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4	Review
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6	Title: Atomic Force Microscopy Studies of Bioprocess Engineering Surfaces – Imaging,
7	Interactions and Mechanical Properties Mediating Bacterial Adhesion.
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9	Sean A James ¹
10	Nidal Hilal ²
11	Chris J Wright ¹
12	
13 14 15 16	 Biomaterials, Biofouling and Biofilms Engineering Laboratory (B³EL, System and Process Engineering Center, College of Engineering, Swansea University, Fabian Way, Swansea SA1 8EN, UK.
17 18 19	 Centre for Water Advanced Technologies and Environmental Research (CWATER), College of Engineering, Swansea University, Fabian Way, Swansea SA1 8EN, UK
20	Correspondence: Chris J Wright, System and Process Engineering Center, Swansea
21	University Bay Campus, Fabian Way, SA1 8EN, Swansea, Wales.
22	E-mail: C.J.Wright@swansea.ac.uk
23	
24	Keywords: Atomic force microscopy, Bacteria, Biofouling, Force measurement,
25	Nanoindentation.
26	
27	Abbreviations: DVLO, Derjaguin, Landau, Verwey, Overbeek theory; XDVLO, Extended
28	Derjaguin, Landau, Verwey, Overbeek theory; EPS, Extracellular Polymeric Substance; LB-
29	EPS, Loosly Bound Extracellular Polymeric Substance; TB-EPS, Tightly Bound Extracellular
30	Polymeric Substance; SMFS, Single Molecule Force Spectroscopy; SCFS, Single Cell Force
31	Spectroscopy.
32	

1 Abstract

2 The detrimental effect of bacterial biofilms on process engineering surfaces is well documented.

3 Thus, interest in the early stages of bacterial biofilm formation; in particular bacterial adhesion 4 and the production of anti-fouling coatings has grown exponentially as a field. During this time, 5 Atomic force microscopy (AFM) has emerged as a critical tool on the evaluation of bacterial 6 adhesion. Due to its versatility AFM offers not only insight into the topographical landscape and 7 mechanical properties of the engineering surfaces, but elucidates, through direct quantification 8 the topographical and biomechnical properties of the foulants The aim of this review is to 9 collate the current research on bacterial adhesion, both theoretical and practical, and outline 10 how AFM as a technique is uniquely equipped to provide further insight into the nanoscale 11 world at the bioprocess engineering surface. .

2 1 Introduction

3 Bacterial contamination of water based industrial infrastructure is unavoidable, consequently, 4 the formation of bacterial biofilm on process engineering surfaces is a substantial issue for 5 many industrial processes. A bacterial biofilm consists of a heterogeneous consortium of sessile 6 organisms embedded within a gel like support matrix formed from exopolysaccharides, proteins, 7 and extracellular DNA, typically adhered to a solid support structure. The formation of a biofilm 8 offers bacteria considerable advantages over the planktonic state, including a higher 9 concentration of nutrients and aqueous gasses, increased proximity to other cells, and 10 protection from biocidal, chemical and biological attack. Additionally, biofilm formation is known 11 to result in numerous detrimental effects on process engineering infrastructure, including 12 reduction in the efficacy of cooling towers, heat exchangers, ion exchangers and, filtration 13 membranes, as well as causing a substantial level of end product spoilage. The combined fiscal 14 impact of bacterial bioburden is likely to amount to many millions, if not billions of pounds of lost 15 income due to reductions in process efficiency and repairs. Biofilms are of particular concern as 16 the innate resistance offered by the structure often result in removal being almost impossible, 17 while simultaneously facilitating the re-contamination of any downstream infrastructure that may 18 have been successfully cleaned.

19

20 Several control strategies are often employed in an effort to control bacterial biofouling, typically 21 this will involve the use of toxic chemical such as sodium hypochlorite, glutaraldehyde, 22 isothiozoline, and chlorine dosing. However, these strategies are non-specific, damaging to 23 infrastructure and often ineffective in the removal of fully established biofilms. Furthermore, 24 tightening of legislation on the use of antimicrobials has further limited their efficacy, therefore 25 highlighting the need for the implementation of new strategies. As bacterial adhesion marks 26 the initiation of biofilm formation the creation novel anti-adhesion compounds and coatings 27 offers promising solutions to the biofouling problem. Typically, these compounds and coatings 28 alter surface chemistry therefore modifying the strength of binding forces facilitating bacterial 29 adhesion. Hence furthering our understanding of the fundamental bacterial substrate 30 interactions that promote primary adhesion is essential, as the application of such knowledge is 31 paramount in the formation and design of these strategies (1-3). Despite this, development of 32 such technologies is limited primarily due to the lack of understanding of the forces governing 33 theses interactions. Theoretical models predicting the interactions between bacteria and 34 substrates are well established, with comprehensive modeling of the van der Waals, 35 electrostatic and Lewis acid-base interactions described in the extended DVLO (XDVLO) theory 36 (4). Despite the comprehensive nature of this model, discrepancies in the predictions when 37 compared to experimentally derived data exposes a number of substantial flaws (5-7), 38 particularly in the description of the interaction between biological moieties.

1 Atomic force microscopy (AFM) offer an answer to this issue allowing unique insights into the 2 interaction processes of biological systems. Application of traditional AFM techniques allow for 3 the unparalleled visualization and, characterization of the substrate topography at both the 4 micro and nanometer scale. This has been applied to the characterization of bioprocess 5 engineering surfaces such as stainless steel to measure surface roughness of different finishes 6 pore size determination of separation membranes and study of cellular surfaces (8-10) AFM is 7 not just an imaging device it also has a force measurement capability that has provided novel 8 insight into the interactions of bioprocess engineering surfaces this includes the direct 9 quantification of forces at the microbial surface and nano-mechanics allowing elucidation of the 10 interplay of all of these factors under in situ conditions (11,12).

11

12 Through the course of this review the authors hopes to highlight the versatility of AFM.
13 Summarize the landmark research that guided the application of the technology in the
14 characterization of substrates, bacterial cells, and the adhesion forces and how these studies,
15 build upon current microbial adhesion models that impact the characterization and optimization
16 of bioprocess engineering surfaces.

17

18 **2 Basic Principles**

19 AFM is a form of high resolution microscopy from the family of scanning probe microscopies 20 (SPM) developed in the mid-1980s the components of which are detailed in figure 1 (13). AFMs 21 defining features are its resolution, demonstrated to be in the order of fractions of a nanometer, 22 and versatility as a force sensor. AFM operates two major modes; imaging and force 23 spectroscopy, the former can be further divided into two major categories of imaging; contact 24 and intermittent-contact, the basic principles employed during operation remain the same 25 regardless of application. Other AFM imaging techniques have been developed that map 26 properties such as conductance and friction across a surface, however it is contact and 27 intermittent contact mode that have mainly been applied to the study of process engineering 28 surfaces.





