Surface functionalization-dependent localization and affinity of SiO₂ nanoparticles within the biofilm EPS matrix

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ABSTRACT

The contribution of the biofilm extracellular polymeric substance (EPS) matrix to reduced antimicrobial susceptibility in biofilms is widely recognised. As such, the direct targeting of the EPS matrix is a promising biofilm control strategy that allows for the disruption of the matrix, thereby allowing a subsequent increase in susceptibility to antimicrobial agents. To this end, surface-functionalized nanoparticles (NPs) have received considerable attention. However, the fundamental understanding of the interactions occurring between engineered NPs and the biofilm EPS matrix has not yet been fully elucidated. An insight into the underlying mechanisms involved when a NP interacts with the EPS matrix will aid in the design of more efficient NPs for biofilm control. Here we demonstrate the use of highly specific fluorescent probes in confocal laser scanning microscopy (CLSM) to illustrate the distribution of EPS macromolecules within the biofilm. Thereafter, a three-dimensional (3D) colocalization analysis was used to assess the affinity of differently functionalized silica NPs (SiNPs) and EPS macromolecules from Pseudomonas fluorescens biofilms. Results show that both the charge and surface functional groups of SiNPs dramatically affected the extent to which SiNPs interacted and localized with EPS macromolecules, including proteins, polysaccharides and DNA. Hypotheses are also presented about the possible physicochemical interactions which may be dominant in EPS matrix-NP interactions. This research not only develops an innovative CLSM-based methodology for elucidating biofilm-nanoparticle interactions but also provides a platform on which to build more efficient NP systems for biofilm control.

Introduction

Bacterial biofilms are highly-structured multicellular accumulations of microbes which are enclosed in a self-produced extracellular polymeric substance (EPS) matrix composed of polysaccharides, proteins/enzymes, lipids and exogenous DNA (eDNA), as well as lysed cell debris [1,2]. The EPS matrix allows for the colonization of both biotic and abiotic surfaces and also contributes to the persistence of bacteria at various interfaces by playing a crucial role in the development and survival of the bacterial population [3]. It does this by serving as a scaffold for growth, by aiding in the protection of cells from environmental stresses, by serving as a nutrient reservoir and also by affecting the diffusion of antibacterial agents throughout the biofilm [4]. In addition, biofilms typically grow heterogeneously within complex environments which further contributes to their unique and intricate three-dimensional (3D) architecture and extensive physicochemical gradients [5].

Biofilm control and eradication is of great importance in the processing industries as their accumulation and persistence can lead to significant economic losses stemming from the induction of corrosion in piping [6], contamination of raw material leading to a decrease in product quality [7] or increasing energy usage [8]. In addition to this, bacterial biofilms also have a significant impact in human health and wellbeing [9]. In the food, agricultural and medical sectors, bacterial contamination in the form of biofilms calls for the development of more intensive and novel infection treatment and disinfection methods to control disease outbreaks and reduce food wastage and contamination [10].

Traditionally, biofilm removal strategies have focused on directly targeting the bacterial cells themselves. However, the protection offered by the EPS matrix is believed to lead to a decrease in the effectiveness of antibiofilm agents [11]. Although many bacterial cell-targeting strategies have been employed for biofilm removal, attention is now shifting from targeting individual bacterial cells within the biofilm, to methods for the degradation of the EPS matrix itself [12]. Therefore, the development of novel approaches for the control of unwanted biofilms must include the analysis of the composition of the EPS matrix in relation to the
interactions which occur between the EPS matrix and external anti-biofilm agents.

Engineered nanoparticles (NPs) have shown great promise in combating and eradicating bacterial biofilms [4]; however, there is currently a lack of fundamental understanding of the mechanisms associated with and also the interactions between the EPS matrix and engineered NPs. The biological and physicochemical interactions occurring between NPs and bacterial biofilms are highly complex and dependent on a myriad of factors including the inherent characteristics of the NPs (e.g., size, charge, surface functionalization), EPS composition and abundance (e.g., biomolecules present, maturity/age, density, bacterial species), and the environmental conditions (e.g., temperature, pH, flow conditions, ionic strength) [1]. A fundamental understanding of the interactions occurring in biofilm-NP systems is of great importance in the use of the nanomaterials for future applications in either the study of, or eradication of persistent biofilms. Currently, few studies have evaluated the fundamental interactions which occur in the EPS matrix when bacterial biofilms are exposed to differentially engineered NPs.

