavior. The effect of IS in the transport of SSNs and PHSNs 110 111 in saturated porous media was evaluated at pH 9.5 and at the same SiO_2 concentration of 100 ppm. The BTCs for SSNs (Figure 5a) showed that the values for relative effluent 112 113 concentration at 9 PVs decreased as the IS increased, from 0.99 at IS 1 mM, to 0.96 at IS 10 114 mM and then 0.92 at IS 100 mM, which was expected and is consistent with the literature 115 [20, 59]. The addition of salt increases the number of counter-ions migrating to the solvation layer of the nanoparticles, thus decreasing the Debye-length and the repulsive electrostatic 116 117 forces. Although the total interaction energies calculated yielded a lowered energy barrier, a 118 weak secondary minimum (-2 kbT) was obtained only at IS of 100 mM.

The values for relative effluent concentrations at 9 PV in the BTCs for PHSN (Figure 5b) also decreased with increasing IS, from 0.94 at IS 1 mM to 0.88 at IS 10 mM and then 0.34 at IS 100 mM. Once again, the mobility PHSNs was much more affected by the change in IS than for SSNs. The similarity in the BTC shape with changes in pH and IS suggests that similar deposition processes are involved. As for the SSNs at pH 9.5 and IS at 100 mM, a weak secondary minimum for PHSNs was calculated from the interaction energy profiles at an IS of 100 mM.

126 Overall, low pH and high IS decreased the mobility of the nanoparticles in porous media. We 127 observed similar trends between SSN and PHSN: (i) As pH decreased, zeta potential also 128 decreased, which is expected because the isoelectric point of bare SiO_2 is around pH = 2 [62-129 64]. (ii) As IS increased, zeta potential decreased as a result of the stabilization of the excess 130 ions in the electrical double layer of each nanoparticle by the counter-ions from the NaCl 131 added. In both cases, a decrease in zeta potential led to a decrease in the intensity of repulsion 132 forces between nanoparticles and between nanoparticles and sand. However, the increased 133 retention of PHSNs compared to the SSNs, which is non-intuitive because both particles have 134 comparable sizes (Figure 1) and surface chemical composition (Figure 2c). The DLVO 135 interaction energies calculated are qualitatively in agreement with the mobility trends for 136 SSN and PHSN, but the calculated values are not consistent in several instances. For 137 example, if we compare the particle-collector interaction resulting energy profiles for Exp. 5 138 (SSNs, pH 9.5, IS 100 mM, Figure 6c) with Exp. 10 (PHSNs, pH 9.5, IS 100 mM, Figure 6d) the heights of energy barriers were comparable, and yet, the steady-state relative effluent 139 140 concentrations were 0.34 and 0.96 for PHSNs and SSNs, respectively. A theoretical analysis 141 of DLVO interaction forces between SSN or PHSN and collector surfaces suggested 142 relatively small differences in the magnitude of the interaction forces [50], and is in 143 agreement with the calculations in this study. It is likely that the surface structure differences 144 caused by the concave asperities in the PHSNs caused by the pores played an important role 145 in the mobility of these particles.

146 Single Collector Contact Efficiencies (η_0) and Particle-Collector Attachment 147 Efficiencies (α_{pc}). The η_0 values as a function of particle radius (a_p) for SSNs (Figure 7a) and 148 PHSNs (Figure 7b) were calculated based on expressions derived elsewhere [51-56]. The expressions and parameters used to estimate the η_0 can be found in Table S3 and Table S4, respectively.

The average η_0 predicted for SSNs ($a_p = 92 \text{ nm}$) and PHSN ($a_p = 103 \text{ nm}$) were 0.007 and 151 152 0.005, respectively. The η_0 were slightly greater for SSNs, which is contrary to observations 153 of lower mobility of PHSNs in the column experiments. To assess the size difference effect 154 between SSNs and PHSNs, the η_0 ratio was plotted as a function of particle size in Figure 7c. 155 For any particle radius below 100 nm, SSNs and PHSNs have very similar predictions for η_0 . 156 For any particle radius above 100 nm, the predicted η_0 for SSNs surpassed the predicted η_0 157 for PHSNs, indicating more chance of retention for SSNs than for PHSNs. Therefore, η_0 158 could not explain why such inconsistencies with the experimental data exist.

159 The α_{pc} values remained below 0.04 for all experimental conditions with SSNs, whereas they 160 reached 0.54 and 0.59 for PHSNs at pH 4.5 and IS 1 mM, and at pH 9.5 and IS 100 mM, 161 respectively, which were the conditions where retention was more pronounced. The α_{pc} 162 values (Table S5) were calculated with Equation (3) below.

$$\alpha_{\rm pc} = -\frac{2}{3} \frac{d_{\rm c}}{(1-n)L\eta_0} \ln\left(\frac{C}{C_0}\right) \tag{3}$$

163

164 Surface Roughness and the DLVO Theory. Deviations from DLVO predictions due to 165 nanoscale physical heterogeneity have been previously reported [65-74]. These deviations 166 happen because standard DLVO theory assumes that particle and collector surfaces are 167 geometrically smooth and homogeneous, when in reality colloid may possess some degree 168 of roughness [67]. Modeling and experimental data have demonstrated that surface roughness 169 reduces the repulsive energy barriers, and in some cases eliminates them altogether [67-69, 170 72]. For instance, Liang, et al. [71] observed enhanced retention of silver nanoparticles in 171 porous media with higher surface roughness (SR). Retention was 6.8-fold higher in relatively 172 rough sand particles (root mean square roughness, $S_q = 524$ nm) when compared to the 173 experimental condition using relatively smooth sand ($S_q = 93$ nm).