Figure 1 Schematic representation of the atomic force microscope

5 During measurement AFM records the nanoscale attractive and repulsive forces of the sample 6 material by the deflection of a sharp tipped cantilever, which is systematically scanned across 7 the surface. These forces are measured via the displacement of a laser reflected off the upper 8 gold-coated side of the cantilever onto a quadratic photosensitive photodiode (PSPD). In 9 contact mode, the AFM cantilever is brought into intimate contact with the sample material, the 10 material is then rastered beneath the cantilever and axial and longitudinal deflections in the 11 cantilever recorded via the photodiode. Intimate contact with the sample material is essential, 12 and maintained through the implementation of a DC feedback amplifier controlling a 13 piezoelectric motor to maintain a set level of deflection in the cantilever. This ensures that the 14 force applied to the sample is constant and controlled to prevent sample damage. The forces in 15 the bent cantilever maintain the imaging tip in direct contact with the surface. The destructive 16 forces applied to the sample as a result of imaging can be mitigated through imaging in a liquid 17 media and/or changing to intermittent-contact imaging. Additionally imaging in liquid allows for 18 the removal of the strong attractive forces (capillary forces) on the AFM tip and cantilever 19 caused by the adsorbed water layers at the surface of the sample further reducing the forces 20 applied to the sample due to the tip. Imaging in a liquid media has the additional advantage of 21 allowing for the characterization of sample materials such as membranes and biological 22 contaminant under process- relevant conditions. However, due to the mobility of some bacterial 23 species and the generally weak forces binding the cells to the surface, immobilization of the 24 cells is often required to prevent destruction of the sample (14-16). Comprehensive review of 25 the methods employed in the immobilization of bacterial cells can be found here (17). However, 26 for the purposes of this review a brief summery shall be supplied. Typically immobilization takes 27 one of two form; mechanical, or chemical. Mechanical immobilization traps the cells within an 28 inert matrix for example, gelatin, agar or membranes (14,18–20). This methodology has been

1 refined to include the use of functionalized surfaces such as lithographically patterned silica and 2 polydimethylsiloxane (PDMS) (15,21-24). Chemical immobilization typically make use of 3 surface chemistry to bind the cell to substrate. The target for immobilization varies and includes 4 surface charge in the case of poly-L-lysine to the crosslinking of carboxyl groups (25-30). While 5 chemical techniques typically result in a high level of immobilization and specificity in orientation 6 care should be taken in selecting an appropriate technique. Chemical fixation, through its very 7 nature alters the surface chemistry of the cell and therefore alters not only the surface 8 properties but also the viability of the cell. Hence, it is critical that an appropriate methodology is 9 selected that takes into consideration the goals of the experimentation.

10 When imaging of soft samples such as biological samples or filtration membranes is required 11 application of contact mode imaging may not be suitable due to the intimate contact of the sharp 12 imaging tip with the surface, in this case intermittent contact or tapping mode is implemented. 13 During tapping mode, the cantilever is driven into oscillation at a frequency close to that of the 14 cantilevers resonance. The oscillation of the cantilever is then monitored as the sample material 15 is scanned with any alterations in the cantilevers oscillation corrected through the 16 implementation of a direct feedback loop, causing the movement of a piezo-scanner, which is 17 used to produce the image. The change in the oscillation as a function of the movement of the 18 piezo-scanner is then recorded as a topography image. Simultaneously, alteration in the phase 19 angle of the oscillations at the tip of the cantilever and the free oscillation of the cantilever can 20 be recorded. This allows for the characterization of the phase angle across the surface; termed 21 the phase contrast. Phase contrast mapping allows for a qualitative analyses of the differences 22 in the mechanical properties across a surface of heterogeneous material. Tapping mode offers 23 a number of advantages in the imaging of softer materials, most predominantly tapping mode 24 minimizes the destructive lateral forces of the technique on the sample material. 25



Figure 2. AFM images of bioprocess engineering surfaces (A) Cyclopore microfiltration membrane $(3.2\mu m^2)$ (B) Humic acid layer fouling an ultrafiltration membrane $(3.5\mu m^2)$ (C) Air bubbles at a membrane surface $(0.75\mu m^2)$ (D) Stainless steel process plant surface. $(10\mu m^2)$.(E) Lawn of Saccharomyces cerevisiae cells (brewing yeast NCYC 1681) showing budding scars $(15 \mu m^2)$ (F) Shewanella oneidensis cell showing protein clusters at surface formed during anaerobic respiration and microbial fuel cell operation $(1.2\mu m^2)$

1 Figure 2 shows a number of example AFM images captured in our laboratory outlining the 2 application of AFM in bioprocess engineering. The first three images are at membrane surfaces 3 showing features important to process performance such as pore size and distribution (Figure 2 4 (A)) Figure 2 (B) shows an ultrafiltration membrane purposely fouled with humic acids; a 5 common foulant of process engineering surfaces consisting of the biproducts of organic matter 6 biodegradation, Figure 2 (C) details bubbles at a membrane surface, which formed under 7 certain operating conditions and are reported to participate in biofouling reduction (31). Finally 8 figure 2 (D) details surface defect in a stainless steel surface as a result of surface polishing. 9 These first four images highlight how AFM is capable of characterisaing a number of surface 10 artifacts associated with fouling. Figure 2 also details AFM images that suggest molecular 11 features of the microbial cell wall, in this particular case protein clusters involved in electron 12 transport under anaerobic conditions (Figure 2 (E)); and yeast budding scars (Figure 2 (F)).

13



Figure 3. AFM Force spectroscopy of a single stationary phase *S. cerevisiae* cell probe against freshly cleaved mica in 0.01M NaCl pH5

14

15

Application of AFM is not limited to the characterization of topographical landscapes, in fact AFMs greatest asset lies in its ability to directly quantify the nanomechanical properties of a material. This is achieved through the generation of force-displacement curves; a graphical representation of the deflection of the cantilever as a function of the tip-sample separation distance as the tip is brought into contact with the sample. Figure 3 shows a typical forcedisplacement curve, in this case a single *S.cerevisiae* cell probe against(stationary phase) freshly cleaved mica. The force-displacement curve is treated as two curves; the approach

1 curve (E,D,F,G) and the retraction curve (A-D). At position E the sample and probe are in 2 complete isolation this continues until position F wherein long range electrostatic forces begin to 3 repel the probe. The probe is then pressed further towards the surface by the extension of the 4 piezo scanner, through the electrostatic repulsion experienced at F until contact with the sample 5 surface is achieved. At this point a number of nano-mechanial properties can be defined, such 6 as elastic and plastic deformation regions and the yield point. The bending of the cantilever 7 continues at the constant, defined approach speed until a predefined loading force is 8 experienced (G). Point A defines the start of the retraction curve, the probe is retracted from the 9 surface and any compression in the probe or surface is present to point B. Point C represents 10 the separation of the probe from the sample material, in this case the probe exhibits some 11 adhesion with the substrate as seen through the deflection of the cantilever in the opposite 12 direction culminating in the snapping of the attractive forces at the apex wherein the cantilever 13 returns to a state of zero deflection (E). This characteristic behavior allows for the generation of 14 a number of key variables, for example, at point C the maximum adhesive force can be defined 15 as the maximum deflection of the cantilever. Similarly, the energy of adhesion can also be 16 defined as the area between the approach and retract curves. With regards to biological 17 samples, this area is often large and not as "clean" as non-biological samples. This is for a 18 variety of reasons, including multiple fracture points caused by the fracture of different 19 macromolecules; discussed in depth later in this review, and the tendency for elastic 20 deformation of biological samples. The nanomechanical properties of the substrate can be 21 derived from the contact regions (F-G and A-B) and the adhesive properties from region B-D, 22 however it is worth noting that these variables are dependent on the loading rate of the 23 measurement.

24

Comparison of force-deflection curves collected from soft biological samples to those taken at samples with significantly higher mechanical properties such as silica can allow for the characterization of nanomechanical properties, such as Young's modulus and turgor pressure. This is achieved by processing of the curves through a Herztian model, which describes the elastic deformation of two homogenous bodies touching under load, assuming a parabolic indenter and that the cantilever is of significantly greater thickness than that of the indentation depth, the force can be is defined by the equation;

32
$$F(\delta)_{parabolic} = \frac{4E\sqrt{R}}{3(1-v^2)}\delta^{3/2}$$

33 where R is the radius of the indenter, δ is the indentation depth, E the Young's modulus and v 34 the poisson ratio. Assuming the prerequisites are met the definition of the indentation depth as a 35 function of the applied force is then possible. However, in reality the first assumption; that the 36 both bodies are homogenous, is often incorrect and in fact the substrate is heterogeneous in 37 nature. Hence, multiple force curves at multiple surface locations this should be considered when analyzing a homogenous substrate.. Furthermore, the act of nanoindentation is by its very definition a destructive process and as a result consecutive indentation of the same geographical location may result in variation.