Thus far, the in-situ analysis of the localization and possible high-affinity interactions between NPs and EPS matrix macromolecules with NPs has remained a challenge. Colocalization CLSM studies have frequently been used to determine whether two molecules associated with the same structure or with each other [13]. For example, determination of the association of a protein with a specific cellular organelle [14], or whether two different proteins associate with the same structure [15]. The repeated observation of the association and nanometer proximity of two fluorescent probes increases the certainty that there is a biological or physicochemical interaction occurring between them. To this end, many fluorescent probes have been used for colocalization analysis [16–18]. Lectins are proteins which are highly specific for carbohydrates binding without altering their structure. Their use in high resolution imaging represents an opportunity for non-destructive in-situ analyses and thus have been widely used for glycoconjugate biofilm investigations when used in combination with other fluorescent probes [19–21]. Therefore, the combination of a fluorescence lectin-binding analysis [22] and colocalization analysis in CLSM could aid in evaluating the localization and affinity of differently functionalized NPs with fluorescently labelled EPS matrix macromolecules.

In this work we describe a highly sensitive confocal laser scanning microscopy (CLSM) imaging system developed to quantitatively identify the colocalization of various engineered fluorescent silica NPs (SiNPs) with specific macromolecules within the EPS matrix of P. fluorescens WCS365 biofilms. Using this system, we were able to accurately assess the accumulation, localization and binding affinities of these SiNPs based on variations in their surface charge and surface functionalization as well as hypothesize the possible physicochemical interactions which may be dominant in EPS matrix-NP interactions.

Results

Nanoparticle characterisation

Characterization was carried out on four groups of fluorescent-labelled SiNPs (Table 1), that were synthesized and subsequently functionalized through alkyl-silane grafting. The as-synthesized SiNPs have a diameter of 54.0 ± 6.9 nm, as determined by TEM (Fig. 1). In Table 1, the size distributions and surface charge of SiNPs are reported, determined by TEM and DLS analysis and Z-potential measurements, respectively. The functionalization of SiNPs does not affect the size of the NPs core, as shown by the size distribution extrapolated from TEM images of the functionalized SiNPs (Fig. S1). However, the effect of silane grafting can be observed in both hydrodynamic ratio and surface charge (Table 1). All the functionalized NPs show an increase in size, determined by DLS, consistent with silane grafting onto the surface. Furthermore, all the NPs present a low polydispersity index (PDI), which indicates high stability of the aqueous dispersion. Bare-SiNPs show the characteristic negative charge of SiNPs, whereas the positive charge measured for Amine-SiNPs is due to the presence of the –NH$_2$ group. Epoxy-SiNPs exhibit a negative Z-potential, but higher compared to the non-functionalized NPs; this can be ascribed to the hydrophobic character of the epoxy-group. On the other hand, Hydroxy-SiNPs show a higher negative surface charge, consistent with a greater stability of the aqueous dispersion given by the 3-(2,3-dihydroxyproproxy) propyl]-trimethoxysilane ligand.

FT-IR spectra were collected on non-dye loaded SiNPs, in order to eliminate the contribution from the internal fluorescent label (Fig. 2). All the spectra show strong characteristic stretching bands for –OH (3300 - 3700 cm$^{-1}$), Si–O–Si (~1100 cm$^{-1}$) and Si–OH (~850 cm$^{-1}$), as well as Si–OH bending (~950 cm$^{-1}$); the peak at 1634 cm$^{-1}$ is attributed to vibrations of water molecules adsorbed on the NPs surface [23] For the Amine-, Epoxy- and Hydroxy-functionalized SiNPs we can observe diagnostic bands for C–H stretching (2800 - 2900 cm$^{-1}$) and bending (1350 - 1500 cm$^{-1}$), ascribable to the alkyl chains of grafted silanes [24]. It must be pointed out that for the Amine-SiNPs a contribution can be considered of the –NH$_2$ groups vibrations to the band at ~1500 cm$^{-1}$. On the Hydroxy-functionalized SiNPs we can observe a peak at ~670 cm$^{-1}$, which can be assigned to C–OH vibrations [25] The same band is not present in the Epoxy-SiNPs, and this is consistent with a successful epoxy-ring opening reaction.

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![Fig. 1. Representative TEM micrograph for as-synthesized Bare-SiNPs, with particle size distribution (nm) reported in the inset. Scale bar is 50 nm.](image-url)
The bottom layer of the biofilm is composed of well-distributed information (Fig. S2-S5). The biofilm was shown to entrap between 20 and 25% of the SiNPs from solution (Fig. S6) meaning that the relative distribution and architecture of EPS macromolecules within the biofilm (Fig. 3C) and are also found in close proximity to bacterial cells (Fig. 5D, red arrowhead). N-Acetylgalactosamine (NAG)-containing polysaccharides stained by the WGA lectin are distributed in both string-like projections protruding from the microcolonies (Fig. 5E, white arrows) as well as in clusters between bacterial cells (Fig. 5F, red arrowhead). Proteins within the EPS matrix which are stained by Sypro® Orange show a variety of morphologies including rounded clusters (Fig. 6A, white arrows) and surrounding the membranes or dead or dying cells (Fig. 6, white arrowheads). Sypro® Orange was unable to stain proteins on the surface of live cells and thus did not produce any significant fluorescent signal on the cell surface (Fig. 6B, red arrowhead). Sypro® Green staining of the biofilm shows both the presence of both eDNA (Fig. 6C, white arrowheads) and genomic DNA within dead/dying cells (Fig. 6C, white arrows) which are distributed into the spaces between live cells (Fig. 6D). SiNPs within the biofilms are shown as cloud-like distributions or in smaller aggregates and can be found in close proximity to polysaccharides, proteins or eDNA but could possibly also be found attached to another unstained component of the biofilm.