Specifically, two phenomena are relevant: (i) the concave asperities in PHSN surface, causedby the presence of pores in the surface, result in the separation distance among particle-

176 particle and/or particle-collector surfaces becoming larger at the pores, when compared to 177 that of SSNs [69] as depicted in Figure 8. Generally, as the separation distance increases, overall repulsive electrostatic forces generally decay faster than van der Waals interactions 178 179 [75], resulting in enhanced attraction to the collector surface. (ii) The roughness caused by 180 the asperities on PHSN surface alter the flow field around the particle during the column 181 experiments, enhancing resisting adhesive torque and diminishing applied hydrodynamic 182 torque [76]. These effects likely led to enhanced colloid immobilization on the collector in 183 this study.

184 It is also important to note that this work has assessed the transport behavior of SiO2 NPs in 185 saturated sand, and thus, will differ from actual agricultural conditions. Agricultural soils are 186 generally unsaturated, with some exceptions, such as when the soil is irrigated or when it 187 rains. Agricultural soil may also contain high organic matter content and other types of 188 contaminants that could complex with the NPs and influence their mobility. These 189 conditions, however, are not replicated in our experimental setting. Nevertheless, the use of 190 saturated sand served as model towards a first step in assessing the fundamental differences 191 in the transport profile of SSNs and PHSNs and how their structure plays a role in their 192 mobility.

193 Conclusions

Herein, we synthesized, characterized, and investigated the mobility of SSNs and PHSNs in a saturated sand-packed column in various pH and IS conditions. As expected, the zeta potential of both sets of particles approached lower absolute values as pH became more acidic and closer to the isoelectric point of SiO_2 nanoparticles. On the other hand, zeta potential approached lower absolute values as IS increased. The decrease in zeta potential indicates a decrease in the repulsive forces between nanoparticles, which in turn led to a small extent of agglomeration, and higher PdI.

During the column experiments, deposition was enhanced by the decrease of pH and increase of zeta potential, as such conditions deteriorate (acidic pH and high salinity) the overall colloidal stability of SiO₂ nanoparticles in suspension. The PHSNs, however, experienced a higher degree of deposition when compared to SSNs. DLVO energy profiles and single 205 collector contact efficiency values were unable to explain why such discrepancies existed,
206 and surface roughness likely contributed to the different extents of deposition. The surface
207 roughness was not factored into the DLVO calculations because of the high complexity of
208 the calculations and the many assumptions needed. The surface roughness related concave
209 asperities of the pores lowered the repulsive energy barriers and led to more deposition.

This study elucidates how nanoparticle architecture can influence their mobility in porous media and highlights the importance of a thorough experimental analysis of the fate and transport of nanoparticles of different architecture that are likely to be discharged in the environment, such those intended for use in nano-enabled agriculture.

214 **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

217 **CRediT author statement**

Vinicius Bueno: Conceptualization, Methodology, Investigation, Writing-Original Draft,
Visualization. Alessandro Bosi: Conceptualization, Methodology, Investigation, WritingReview & Editing. Tiziana Tosco: Conceptualization, Investigation, Writing Review &
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Figure 1. TEM images of (a and b) SSNs and (c and d) PHSNs. The images on the right exhibit more magnified images of singular (b) SSN and (d) PHSN to show the different features between both structures. Although both structures are spherically shaped, PHSNs feature a hollow core and pores. The latter is directly responsible for the increased surface area and roughness of PHSNs when compared to SSNs.



Figure 2. Particle size distribution at pH 9.5 and IS 1 mM of (a) SSNs and (b) PHSNs. The

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240 width of the PHSN size distribution is broader than that for SSN, indicating a higher PdI as
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confirmed in Table 1.



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Figure 3. BTCs at fixed IS 1 mM and varying pH for (a) SSNs and (b) PHSNs. The shaded zones represent the standard deviation of the particle concentration run in triplicates. Wider shaded areas represent greater standard deviation among replicates. The conditions with no visible shaded areas represent runs with not relevant difference among replicates.



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Figure 4. DLVO energy profiles at fixed IS 1 mM and varying pH for (a) SSN particleparticle interactions, (b) PHSN particle-particle interactions, (c) SSN particle-collector interactions, and (d) PHSN particle-collector interactions.



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Figure 5. BTCs at fixed pH 9.5 and varying IS for (a) SSNs and (b) PHSNs. The shaded zones represent the standard deviation of the particle concentration run in triplicates. Wider shaded areas represent greater standard deviation among replicates. The conditions with no visible shaded areas represent runs with not relevant difference among replicates.





Figure 6. DLVO energy profiles at fixed pH 9.5 and varying IS for (a) SSN particle-particle
interactions, (b) PHSN particle-particle interactions, (c) SSN particle-collector interactions,
and (d) PHSN particle-collector interactions.





Figure 7. Estimates of η_0 for (a) SSNs and (b) PHSNs, and (c) the SSN-PHSN η_0 ratio.



Figure 8. Schematic representation of the distance between sites in the SSN (a) and PHSN (b), and a hypothetical surface. For SSNs, the distance from point *a* to the surface, h_a , and the distance from a relatively distant point *a*' to the same surface, $h_{a'}$, are approximately the same. Meanwhile, in PHSNs, the distances h_b , h_c and h_d may vary significantly, even if they are at the same distance as points *a* and *a*' in scheme (a). Effectively, the overall separation distance between rough spherical surfaces and a hypothetical surface is larger than the distance between smooth spherical surfaces and a hypothetical surface.

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