4 It is worth noting that with the interpretation of force curves determination of the contact point is 5 often necessary. This is particularly challenging for biological samples for several reasons. 6 Firstly, there is the definition of contact, wherein contact of any description defines the zero 7 point. This would be easily discernable if both objects were essentially smooth on the given 8 scale of measurement; in this case the nanometer scale. However, this is far from the case and 9 in fact both topographies, especially the biological one are likely to respectively rough due to the 10 presence of surface macromolecules to name one obstruction. Methods for the calculation of 11 the precise contact point have been developed, wherein scattered laser light shone between the 12 colloid and the sample is detected using a near-field scanning optical microscope or inverted 13 light microscope however they are far from simply employed. Secondly, on approaching the 14 sample if the attractive forces exceed the spring constant of the cantilever then the cantilever 15 will snap into contact with the sample, and then further drawn into the sample by the attractive 16 forces. This results in two issues, firstly the contact point is almost impossible to define and 17 secondly it is then necessary to assume that no indentation of the sample has occurred. While 18 this is easily mitigated through selection of a cantilever with an appropriate spring constant it 19 does limit resolution.

20

21 **3 Bacterial Adhesion and Biofouling**

22 **3.1 Mechanisms and Theoretical Models**

23 Surface adhesion is a multistage event, consisting of initial or reversible adhesion followed by 24 permanent or irreversible adhesion. Reversible adhesion defines the most crucial step in 25 adsorption of the cell to the substrate, involving the simple, non-specific interaction of a number 26 of fundamental physical effects including; van der Waals forces, Brownian motion, electrostatic, 27 hydrophobic and acid-base interactions. While irreversible adhesion is more complex as it is the 28 product of the characteristics of the cell, and predominately mediated by polymeric structures on 29 the bacterial surface such as pili, flagella, adhesins and capsule components (slime layer, 30 glycocalyx).

31

32 Initially, attempts to model bacterial adhesion focused on the use simple colloidal models such 33 as Derjaguin-Landau-Verwey-Overbeek (DVLO) (16,32,33). However experimental results have 34 typically been shown to correlate poorly to predictions made by this method, as the DVLO 35 theory only considers electrostatic and van der Waals forces (5). This may be a result of the 36 inherent assumptions of DVLO theory, which suggests that pH and ionic strength are the 37 defining features of bacterial adhesion in solution; while ignoring the effect of hydrophobicity, 38 or, the incorrect estimation of key variables (6,7). The result of these discrepancies resulted in 39 the combination of the DLVO theory and surface thermodynamics. While this remediated some

1 of the discrepancies of the DVLO theory a number of other phenomenon were still unaccounted 2 for, predominantly as a result of the assumptions of thermodynamic theory in which the reaction 3 is assumed to be reversible, and that a new intimate bacterium substrate interface is formed 4 (34-37). van Oss further extended the DVLO theory to include hydrophobic interactions (4). 5 Extended DVLO theory (XDVLO) as it became known, expressed the total adhesion energy of a 6 spherical object; the cell, against a semi-infinite plane; the substrate at a discrete distance to be 7 a result of the sum of the van der Waals, electrostatic and lewis acid-base interactions and is 8 defined by the equation:

9

$$\Delta G^{TOT}(d) = \Delta G^{LW}(d) + \Delta G^{EL}(d) + \Delta G^{AB}(d)$$

10 wherein $\Delta G^{LW}(d)$, $\Delta G^{EL}(d)$ and $\Delta G^{AB}(d)$ are the free energies of the Lischitz-van der Waals, 11 electrostatic and Lewis acid-base interactions respectively at a given distance. At a discrete 12 distance d, $\Delta G^{LW}(d)$, $\Delta G^{EL}(d)$ and $\Delta G^{AB}(d)$ can be defined as:

13
$$\Delta G^{LW}(d) = -\frac{A}{6} \left[\frac{a}{d} + \frac{a}{d+2a} + \ln\left(\frac{d}{d+2a}\right) \right]$$

14
$$\Delta G^{EL}(d) = \pi \varepsilon a(\zeta_1^2 + \zeta_2^2) \left[\frac{2\zeta_1 \zeta_2}{\zeta_1^2 + \zeta_2^2} ln \frac{1 + exp(-\kappa d)}{1 + exp(-\kappa d)} + ln\{1 - exp(-2\kappa d)\} \right]$$

15
$$\Delta G^{AB}(d) = 2\pi a \lambda \Delta G^{AB}_{adh} \exp\left[\frac{(d_0 - d)}{\lambda}\right]$$

16 wherein *a* is the radius of the sphere, *A* is the Haymaker constant and defined as:

 $A = -12\pi d_0^2 \Delta G_{adh}^{LW}$

18 ζ is the zeta potential, ε is the permittivity of the medium K is the electric double layer thickness, 19 λ is the correlation length if molecules in a liquid medium and d_0 is the closest separation 20 distance between the sphere and plane. Finally, ΔG_{adh}^{LW} and ΔG_{adh}^{AB} are the free energies of 21 adhesion for the Lischitz-van der Waals, and Lewis acid-base interactions as defined by the 22 LW-AB approach. Through the implementation of this model a series of separation distance 23 against free energy graphs can be created theorizing the interactions between the cell and the 24 substrate. Through interpretation of these graphs maximum and minimum energy requirements 25 can be deduced for a given separation distance and hence favorability of the adhesion event. 26 However, while the theory proves to be a powerful tool in the modeling of the initial stages of 27 bacterial adhesion the model still fails to take into consideration the biological aspect of the 28 interaction such as the effects due to pilli, flagella and fails to account for non-spherical cell 29 shapes.. AFM offers a unique opportunity to remedy these issues. Firstly, through the 30 application of force spectroscopy; to be discussed in the section 4, AFM allows for the direct 31 quantification and comparison of experimentally derived approach curves and the theoretical 32 separation distance against free energy curves predicted by these models. Secondly, through 33 the characterization of the folding/unfolding pathways of surface macromolecules and the use of 34 non-spherical cell probes further refinement of the model through the implementation of a fourth 35 biological factor in the XDVLO theory may be possible. Consequently, the implementation of a 1 more rigorous theoretical framework will be possible for predicting and preventing biofouling at

- 2 bioprocess engineering surfaces
- 3

4 **3.2 Substratum Characteristics**

5 3.2.1 Surface Roughness

6 The effect of substratum characteristics on the adhesion of bacteria has been a point of interest 7 ever since the discovery of the detrimental effects of biofilms, with both surface roughness and 8 nanomechanical properties being implicated with increasing retention of bacteria [31]. However, 9 due to its complex nature elucidating the complete mechanism of bacterial attachment and 10 biofouling of bioprocess surfaces remains a challenge. As shown by the XDVLO theory the 11 physiochemical interactions are understood to a degree although not comprehensively; the 12 impact of surface topography on the adhesive characteristics remains less well defined. To 13 begin to elucidate the mechanism by which bacterial adhesion is influenced by surface 14 roughness a method for quantifying surface roughness is required. A number of techniques are 15 available to researchers to accomplish this such as stylus and optical type profilometers and 16 dynamic contact angle. These techniques offer a number of advances such as being relatively 17 simple to implement and not particularly resource intensive, but their resolution is limited to the 18 microscale and offer no further insight into the substrate characteristics. As a result, more 19 researchers are applying AFM to the characterization of such surfaces. AFM offers a number of 20 key features not achievable through other means, most notably its ability to resolve the surface 21 roughness at the nanometer scale: a scale that has grown in interest over the last few years 22 (38,39).