**Colocalization of SiNPs with EPS matrix macromolecules**

Under semi-quantitative and qualitative instances, the colocalization of two probes in separate fluorescent channels (green and red) can be subjectively (and incorrectly) identified by the appearance of a yellow colour when images are superimposed or “merged” [27]. Therefore a systematic colocalization analysis using image processing that compares the relative position and intensity of each pixel would give the most reliable and reproducible results [13]. The terminology for describing colocalization data considers weak colocalization ranging from 0 to 0.49, moderate colocalization as 0.5–0.79, and strong colocalization as 0.8–1.0 [28]. As shown in Fig. 7, Epoxy-SiNPs display the highest overall colocalization with EPS macromolecules compared to the other SiNPs. The highest colocalization ratio was found in interactions between NAG-containing polysaccharides with Epoxy-SiNPs; the only binding pair to show a strong colocalization ratio. Moderate colocalization ratios were found between both Mannose-containing polysaccharides with Epoxy-SiNPs as well as between NAG-containing polysaccharides with Amine-SiNPs. All other SiNPs and probe combinations shows weak colocalization. Colocalization ratios are particularly low for proteins and polysaccharides interacting with Bare-SiNPs, Galactose-containing polysaccharides with Amine-SiNPs and Mannose-containing polysaccharides with Hydroxy-SiNPs.[28]

**Protein binding affinity (in-vitro analysis)**

The protein corona formed when biofilms are exposed to extracted EPS proteins in-vitro (Fig. S7) showed that changes in the SiNPs surface chemistry can alter the amount and type of protein which associates with the SiNP surface. To evaluate the level of binding of various proteins to differently functionalized SiNPs and how protein charge and pl affects this, the SiNPs were exposed to either an EPS protein extract or two model proteins, BSA and RNase, each with different surface chemistry and charge (Fig. 8). The in-silico mapping of the electrostatic surface potentials (at physiological pH) of the model proteins demonstrated an overall negative charge for BSA while RNase showed an overall positive charge (Fig. S8). Bare-SiNPs and Hydroxy-SiNPs showed no significant difference in their protein binding affinity ranging between 20 and 25% of the total protein in solution. In terms of positively charged Amine-SiNPs, there was a significant increase in the binding affinity of BSA while RNase showed a decreased binding affinity and the EPS-extract showed an affinity intermediate to the model proteins. Epoxy-SiNPs showed an overall significant increase in the amount of protein bound as compared to the other SiNPs.
Carbohydrate binding affinity (in-vitro analysis)

To identify the contribution of specific carbohydrates to the level of colocalization shown, SiNPs were exposed to each mono- or polysaccharide independent of the biofilm. Amine-SiNPs were shown to have the highest overall percentage of monosaccharide molecules bound to their surface while glucose was shown to be the monosaccharide which bound to all SiNPs with the highest proportion and galactose showing the lowest overall affinity (Fig. 9A). Both NAG and alginate polysaccharides showed no significant difference (P < 0.05) in binding preference towards each SiNPs but NAG did have a higher binding affinity as compared to alginate (Fig. 9B).

Discussion

Distribution of SiNPs within the EPS matrix

Changes in the NP surface chemistry have been shown to affect how they interact with the biofilm as a whole. For instance, Li et al. demonstrated that the control of NP distribution and penetration into bacterial biofilms is determined by variation of surface properties. This research showed the penetration of only positively charged functionalized CdSe–ZnS core–shell NPs (24 nm diameter) into Escherichia coli biofilms [29], with the changes in the distribution of the NPs being governed by their charge and hydrophobicity. In the current study, even though the entrapment of SiNPs by the biofilm did not vary significantly, the distribution patterns and binding affinity of the SiNPs in CLSM imaging varied greatly, further highlighting the need for in-depth high-resolution analyses. Positively charged Amine-SiNPs showed the highest overall entrapment and accumulation in the biofilm (Fig. 9B) but all functionalized SiNPs were nevertheless able to fully penetrate the biofilms microcolonies regardless of their overall charge or surface groups present (Fig. 4). Changes in the localization and distribution patterns of SiNPs in the EPS matrix based on their surface chemistry displays a contribution of these groups to the affinity of SiNPs to different macromolecules in the EPS matrix and not their propensity to permeate the biofilm.