23 Several studies have been completed detailing the effect of surface roughness on bacterial 24 adhesion (10-18). The consensus of the aforementioned studies being that on the micro-scale, 25 bacterial adhesion is at its peak when the arithmetic roughness (Ra) of the surface nears the 26 diameter of the cell. This correlation between surface adhesion and surface roughness has 27 been attributed to a number of factors, including the maximization of surface contact area, the 28 accumulation of organic contaminants and protection from sheer stresses (40-42). 29 Consequently, modification of substrate Ra has become a primary focus in the formation of 30 antifouling strategies.

31 Interest in minimizing the surface roughness of substrate materials has resulted in a growth in 32 the characterization and application of nanoscale topographies (43–47). However, nanoscale 33 interactions have proved to be more complex with a number of fundamental contradictions 34 arising (48-54). For example, a significant increase in the adhesion of Escherichia coli and 35 Pseudomonas aeruginosa on nanorough thin film titanium substrates was observed in 36 comparison to a flat reference samples(46). The authors attribute this to the physical stimulus of 37 the nanotextured surface. However, the authors proceed to describe the absence of flagella on 38 P. aeruginosa cells bound to same surface, suggesting changes in cell surface characteristics. 39 Conversely, in direct opposition to the previously mentioned study Ivanova et al. reported that on comparable titanium surfaces adhesion of *P aeruginosa* and *S. aereus* were inversely proportional to the roughness of the surface (43,47). However it is worth noting that a number of the studies performed on titanium contain quantities of TiO₂, a known antibacterial. It is therefore difficult to distinguish alterations in adhesion caused by the nanoscale roughness of

5 the surface as opposed to those caused by the antimicrobial activity.

In a more recent study the effect of nanotapography and grain size, on the adhesion of *E. coli*, *P. aeruginosa, S. epidermidus* and *S. aureus* was observed on shot peened SS316L stainless steel (55). During the study the authors suggest that increases in the nanoscale roughness of material notably increased the adhesion rate of both *S. aureus* and *S. epidermidus* while having no significant effect on the adhesion rate of the two Gram negative species. The research group were additionally able to conclude that the refinement of the grain size, as a result of shot peening did not significantly alter adhesion rates.

13

14 **3.2.3 Conditioning Layers**

15 Adsorption of organic matter to surfaces in aqueous environments and the corresponding 16 impact on bacterial adhesion rates has been well documented (56-62). A major contributing 17 factor to the formation of an organic conditioning layer is the deposition of extracellular 18 polymeric substance (EPS). EPS is not a defined mixture but rather the collective name given to 19 a number of soluble macromolecules produced by bacteria. EPS has been demonstrated to 20 consist of a temporally dynamic mix of polysaccharides, DNA, lipids, proteins and humic 21 substances (63-66). EPS can be loosely grouped into two forms when viewed with regards to 22 planktonic cells, loosely bound (LB-EPS) and tightly bound (TB-EPS), both varying in 23 composition and the former being the primary form of EPS conditioning layers and composed 24 primarily of bacterial proteins and the latter bacterial capsids (67,68). While the formation of 25 organic conditioning layers effects all surfaces within a process engineering environment, by far 26 the most researched area is in the organic fouling of membranes (69–71).

Application of membrane separation technologies has become a substantial part of a number of industrial processes including membrane bioreactors, desalination plants and food processing. However, with all sizes of membrane; microfiltration, ultrafiltration, nanofiltration and reverse osmosis (RO), biofouling is of particular concern as colonization of the membrane will typically result in reductions in membrane separation efficiency and eventually biodeteriation of the separation membrane surface and surrounding infrastructure (72).

In a recent study, Suwarno *et al.*, characterized the effects of a number of conditioning layer contaminants on bacterial adhesion of RO membranes (73). During the study, the researchers observed no trend in organic fouling and surface roughness with an increase in surface roughness of 19nm when the membrane was fouled with alginic acid, this was compared to a 21nm and 15nm decrease in surface roughness for a bovine serum albumin fouled membrane and membrane bioreactor permeate fouled membrane, respectively. Suwarno *et al.*, continued to use AFM to characterize the adhesive properties of the membranes, during the force

1 measurement study it was found that the adhesive force exhibited at the membrane surface 2 was significantly increased from the baseline when exposed to the model foulants; alginic acid 3 and bovine serum albumin [55]. However, the change in adhesion was less pronounced with 4 contamination of membrane bioreactor permeate; an increase of 110nN was observed. The 5 group attributed this to be a result of non-uniform distribution of the foulant material. The 6 research identified an increase in bacterial adhesion as a result of the biopolymer fouling. 7 Bacterial adhesion to the membrane bioreactor permeate fouled membranes was found to be in 8 the order of four times higher than that of the virgin membranes indicating that these changes in 9 surface roughness and adhesion correlate to an increase in bacterial retention.

10 **3.3 Force Spectroscopy**

11 Implementation of AFM for the characterization of bacterial nano-mechanics has been widely 12 accepted as an essential tool in the microbiological community. Eager adoption of AFM by the 13 community is the result of the ability of AFM to resolve the nano-mechanical properties of cells 14 on all levels, from single molecules to consortia of multiple cells such as biofilms in the nano 15 and picoNewton range (74). Understanding the nano-mechanics of bacterial cells with regards 16 to the cellular capsid, elastic modulus and turgor pressure is of particular importance in the 17 further refinement of the understanding of reversible attachment. While, characterization of 18 membrane bound polyproteins, adhesins and cellular organelles will further understanding of 19 the mechanisms of irreversible attachment.

20

21 **3.3.1 Single Molecule Force Spectroscopy (SMFS)**

Surface macromolecules are essential for the ubiquitous successe of microbes within the environment, mediating a number of physiological processes including adhesion. Chemical functionalization of the cantilever has allowed AFM to be applied in the characterisation of a number of surface bound receptors important in adhesion under *in situ* conditions including lipopolysaccharides, pilli and adhesins (75–82). The study of surface macromolecules has facilitated the rapid development of SMFS techniques, and ensured a comprehensive understanding of the fundamental mechanics governing tip-molecules interactions (83–85).

29 Microbial surface proteins can be generally broken down into two groups: functional surface 30 proteins such as adhesins and surface bound long chain macromolecules such as pili. AFM 31 characterisation of long chain surface macromolecules is complex. However, through the use of 32 established models such as the Worm Like Chain (WLC) and Freely Jointed Chain (FJC) 33 models a wealth of information on adhesion can be ascertained. Use of the WLC and FJC 34 models allow for the definition of the entropic elasticity of the molecules and as the result the 35 contour length (L_0) (86,87). If the structure of the target macromolecule is unknown, definition of 36 L_o allows for the confirmation of the experimentally derived data through the use of a normal 37 (Gaussian) distribution, while simultaneously partially elucidating the unbinding pathway 38 (88,89). This method for the definition of L_0 was further expanded through the work of Farrance 39 et al., wherein an idealized theoretical tethered surfaces and the probability of two such

1 surfaces contacting is used to predict the distribution of experimentally derived data (90). In a

2 recent example, SMFS was applied to demonstrate the role of *P. aeruginosa* type IV pili (91).

3 During the study the group untilised a *pilT* mutant to demonstrate low-affinity, high-avidity model

- 4 for type IV pili adhesion through the application of the WLC model. The group concluded that
- 5 each subunit of PilA contained a adhesintope capable of binding the surface allowing adhesion
- 6 along the full length of the molecule.
- 7

8 **3.3.2 Single Cell Force Spectroscopy**

9 Pioneered through the construction of a S. cerevisiae cell probes by Bowen et al., single-cell 10 force spectroscopy (SCFM) has become a fundamental technique in adhesive force 11 characterization (92). This technique is of particular importance in the field of bacterial adhesion 12 as it allows for the simultaneous quantification of all factors governing adhesion under in situ 13 conditions, therefore reducing error as the result of the interpretation of discrete elements. 14 Selection of an appropriate immobilization technique is imperative in the construction of an AFM 15 cell probe (Figure 4). As a result a number of techniques have been utilized to functionalize 16 cantilevers including chemical fixation, electrostatic and wet adhesives (93-100). With the 17 optimum method allowing for the immobilization of the target cell with minimal effect of the 18 technique of the viability, and physiochemical properties of the immobilized cell. Further 19 consideration needs to be taken in to the size of the target cell, with larger cells; such as S 20 cerivisae (~10µm) being easily immobilized directly to the cantilever surface. While 21 immobilization of smaller cells and spores; such as bacteria (~1µm) require further steps. In one 22 such method to immobilize a single bacterial cell, a colloidal sphere was immobilized onto the 23 tip a AFM cantilever. This colloid was then further functionalized with polydopamine to facilitate 24 the immobilization of a single bacterium through AFM micromanipulation (101). This technique 25 offered a number of significant advantages over comparable techniques for the immobilization 26 of single cells. Firstly, through the use of confocal microscopy the technique was shown to 27 immobilize bacterial cells with minimal effect of the target cells membrane integrity, and as a 28 result the viability of that cell. Additionally, the technique allows for the immobilization of the 29 target cell in a highly organized manner further increasing the repeatability of the 30 measurements. Finally the technique was found to be applicable to a number of target species 31 (99).

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Figure 4. SEM image of a cell probe A) *Saccharomyces cerevisiae* B) *Aspergillus niger*

4 5

6 In a recent study, SCFS was applied to characterize the effect of antifouling polymer brushes on 7 the adhesion of Yersinia pseudotuberculosis (102). During the study the group compared the 8 adhesive forces, energy of adhesions and the rupture distance of the brushes against 9 polytetrafluoroethylene (PTFE), polystyrene and glass controls. It was found that the adhesive 10 forces exhibited by the bacteria where sustainably reduced, with a 22% reduction observed for 11 the oligo(ethyleneglycol) methyl ether methacrylate (MeOEGMA) and oligo(ethyleneglycol) 12 methacrylate (HOEGMA) brushes and no less than a 95% reduction for the N-(2-hydroxypropyl) 13 methacrylamide (poly(HPMA)), (3-acryloylamino-propyl)- (2-carboxyethyl)-dimethyl-ammonium 14 (CBAA), [2-(methacryloyloxy)ethyl]- dimethyl-(3-sulfopropyl)ammonium hydroxide (SBMA), 2-15 methacryloyloxyethyl phosphorylcholine (PCMA), and, Poly(2-hydroxyethyl methacrylate) (poly-16 (HEMA)) brushes. Similarly the energy of adhesion showed a significant decrease from 450, 17 550 and 7000aJ for the glass, PTFE and PS respectively to 100aJ for the oligo(ethylene glycol) 18 methactylates and 10aJ for the polyzwitterioic and poly(HPMA) brushes. Further analyses of the 19 force-distance curves highlighted a decrease in the number of rupture events with a 20 corresponding increase in the event distance. A further comprehensive study conducted by 21 Aguayo et al., investigated the impact of nanopatterned polycarbonate on S. aureus adhesion 22 (103). During the study the group observed an increase in both the adhesive force and energy 23 of adhesion with increasing contact time and nanopatterning. Interestingly, the number and 24 location of rupture events exhibited remained relatively constant, the researchers suggested 25 that the surface and/or capsular receptors were involved in adhesion to both substrates

26

27 4 Conclusions

28 In conclusion, AFM has become an essential tool for the study of bioprocess engineering 29 surfaces. This is exemplified by the comprehensive insight it has provided into the nanoscale 30 forces involved in bacterial adhesion and their mechanical properties. Through the 31 characterization of nanoscale surface topography AFM has allowed for the elucidation of the 32 effect of surface roughness on bacterial adhesion rates adding further experimental evidence to 33 the predictions of adhesion models and in turn informing potential anti-fouling strategies. 34 Studies into microbial conditioning have demonstrated the heterogeneity of the conditioning 35 layer while demonstrating the importance of this layer in bacterial adhesion and alluding to 36 potential methods to prevent, remove or mitigate the effect. Conversely AFM has been used to 37 demonstrate the efficacy of anti-fouling layers. Through the functionalisation of probes attached 38 to AFM cantilevers, SMFS has revolutionised our understanding of the complex interplay 39 between microbial surface macromolecules under in situ conditions providing valuable insight

1 into new strategies of preventing the initiation of irreversible attachment. SCFM has now 2 become a mainstay in microbial applications of AFM, and the study of microbial colonisation of 3 bioprocess engineering surfaces. SCFM studies has demonstrated the efficacy of a number of 4 antifouling strategies while facilitating whole cell studies to further future modelling. AFM is not 5 without its limitations and most applications of the technology still require the immobilisation of 6 the cells prior to measurement, potentially influencing the validity of the measurement. Despite 7 this AFM is still an essential addition to the family of instruments used in the characterization of 8 bioprocess engineering surfaces and bacteria. The technology promises further analytical 9 capabilities for the study of microbial systems with the only limitation being the imagination 10 needed to invent more ingenious functionalised cantilever probes. 11

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13 Conflict of interest

- 14 The authors declare no financial or commercial conflict of interest.
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1 5 References

2	1.	Valle J, Burgui S, Langheinrich D, Gil C, Solano C, Toledo-Arana A, et al. Evaluation of
3		Surface Microtopography Engineered by Direct Laser Interference for Bacterial Anti-
4		Biofouling. Macromol Biosci [Internet]. 2015 Aug;15(8):1060–9. Available from:
5		http://doi.wiley.com/10.1002/mabi.201500107
6	2.	Chan WF, Marand E, Martin SM. Novel zwitterion functionalized carbon nanotube
7		nanocomposite membranes for improved RO performance and surface anti-biofouling
8		resistance. J Memb Sci. 2016;509:125–37.
9	3.	Pechook S, Sudakov K, Polishchuk I, Ostrova I, Zakin V, Pokroy B, et al. Bioinspired
10		Passive Anti-biofouling Surfaces Preventing Biofilm Formation. J Mater Chem B.
11		2014;4(8):1166–9.
12	4.	Van Oss CJ. Energetics of Cell-Cell and Cell -Biopolymer Interactions. Cell Biophys.
13		1989;14:1–16.
14	5.	Poortinga AT, Bos R, Norde W, Busscher HJ. Electric double layer interactions in
15		bacterial adhesion to surfaces. Surface Science Reports. 2002. 1-32 p.
16	6.	Dan N. The effect of charge regulation on cell adhesion to substrates: Salt-induced
17		repulsion. Colloids Surfaces B Biointerfaces. 2003;27(1):41-7.
18	7.	Duval JFL, Busscher HJ, Van De Belt-Gritter B, van der Mei HC, Norde W. Analysis of
19		the Interfacial Properties of Fibrillated and Nonfibrillated Oral Streptococcal Strains from
20		Electrophoretic Mobility and Titration Measurements: Evidence for the Shortcomings of
21		the "Classical Soft-Particle Approach." Langmuir [Internet]. 2005 Nov;21(24):11268-82.
22		Available from: http://pubs.acs.org/doi/abs/10.1021/la051735q
23	8.	Bowen WR, Lovitt RW, Wright CJ. Atomic force microscope studies of stainless steel :
24		Surface morphology and colloidal particle adhesion. 2001;6:623–9.
25	9.	Francis LW, Lewis PD, Gonzalez D, Ryder T a, Webb G, Joels L a, et al. Progesterone
26		induces nano-scale molecular modifications on endometrial epithelial cell surfaces. Biol
27		Cell [Internet]. 2009;101(8):481–93. Available from:
28		http://www.ncbi.nlm.nih.gov/pubmed/19236310
29	10.	Johnson D, Hilal N. Characterisation and quantification of membrane surface properties