Fig. 3. CLSM images of 72h P. fluorescens WCS365 biofilms after exposure to (A) Bare-SiNPs, (B) Amine-SiNPs, (C) Epox-SiNPs and (D) Hydroxy-SiNPs. Biofilm cells were stained with Syto9 (green) and SiNPs were labelled with RITC (red). The central images show the horizontal (xy) section 0.5 μm above the substrate surface. Upper and side panels represent z-stack images of the xz and yz planes, respectively. The yellow lines indicate the position of xz and yz planes on the xy section images. White arrows and arrowheads indicate the distribution patterns of SiNPs and white dashed squares represents the centre of the microcolonies. Representative images from three independent experiments are displayed. Scale bar represents 20 μm. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
biofilm structure while eDNA has been shown to provide a structural scaffold for have been shown to cross-link and form complexes with other macromolecules. For example, polysaccharides like PEL and PSL have been shown to cross-link and form complexes with other macromolecules. The structure-to-function relationship of the surface of elucidate the composition of bacterial biofilm secreted by metabolically active cells. DNA in the EPS matrix is either present in the EPS matrix after being released as genomic DNA from lysed cells or as increased likelihood of proteins forming complexes with other EPS molecules with the EPS matrix. While BSA has a high affinity for this positively charged biofilm EPS matrix as well as limited EPS proteinaceous material 

Identification and spatial organisation of EPS matrix macromolecules

The EPS matrix must be assessed both in terms of its inherent 3D architecture and the role of each molecule within the structure, albeit only qualitatively. If a particular molecule is to be targeted for degradation and subsequent biofilm removal, then molecules involved in the structural rigidity and integrity of the biofilm should be identified and the surface of “smart” NPs should be tailored in order to direct the NP towards its target. To this end, the structure-to-function relationship of many biofilm-associated macromolecules have previously been elucidated. For example, polysaccharides like PEL and PSL have been shown to cross-link and form complexes with other macromolecules for the maintenance and development of the microcolony structure while eDNA has been shown to provide a structural scaffold for biofilm development. Few studies have used differential staining to elucidate the composition of bacterial biofilms. Super-resolution microscopy of Vibrio cholerae biofilms using an in vivo labeling strategy allowed for the visualization of the molecular architecture and composition of these biofilms. In the case of P. fluorescens biofilm, no studies have previously used the highly specific nature of fluorescent probes to assess the affinity of functionalized NPs with various EPS macromolecules. For the current P. fluorescens biofilms, the string-like topography of both NAG- and Mannose-containing polysaccharide EPS molecules suggests that they may be useful for stabilizing the 3D structure of the biofilm and can also provide a scaffold on which the biofilm can develop in its early stages of attachment. Galactose-containing polysaccharides can be produced as part of the differentiation in maturation of the biofilm while the variation of morphology and topography of proteins within the EPS suggests the increased likelihood of proteins forming complexes with other EPS biomolecules with the biofilm. The eDNA in biofilms is either present in the EPS matrix after being released as genomic DNA from lysed cells or as eDNA secreted by metabolically active cells. DNA in the EPS matrix has been shown to be essential in intercellular adhesion and furthering biofilm stability. The binding of fluorescently-labelled lectins to multiple targets within biofilms or bioaggregates has been previously demonstrated, especially in the binding of lectins to glycoconjugates and proteins both on the bacterial cell surface as well as within the EPS matrix. However, in the current study, the signal produced by EPS molecule-specific stains was shown to occur independently to that of the signal from live bacterial cells, thus mitigating the possible effects of cell-surface binding and therefore erroneous colocalization results.

In-vitro binding affinity analyses

The use of simplified in-vitro models in biofilm research have been instrumental in addressing some fundamental questions with regard to interaction studies but cannot always be compared to in-situ models which are closer resemble a natural environment. The interactions involved within biofilms are complex; which makes them difficult to study under in-vitro conditions. Both methodologies are important to include so as to increase the ease of transferance to clinical or industrial settings. The protein concentration per mass of P. fluorescens biofilm EPS extracts have been shown to be at least ten times higher than the carbohydrate concentration per mass. Proteins within the biofilm are often closely associated with (or form complexes) with other proteins, polysaccharides and eDNA and this can result in structurally unique niches with multiple exposed surface functional groups for SiNPs to interact with. Due to the increased concentration of proteins in these biofilms, an interaction of SiNPs with a protein component (or more likely protein complex) of the EPS matrix may therefore have a higher probability of occurring. The affinity of proteins for SiNPs was studied using a protein extract precipitated from the biofilm EPS matrix as well as BSA and RNase as model proteins with varying charge and pi. As compared to the other SiNPs, Amine-SiNPs show a unique protein binding pattern. While BSA has a high affinity for this positively charged surface, a significantly lower amount of RNase molecules were bound to the Amine-SiNPs surface. As this phenomenon is only observed in positively charged SiNPs, it suggests a dominant electrostatic contribution. To further demonstrate the implication of surface charge and functionalization in the affinity of proteins for the SiNPs, the exposure of different SiNPs to an EPS protein extract showed a significant variation in the protein corona formed (Fig. S7). On the other hand, the higher binding affinity of EPS proteins for Epoxy-SiNPs when compared to the other SiNPs was again possibly due to the reactivity of the functional groups at