1		using atomic force microscopy: A comprehensive review. Desalination. 2015. p. 149-64.
2	11.	Powell LC, Pritchard MF, Emanuel C, Onsøyen E, Rye PD, Wright CJ, et al. A
3		nanoscale characterization of the interaction of a novel alginate oligomer with the cell
4		surface and motility of Pseudomonas aeruginosa. Am J Respir Cell Mol Biol.
5		2014;50(3):483–92.
6	12.	Bowen WR, Lovitt RW, Wright CJ. Atomic Force Microscopy Study of the Adhesion of
7		Saccharomyces cerevisiae. J Colloid Interface Sci [Internet]. 2001;237(1):54-61.
8		Available from: http://www.ncbi.nlm.nih.gov/pubmed/11334514
9	13.	Binnig G, Quate C, Gerber C. Atomic Force Microscope [Internet]. Physical Review
10		Letters. 1986. p. 930–3. Available from:
11		http://link.aps.org/doi/10.1103/PhysRevLett.56.930%5Cnpapers2://publication/uuid/551F
12		6513-EAC6-4BC0-A144-AB381B706D40
13	14.	Touhami A, Jericho MH, Beveridge TJ. Atomic force microscopy of cell growth and
14		division in Staphylococcus aureus. J Bacteriol. 2004;186(11):3286–95.
15	15.	Kailas L, Ratcliffe EC, Hayhurst EJ, Walker MG, Foster SJ, Hobbs JK. Immobilizing live
16		bacteria for AFM imaging of cellular processes. Ultramicroscopy. 2009;109(7):775-80.
17	16.	Marshall KC, Stout R, Mitchell R. Mechanism of the Initial Events in the Sorption of
18		Marine Bacteria to Surfaces. Journal of General Microbiology. 1971. p. 337-48.
19	17.	James SA, Powell LC, Wright CJ. Atomic Force Microscopy of Biofilms- Imaging,
20		Interactions, and Mechanics. In: Dhanasekaran D, editor. Microbial Biofilms - Importance
21		and Applications. InTech; 2016.
22	18.	Gad M, Ikai A. Method for immobilizing microbial cells on gel surface for dynamic AFM
23		studies. Biophys J. 1995;69(6):2226-33.
24	19.	Kasas S, Ikai A. A method for anchoring round shaped cells for atomic force microscope
25		imaging. Biophys J. 1995;68(5):1678–80.
26	20.	Ahimou F, Touhami A, Dufrêne YF. Real-time imaging of the surface topography of living
27		yeast cells by atomic force microscopy. Yeast. 2003;20(1):25-30.
28	21.	Yao X, Walter J, Burke S, Stewart S, Jericho MH, Pink D, et al. Atomic force microscopy
29		and theoretical considerations of surface properties and turgor pressures of bacteria.

1		Colloids Surfaces B Biointerfaces. 2002;23(2–3):213–30.
2	22.	Vadillo-Rodríguez V, Busscher HJ, Norde W, De Vries J, Dijkstra RJB, Stokroos I, et al.
3		Comparison of atomic force microscopy interaction forces between bacteria and silicon
4		nitride substrata for three commonly used immobilization methods. Appl Environ
5		Microbiol. 2004;70(9):5441–6.
6	23.	Cerf A, Cau JC, Vieu C, Dague E. Nanomechanical properties of dead or alive single-
7		patterned bacteria. Langmuir. 2009;25(10):5731–6.
8	24.	Formosa C, Pillet F, Schiavone M, Duval RE, Ressier L, Dague E. Generation of living
9		cell arrays for atomic force microscopy studies. Nat Protoc [Internet]. 2014 Dec
10		31;10(1):199–204. Available from:
11		http://www.nature.com/doifinder/10.1038/nprot.2015.004
12	25.	Bolshakova A V., Kiselyova OI, Filonov AS, Frolova OY, Lyubchenko YL, Yaminsky I V.
13		Comparative studies of bacteria with an atomic force microscopy operating in different
14		modes. In: Ultramicroscopy. 2001. p. 121-8.
15	26.	Micic M, Hu D, Suh YD, Newton G, Romine M, Lu HP. Correlated atomic force
16		microscopy and fluorescence lifetime imaging of live bacterial cells. Colloids Surfaces B
17		Biointerfaces. 2004;34(4):205-12.
18	27.	Arnoldi M, Fritz M, Bäuerlein E, Radmacher M, Sackmann E, Boulbitch A. Bacterial
19		turgor pressure can be measured by atomic force microscopy. Phys Rev E - Stat
20		Physics, Plasmas, Fluids, Relat Interdiscip Top. 2000;62(1 B):1034–44.
21	28.	Parikh SJ, Chorover J. ATR-FTIR spectroscopy reveals bond formation during bacterial
22		adhesion to iron oxide. Langmuir. 2006;22(20):8492–500.
23	29.	Doktycz MJ, Sullivan CJ, Hoyt PR, Pelletier DA, Wu S, Allison DP. AFM imaging of
24		bacteria in liquid media immobilized on gelatin coated mica surfaces. Ultramicroscopy.
25		2003;97(1–4):209–16.
26	30.	Camesano TA, Liu Y, Datta M. Measuring bacterial adhesion at environmental interfaces
27		with single-cell and single-molecule techniques. Adv Water Resour. 2007;30(6-7):1470-
28		91.
29	31.	Johnson DJ, Al Malek SA, Al-Rashdi BAM, Hilal N. Atomic force microscopy of

- 1 nanofiltration membranes: Effect of imaging mode and environment. J Memb Sci. 2 2012;389:486-98. 3 32. Verwey EJW. Theory of the stability of lyophobic colloids. The Journal of physical and 4 colloid chemistry. 1947. p. 631-6. 5 33. Derjaguin B, Landau L. Theory of the stability of strongly charged lyophobic sols and of 6 the adhesion of strongly charged particles in solutions of electrolytes. Prog Surf Sci. 7 1993;43(1-4):30-59. 8 34. Fletcher M, Pringle JH. The effect of surface free energy and medium surface tension on 9 bacterial attachment to solid surfaces. J Colloid Interface Sci. 1985;104(1):5-14. 10 35. Pratt-Terpstra IH, Weerkamp a H, Busscher HJ. Adhesion of oral streptococci from a 11 flowing suspension to uncoated and albumin-coated surfaces. J Gen Microbiol [Internet]. 12 1987;133(11):3199–206. Available from: http://www.ncbi.nlm.nih.gov/pubmed/3446750 13 36. Rutter PR, Vincent B. Physicochemical Interactions of the Substratum, Microorganisms, 14 and the Fluid Phase. In: Marshall KC, editor. Microbial Adhesion and Aggregation: 15 Report of the Dahlem Workshop on Microbial Adhesion and Aggregation Berlin 1984. 16 January 15--20 [Internet]. Berlin, Heidelberg: Springer Berlin Heidelberg; 1984. p. 21-38. 17 Available from: http://dx.doi.org/10.1007/978-3-642-70137-5_3 18 37. Pratt-Terpstra IH, Weerkamp AH, Busscher HJ. On a relation between interfacial free 19 energy-dependent and noninterfacial free energy-dependent adherence of oral 20 streptococci to solid substrata. Curr Microbiol [Internet]. 1988 Nov;16(6):311-3. 21 Available from: http://link.springer.com/10.1007/BF01568537
- 38. Truong VK, Lapovok R, Estrin YS, Rundell S, Wang JY, Fluke CJ, et al. The influence of
 nano-scale surface roughness on bacterial adhesion to ultrafine-grained titanium.
 Biomaterials. 2010;31(13):3674–83.
- 39. Arnold JW, Bailey GW. Surface finishes on stainless steel reduce bacterial attachment
 and early biofilm formation: scanning electron and atomic force microscopy study. Poult
 Sci. 2000;79(12):1839–45.
- 40. Boyd RD, Verran J, Jones M V., Bhakoo M. Use of the atomic force microscope to
 determine the effect of substratum surface topography on bacterial adhesion. Langmuir.