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**Fig. 4.** Penetration profile of P. fluorescens WCS365 biofilms after exposure to (A) Bare-SiNPs, (B) Amine-SiNPs, (C) Epoxy-SiNPs and (D) Hydroxy-SiNPs. The x-axis shows the depth of penetration of biofilms, where 0 μm represents the top layer of biofilm and the dashed vertical line (at approximate average thickness of the 14.7 μm) represents the bottom layer of biofilm. The y-axis is the normalized fluorescence intensity of red channel.
the Epoxy-SiNPs surface [24]. The affinity of the carbohydrate component of EPS matrix to SiNPs was also further analyzed through in-vitro studies, using individual monosaccharide and polysaccharide solutions. The contribution of SiNP surface charge became more evident for the binding of sugar molecules to the surface of SiNPs with the highest overall binding shown to be on the surface of Amine-SiNPs.

Colocalization of SiNPs with EPS matrix macromolecules

Differences in the colocalization of SiNPs which have relatively similar surface chemistry (Epoxy- and Hydroxy-SiNPs) or SiNPs with a similar charge (Bare-, Epoxy- and Hydroxy-SiNPs) with each respective EPS macromolecule further highlights the extent to which a comparatively small change in the surface chemistry of the SiNPs can dramatically affect the SiNPs binding affinity. Given the complex and heterogeneous nature of the biofilm matrix, complex physicochemical interactions are expected to govern the affinity of SiNPs to EPS macromolecules; however, we can hypothesize that certain distinct characteristics of both the SiNPs and EPS macromolecules in question may be dominant in changing their affinity and therefore stabilizing their interactions. For example, higher levels of colocalization between Epoxy-SiNPs and NAG-containing polysaccharides may be due to the electrostatic contribution of the epoxy functional groups [24] interacting with the amino functional groups on the partially decylated NAG polysaccharides, with possible covalent and hydrogen bonding interactions occurring (Fig. 10A). The increased colocalization between Epoxy-SiNPs and Mannose-containing polysaccharides may also be stabilized by hydrogen bonding between functional groups. Similarly, the electrostatic contributions of the amino,
of these specific interactions may require additional fluorescent labelling. Void or black areas in CLSM images are not always representative of empty space but rather as areas which have not been fluorescently labelled and may contain other macromolecules. Furthermore, the hydroxyl and carbonyl groups of NAG-containing polysaccharides with the amino groups Amine-SiNPs may be stabilized under the same intermolecular forces in these conditions (Fig. 10B).

While the electrostatic contributions of macromolecules and SiNPs may stabilize and increase the likelihood of physicochemical interactions, these same contributions may lead to repulsion forces between similarly charged entities. For instance, the negative charge on the surface of Bare-SiNPs and the overall negative charge of proteins and eDNA within the matrix may lead to lower colocalization between these binding pairs (Fig. 5). In addition, weak colocalization ratios are not necessarily indicative of a lack of interaction but rather, the discernment of these specific interactions may require additional fluorescent labelling.

The complexity of interactions already occurring between different macromolecules within natural biofilms have been elucidated previously [51], while investigations of inherently complex interactions between functionalized NPs and the biofilm matrix is still in its infancy. In order to optimize and contribute to the development of novel approaches for biofilm eradication and control methods, specifically with the aid of engineered NPs, the understanding of the complex interaction which occur in these systems is vital. This study has demonstrated the successful use of a combination of highly specific fluorescent probes and 3D-CLSM imaging to identify the distribution and topography of various macromolecules (including proteins, polysaccharides and DNA) within P. fluorescens WCS365 biofilms. Building on these results, colocalization and binding analyses revealed various high affinity interactions which may be occurring between functionalized SiNPs and these EPS matrix macromolecules. Although highly specific, the design of unique NP systems which have high selectivity for certain EPS macromolecules have previously been based on a more traditional “lock-and-key” or ligand-protein interactions [52], and often require a great deal of optimization. Alternatively, the use of a colocalization analysis approach applied to “smart” NP design may lead to better assessment of NP-biofilm macromolecule interactions and more effective biofilm control measures. For example, if NAG is to be targeted for enzyme degradation, a NP-enzyme conjugate with an overall positive charge may provide more efficient targeted removal. The contribution of this work can add to the fundamental understanding of NP-biofilm matrix interactions, to enable more versatile and efficient NP design and application and provide new avenues for the development of advanced approaches to the prevention, control and eradication of unwanted biofilms.

**Conclusion**

The complexity of interactions already occurring between different macromolecules within natural biofilms have been elucidated previously [51], while investigations of inherently complex interactions between functionalized NPs and the biofilm matrix is still in its infancy. In order to optimize and contribute to the development of novel approaches for biofilm eradication and control methods, specifically with the aid of engineered NPs, the understanding of the complex interaction which occur in these systems is vital. This study has demonstrated the successful use of a combination of highly specific fluorescent probes and 3D-CLSM imaging to identify the distribution and topography of various macromolecules (including proteins, polysaccharides and DNA) within P. fluorescens WCS365 biofilms. Building on these results, colocalization and binding analyses revealed various high affinity interactions which may be occurring between functionalized SiNPs and these EPS matrix macromolecules. Although highly specific, the design of unique NP systems which have high selectivity for certain EPS macromolecules have previously been based on a more traditional “lock-and-key” or ligand-protein interactions [52], and often require a great deal of optimization. Alternatively, the use of a colocalization analysis approach applied to “smart” NP design may lead to better assessment of NP-biofilm macromolecule interactions and more effective biofilm control measures. For example, if NAG is to be targeted for enzyme degradation, a NP-enzyme conjugate with an overall positive charge may provide more efficient targeted removal. The contribution of this work can add to the fundamental understanding of NP-biofilm matrix interactions, to enable more versatile and efficient NP design and application and provide new avenues for the development of advanced approaches to the prevention, control and eradication of unwanted biofilms.

**Methodology**

**Materials**

The following reagents were all purchased from Merck, Ireland, and used as supplied: Acetic Acid, N-Acetylglucosamine (GlcNac), Ammonium Chloride (NH₄Cl), Ammonium Hydroxide (NH₄OH), (3-Aminopropyl) Triethoxysilane (APTES), Bovine Serum Albumin (BSA) Calcium...
Chloride (CaCl₂), Cyclohexane, Dowex® cation exchange resin, Ethanol (EtOH), D-Galactose(Gal), 3-Glycidoxypropyltrimethoxysilane (GPTMS), D-Glucose (Glc), Glycerol, King B Agar, Magnesium Sulfate (MgSO₄), D-Mannose (Man), Mowiol 4–8, Ribonuclease 1 (RNase), Rhodamine B Isothiocyanate (RITC) and Tetraethyl Orthosilicate (TEOS). Potassium Phosphate Dibasic was purchased from Honeywell, Fluka™, (Ireland). Pierce™ silver staining kit, Sypro® Orange Protein Gel Stain and Sytox® Green were purchased from ThermoFischer Scientific (Illinois, USA). Flurescein (FITC)-labelled Lectin Kit I was purchased from Vector Laboratories (California, USA). Grade 1 pure water (18.2 MΩ) was obtained from an Elga Process Water System (Biopure 15 and Pureflex 2, Veolia, Ireland).

**Bacterial culture and maintenance**

The bacterial strain used in this investigation was *P. fluorescens* WCS365. Bacterial cultures were stored at −80 °C in 25% (w/v) glycerol. For cultivation, thawed aliquots were streaked onto King B agar plates and incubated for 24 h at 30 °C. A single bacterial colony was used to inoculate 50 mL of sterile King B medium in a 250 mL Erlenmeyer flask and incubated at 30 °C with shaking at 200 RPM overnight (16–18 h) to an approximate optical density (OD) of 0.6 at 600 nm of between 2.3 and 2.6. The overnight cultures were then diluted to a final OD₀₆₀₀ of 1.0 using fresh sterile King B medium.

**Biofilm growth**

Biofilms were prepared as per Safari et al. (2014) with minor modifications [57]. A 5 mL volume of the OD₀₆₀₀ of 1 diluted culture, supplemented with CaCl₂ to a final concentration of 1.5 mmol L⁻¹, was added to sterile 50 mL centrifuge tube containing a glass coverslip (24 mm × 50 mm) and plugged with sterile cotton wool. Tubes were then incubated for 72 h at 30 °C with shaking at 100 RPM.

**EPS extraction and protein precipitation**

EPS was extracted from 72 h old biofilms grown on glass coverslips using cation-exchange resin (CER) extraction as per Jachlewski et al. with minor modifications [58]. Briefly, the biofilms were gently washed with PBS buffer and then resuspended in 0.5 mL of a NaCl 0.9% (w/v) solution. After sonication for 2 min, the cation exchange Dowex® resin was added (1%, w/v) and the EPS extracts were shaken at 300 RPM for 2 h at 4 °C in the dark. An EPS solution was obtained after centrifugation (20000 g for 20 min at 4 °C) and filtration (Millipore® 0.22 µm pore membrane) of the supernatant. EPS extracts were maintained at −80 °C until their use.

Proteins from an EPS extract were precipitated based on an ammonium sulfate salting-out method with minor modifications [59]. Briefly, to 1 mL of EPS extract, (NH₄)₂SO₄ was added dropwise over a period of 1 h. An increased turbidity was observed after adding 0.49 g of the salt. The EPS extract was then left at 4 °C overnight for complete precipitation and then centrifuged at 20000 g for 20 min at 4 °C. The supernatant was discarded, and the protein pellet was dispersed in the same volume of H₂O. The EPS protein solution was then kept at 4 °C until its use.

**Sample preparation for CLSM**

Various fluorescent probes (Table 2) were used for the identification and subsequent colocalization analysis of polysaccharides, proteins and eDNA within the biofilms. The polysaccharides present in *P. fluorescens* WCS365 biofilms have not yet been identified. Polysaccharides produced by other *Pseudomonas* include alginate [60], PEL (a partially deacetylated N-acetyl-d-glucosamine (NAG) and N-acetyl-d-galactosamine polysaccharide (NAC)) and PSL (a Mannose, Galactose and L-rhamnose containing polysaccharide) [61]. Therefore, the specific lectins used in the current study were chosen based on their prominence, highly specific binding and also their overall abundance within the biofilms from the results of a fluorescein lectin-binding analysis [22] (data not shown). To this end, fluorescently-labelled (FITC) lectins and various other probes were used to label the individual components within the biofilm independent of other molecules.

**Staining**

Lectin stock solutions were diluted in H₂O to a final concentration of 20 µg mL⁻¹. A 1000X dilution of Sypro® Orange was used for the identification of proteins within the biofilm, while eDNA was stained with a 5000X dilution of Sytox™ Green. Each biofilm-coated glass coverslip was carefully removed from the centrifuge tubes, and gently rinsed three times in H₂O. The coverslip was then placed horizontally on a sample holder and 150 µL of each respective fluorescent probe solution was added directly to the biofilm, followed by incubation...
in the dark for 15 min. After incubation, the biofilms were gently rinsed three times in H2O to remove all unbound probes. 

**Exposure to SiNPs:** Suspensions of SiNPs were prepared by diluting stock solutions to a working concentration of 0.5 mg mL\(^{-1}\) using H2O. The working solutions were sonicated for 1 h prior to use. Subsequent to fluorescent probe staining of each biofilm component, 150 μL of each respective SiNP solution was placed on top of the biofilm as described above. The biofilms were rinsed three times in H2O to remove all unbound SiNPs. Each stained biofilm-coated coverslip was mounted onto a glass microscope slide (25 mm x 75 mm x 1 mm) using Tris-buffered Mowiol 4-88 (pH 8.5) mounting medium as per Pulaz et al. [5] (Fig. 59). The prepared slides were then allowed to dry for at least 1 h at room temperature while protected from light. Horizontal plane z-stack images were acquired with an Olympus FluorView FV1000 CLSM attached to an inverted Olympus IX81 microscope with a 60x/1.35 NA UPL SAPO oil immersion objective (Olympus Optical, Tokyo, Japan). The microscope was equipped with 405 nm, 488 nm, 543 nm and 633 nm laser lines.

**Image acquisition, processing and analysis**

All images were acquired equally; 1x digital zoom, a scanning speed of 2.0 μs/pixel, with 2x Kalman line averaging and sequential channel acquisition. At least three image stacks, with a z-step of 1 μm/pixel radius) prior to all analysis. To quantify the penetration profiles of the SiNPs, fluorescence signals from z-stack CLSM images were used to determine the location and penetration of SiNPs. All colocalization analysis were performed using the “JaCoP” plugin [27] in Fiji image processing software. Mander’s correlation coefficients were used for reporting colocalization between SiNPs and EPS matrix macromolecules and therefore the colocalization ratios were independent of signal intensity and overall abundance of fluorescence signal.

**SiNPs entrapment by the biofilm**

A method adapted from by Nevius et al. [63,71] was used to determine the influence on SiNP surface modification on the entrapment of SiNPs into the biofilm. Briefly, biofilms were cultured in black 96-well plates at 30°C for 24 h with orbital shaking at 125 RPM. After growth, biofilms were rinsed three times in H2O to remove loosely attached cells. SiNP suspensions (0.5 mg mL\(^{-1}\)) were added to each biofilm-containing well as well as control wells and incubated for 15 min. The total relative entrapment of SiNPs by biofilms was calculated by measuring the fluorescent intensity of SiNPs which have not yet attached to biofilms (and therefore present in the supernatant) and comparing this to SiNP controls (wells in which NPs have not been exposed to the biofilm). The overall entrapment of SiNPs can be calculated using the difference between the two fluorescence intensity values (n = 6). The SiNPs fluorescence intensity (λex = 550, λem = 585 nm) was measured using a plate reader (SpectraMax iD3, Molecular devices).

**In-vitro binding interactions**

While in-situ experimental work provides an insight into the bacterial biofilm system, in-vitro analyses may provide an insight into the general mechanisms involved in various other applications. For protein binding affinity analysis, 500 μg mL\(^{-1}\) of a precipitated EPS protein extract, BSA or a RNAse solution was exposed to each SiNP [2 mg mL\(^{-1}\)] for 15 min, similarly as above. BSA and RNAse were chosen as model proteins based on their differing surface chemistry to analyse the effect of overall surface charge on the protein attachment to the NP surface. In order to quantify the protein concentration in the supernatant and therefore the concentration of protein bond to the SiNPs surface, the Lowry method for protein quantification was used [64]. For the analysis of the protein corona formed on each NP after exposure to an EPS protein extract, the total precipitated EPS proteins as well as the protein corona formed on each respective SiNP was analyzed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) according to the method described by Laemmli [65]. Ten microliters of supernatant samples were mixed with 10 μL sample buffer (125 mMol L\(^{-1}\) Tris pH 6.8; 4% SDS; 2% β-mercaptoethanol; 0.25 mol L\(^{-1}\) EDTA; 20% glycerol; 0.04% bromophenol blue), separated by SDS-PAGE (12% w/v) and visualized by silver staining according to the manufacturer’s instructions (Pierce Silver stain kit, Thermo Fischer Scientific).

For carbohydrate binding affinity analysis, Mannose, Galactose, Glucose, NAG and sodium alginate stock solutions of 500 mMol L\(^{-1}\) were prepared in H2O. Based on serial dilutions in H2O, standard curves ranging from 0.03125 to 10 mMol L\(^{-1}\) for Mannose, Galactose, Glucose and Alginate was generated. For NAG, a standard curve ranging from 3.125 to 250 mMol L\(^{-1}\) was prepared. For binding analysis, 250 μL of Mannose [5 mMol L\(^{-1}\)], Galactose [5 mMol L\(^{-1}\)], Glucose [5 mMol L\(^{-1}\)], Alginate [5 mMol L\(^{-1}\)] or NAG [100 mMol L\(^{-1}\)] was added to 250 μL of each SiNPs [2 mg mL\(^{-1}\)] solution, respectively, and incubated at 1000 RPM at 25°C for 15 min. To assess the concentration of carbohydrates bound to the SiNPs, the SiNP-carbohydrate solution was centrifugated at 20000 g for 20 min and three 50 μL aliquots were removed from the supernatant for analysis. A phenol-sulphuric acid total carbohydrate assay was performed on each aliquot and compared to each respective standard curve [66].

**Simulation and visualization of protein surface charge distribution**

The electrostatic surface potentials of BSA and RNAse was calculated using the Adaptive Poisson-Boltzmann Solver (APBS) software [67], after assigning protonation states to residues at a neutral pH, using PROPKA [68]. Charge distributions were visualized using 3Dmol [69] with the program internal threshold for colouring electrostatic potential set to −2.0 kcal mol\(^{-1}\) [2]; blue: +2.0 kcal mol\(^{-1}\) [2]. The protein crystal structures of BSA (protein data bank entry ID 1V03) and RNAse (protein data bank entry ID 5D97) were taken from the RCSB Protein Data Bank (http://www.rcsb.org/) [70].

**Statistical analysis**

All data presented is representative of the mean ± standard deviation of at least three separate experiments, each of which contained at least 3 replicates. A one-way ANOVA followed by Tukey’s HSD pairwise comparison (P < 0.05) was used to assess statistical significance using IBM SPSS Statistics for windows IBM SPSS Statistics for Windows version 26.0. (IBM Corp., Armonk, N.Y., USA). Where qualitative images are
shown, experiments were performed in triplicate, and a representative image is shown.

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Declaring of competing interest

All authors declare they have no conflict of interest.

CRediT authorship contribution statement

Dishon Wayne Hiebner: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Validation, Writing - original draft, Writing - review & editing. Caio Barros: Investigation, Writing - review & editing. Laura Quinn: Writing - review & editing. Stefania Vitale: Formal analysis, Investigation, Validation, Writing - review & editing. Eoin Casey: Conceptualization, Writing - review & editing, Funding acquisition, Resources, Supervision, Project administration.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.biocel.2020.100029.

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