1 2002;18(6):2343–6.

2	41.	Whitehead KA, Colligon J, Verran J. Retention of microbial cells in substratum surface
3		features of micrometer and sub-micrometer dimensions. Colloids Surfaces B
4		Biointerfaces. 2005;41(2-3):129-38.
5	42.	Bohinc K, Dražić G, Abram A, Jevšnik M, Jeršek B, Nipič D, et al. Metal surface
6		characteristics dictate bacterial adhesion capacity. Int J Adhes Adhes. 2016;68:39-46.
7	43.	Ivanova EP, Truong VK, Wang JY, Bemdt CC, Jones RT, Yusuf II, et al. Impact of
8		nanoscale roughness of titanium thin film surfaces on bacterial Retention. Langmuir.
9		2010;26(3):1973–82.
10	44.	Lüdecke C, Bossert J, Roth M, Jandt KD. Physical vapor deposited titanium thin films for
11		biomedical applications: Reproducibility of nanoscale surface roughness and microbial
12		adhesion properties. Appl Surf Sci. 2013;280:578-89.
13	45.	Truong VK, Rundell S, Lapovok R, Estrin Y, Wang JY, Berndt CC, et al. Effect of
14		ultrafine-grained titanium surfaces on adhesion of bacteria. Appl Microbiol Biotechnol.
15		2009;83(5):925–37.
16	46.	Singh AV, Galluzzi M, Borghi F, Indrieri M, Vyas V, Podestà A, et al. Interaction of
17		bacterial cells with cluster-assembled nanostructured titania surfaces: an atomic force
18		microscopy study. J Nanosci Nanotechnol [Internet]. 2013;13(1):77-85. Available from:
19		http://www.ncbi.nlm.nih.gov/pubmed/23646700
20	47.	Ivanova EP, Truong VK, Webb HK, Baulin VA, Wang JY, Mohammodi N, et al.
21		Differential attraction and repulsion of Staphylococcus aureus and Pseudomonas
22		aeruginosa on molecularly smooth titanium films. Sci Rep [Internet]. 2011;1:165.
23		Available from:
24		http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=3240996&tool=pmcentrez&re
25		ndertype=abstract
26	48.	Singh AV, Vyas V, Patil R, Sharma V, Scopelliti PE, Bongiorno G, et al. Quantitative
27		Characterization of the Influence of the Nanoscale Morphology of Nanostructured
28		Surfaces on Bacterial Adhesion and Biofilm Formation. Bansal V, editor. PLoS One
29		[Internet]. 2011 Sep 26;6(9):e25029. Available from:

http://dx.plos.org/10.1371/journal.pone.0025029

2	49.	Mitik-Dineva N, Wang J, Stoddart PR, Crawford RJ, Ivanova EP. Nano-structured
3		surfaces control bacterial attachment. In: Proceedings of the 2008 International
4		Conference on Nanoscience and Nanotechnology, ICONN 2008. 2008. p. 113-6.
5	50.	Park MR, Banks MK, Applegate B, Webster TJ. Influence of nanophase titania
6		topography on bacterial attachment and metabolism. Int J Nanomedicine.
7		2008;3(4):497–504.
8	51.	Mitik-Dineva N, Wang J, Truong VK, Stoddart PR, Malherbe F, Crawford RJ, et al.
9		Differences in colonisation of five marine bacteria on two types of glass surfaces.
10		Biofouling [Internet]. 2009 Jul 3;25(7):621–31. Available from:
11		http://www.tandfonline.com/doi/abs/10.1080/08927010903012773
12	52.	Díaz C, Schilardi PL, Dos Santos Claro PC, Salvarezza RC, Fernández Lorenzo De
13		Mele MA. Submicron trenches reduce the Pseudomonas fluorescens colonization rate
14		on solid surfaces. ACS Appl Mater Interfaces. 2009;1(1):136–43.
15	53.	Schaer-Zammaretti P, Ubbink J. Imaging of lactic acid bacteria with AFM - Elasticity and
16		adhesion maps and their relationship to biological and structural data. Ultramicroscopy.
17		2003;97(1–4):199–209.
18	54.	Puckett SD, Taylor E, Raimondo T, Webster TJ. The relationship between the
19		nanostructure of titanium surfaces and bacterial attachment. Biomaterials.
20		2010;31(4):706–13.
21	55.	Bagherifard S, Hickey DJ, de Luca AC, Malheiro VN, Markaki AE, Guagliano M, et al.
22		The influence of nanostructured features on bacterial adhesion and bone cell functions
23		on severely shot peened 316L stainless steel. Biomaterials. 2015;73:185–97.
24	56.	Neihof R, Loeb G. DISSOLVED ORGANIC MATTER IN SEAWATER AND THE
25		ELECTRIC CHARGE OF IMMERSED SURFACES. J Mar Res. 1974;32(1):5–12.
26	57.	Schneider RP, Leis A. Conditioning Films in Aquatic Environments. In: Encyclopedia of
27		Environmental Microbiology [Internet]. Hoboken, NJ, USA: John Wiley & Sons, Inc.;
28		2003. Available from: http://doi.wiley.com/10.1002/0471263397.env036
29	58.	Dufrene YF, Vermeiren H, Vanderleyden J, Rouxhet PG. Direct evidence for the

involvement of extracellular proteins in the adhesion of Azospirillum brasilense.

2 Microbiol. 1996;142:855–865.

- Gómez-Suárez C, Pasma J, van der Borden AJ, Wingender J, Flemming HC, Busscher
 HJ, et al. Influence of extracellular polymeric substances on deposition and redeposition
 of Pseudomonas aeruginosa to surfaces. Microbiology. 2002;148(4):1161–9.
- 6 60. Hwang G, Liang J, Kang S, Tong M, Liu Y. The role of conditioning film formation in
- 7 Pseudomonas aeruginosa PAO1 adhesion to inert surfaces in aquatic environments.
- 8 Biochem Eng J [Internet]. Elsevier B.V.; 2013;76:90–8. Available from:
- 9 http://dx.doi.org/10.1016/j.bej.2013.03.024
- Hwang G, Kang S, El-Din MG, Liu Y. Impact of conditioning films on the initial adhesion
 of Burkholderia cepacia. Colloids Surfaces B Biointerfaces. 2012;91(1):181–8.
- 12 62. Hwang G, Kang S, El-Din MG, Liu Y. Impact of an extracellular polymeric substance
- 13 (EPS) precoating on the initial adhesion of Burkholderia cepacia and Pseudomonas
- 14 aeruginosa. Biofouling [Internet]. 2012;28(6):525–38. Available from:
- 15 http://www.ncbi.nlm.nih.gov/pubmed/22686692
- 16 63. Wingender J, Strathmann M, Rode A, Leis A, Flemming H-C. Isolation and biochemical
- 17 characterization of extracellular polymeric substances from Pseudomonas aeruginosa.
- 18 In: Methods in Enzymology [Internet]. 2001. p. 302–14. Available from:
- 19 http://www.sciencedirect.com/science/article/pii/S0076687901365977
- 20 64. Conrad A, Kontro M, Keinänen MM, Cadoret A, Faure P, Mansuy-Huault L, et al. Fatty
- 21 acids of lipid fractions in extracellular polymeric substances of activated sludge flocs.
- 22 Lipids [Internet]. 2003;38(10):1093–105. Available from:
- 23 http://link.springer.com/10.1007/s11745-006-1165-y
- 24 65. Flemming H-C, Wingender J. Extracellular Polymeric Substances (EPS): Structural,
- 25 Ecological and Technical Aspects. In: Encyclopedia of Environmental Microbiology
- 26 [Internet]. Hoboken, NJ, USA: John Wiley & Sons, Inc.; 2003. Available from:
- 27 http://doi.wiley.com/10.1002/0471263397.env292
- 28 66. Flemming H, Wingender J. The biofilm matrix. Nat Rev Microbiol [Internet].
- 29 2010;8(9):623–33. Available from:

http://dx.doi.org/10.1038/nrmicro2415%5Cnhttp://www.ncbi.nlm.nih.gov/pubmed/206761

- 2
- 45 3 67. Beech I, Hanjagsit L, Kalaji M, Neal AL, Zinkevich V. Chemical and structural 4 characterization of exopolymers produced by Pseudomonas sp. NCIMB 2021 in 5 continuous culture. Microbiology. 1999;145(6):1491-7. 6 68. Yang Y, Wikieł AJ, Dall'Agnol LT, Eloy P, Genet MJ, Moura JJG, et al. Proteins 7 dominate in the surface layers formed on materials exposed to extracellular polymeric 8 substances from bacterial cultures. Biofouling [Internet]. 2016;32(1):95-108. Available 9 from: http://www.tandfonline.com/doi/full/10.1080/08927014.2015.1114609 10 Zaky A, Escobar I, Motlagh AM, Gruden C. Determining the influence of active cells and 69. 11 conditioning layer on early stage biofilm formation using cellulose acetate ultrafiltration 12 membranes. Desalination. 2012;286:296-303. 13 70. Pasmore M, Todd P, Smith S, Baker D, Silverstein J, Coons D, et al. Effects of 14 ultrafiltration membrane surface properties on Pseudomonas aeruginosa biofilm initiation 15 for the purpose of reducing biofouling. J Memb Sci [Internet]. 2001 Nov;194(1):15-32. 16 Available from: http://linkinghub.elsevier.com/retrieve/pii/S0376738801004689 17 71. Liu Y, Mi B. Effects of organic macromolecular conditioning on gypsum scaling of 18 forward osmosis membranes. J Memb Sci. 2014;450:153-61. 19 72. Kochkodan V, Hilal N. A comprehensive review on surface modified polymer 20 membranes for biofouling mitigation. Desalination. 2015. p. 187-207. 21 73. Suwarno SR, Hanada S, Chong TH, Goto S, Henmi M, Fane AG. The effect of different 22 surface conditioning layers on bacterial adhesion on reverse osmosis membranes. 23 Desalination [Internet]. Elsevier B.V.; 2016;387:1–13. Available from: 24 http://www.sciencedirect.com/science/article/pii/S0011916416300868 25 74. Dufrene YF. Recent progress in the application of atomic force microscopy imaging and 26 force spectroscopy to microbiology. Curr Opin Microbiol. 2003;6:317-23. 27 75. Abu-Lail NI, Camesano TA. Elasticity of pseudomonas putida KT2442 surface polymers 28 probed with single-molecule force microscopy. Langmuir. 2002;18(10):4071-81. 29 76. Touhami A, Nysten B, Dufrêne YF. Nanoscale mapping of the elasticity of microbial cells

1		by atomic force microscopy. Langmuir. 2003. p. 4539–43.
2	77.	Bayer ME, Carlemalm E, Kellenberger E. Capsule of Escherichia coli K29:
3		Ultrastructural preservation and immunoelectron microscopy. J Bacteriol.
4		1985;162(3):985–91.
5	78.	Bullitt E, Makowski L. Bacterial adhesion pili are heterologous assemblies of similar
6		subunits. Biophys J. 1998;74(1):623–32.
7	79.	Helenius J, Heisenberg C-P, Gaub HE, Muller DJ. Single-cell force spectroscopy. J Cell
8		Sci [Internet]. 2008 May 6;121(11):1785–91. Available from:
9		http://jcs.biologists.org/cgi/doi/10.1242/jcs.030999
10	80.	Savage DC, Fletcher M, editors. Bacterial Adhesion [Internet]. Boston, MA: Springer US;
11		1985. Available from: http://link.springer.com/10.1007/978-1-4615-6514-7
12	81.	Williams V, Fletcher M. Pseudomonas fluorescens adhesion and transport through
13		porous media are affected by lipopolysaccharide composition. Appl Environ Microbiol.
14		1996;62(1):100–4.
15	82.	Berne C, Ma X, Licata NA, Neves BRA, Setayeshgar S, Brun Y V., et al. Physiochemical
16		properties of caulobacter crescentus holdfast: A localized bacterial adhesive. J Phys
17		Chem B. 2013;117(36):10492–503.
18	83.	Dupres V, Alsteens D, Andre G, Verbelen C, Dufrêne YF. Fishing single molecules on
19		live cells. Nano Today. 2009. p. 262–8.
20	84.	Engel A, Müller DJ. Observing single biomolecules at work with the atomic force
21		microscope. Nat Struct Biol. 2000;7(9):715-8.
22	85.	Engel A, Gaub HE, Müller DJ. Atomic force microscopy: A forceful way with single
23		molecules. Curr Biol. 1999;9(4).
24	86.	Fredrickson GH. The theory of polymer dynamics. Curr Opin Solid State Mater Sci.
25		1996;1(6):812–6.
26	87.	Flory PJ, Volkenstein M. Statistical mechanics of chain molecules. Biopolymers
27		[Internet]. 1969;8(5):699–700. Available from:
28		http://doi.wiley.com/10.1002/bip.1969.360080514%5Cnpapers2://publication/doi/10.1002
29		/bip.1969.360080514

1	88.	Sandal M, Valle F, Tessari I, Mammi S, Bergantino E, Musiani F, et al. Conformational
2		equilibria in monomeric alpha-synuclein at the single-molecule level. PLoS Biol.
3		2008;6(1):e6.
4	89.	Dougan L, Li J, Badilla CL, Berne BJ, Fernandez JM. Single homopolypeptide chains
5		collapse into mechanically rigid conformations. Proc Natl Acad Sci U S A.
6		2009;106(31):12605–10.
7	90.	Farrance OE, Paci E, Radford SE, Brockwell DJ. Extraction of Accurate Biomolecular
8		Parameters from Single-Molecule Force Spectroscopy Experiments. ACS Nano
9		[Internet]. 2015 Feb 24;9(2):1315–24. Available from:
10		http://pubs.acs.org/doi/abs/10.1021/nn505135d
11	91.	Friedlander RS, Vogel N, Aizenberg J. Role of Flagella in Adhesion of Escherichia coli to
12		Abiotic Surfaces. Langmuir. 2015;31(22):6137–44.
13	92.	Bowen WR, Hilal N, Lovitt RW, Wright CJ. Direct measurement of the force of adhesion
14		of a single biological cell using an atomic force microscope. Colloids Surfaces A
15		Physicochem Eng Asp [Internet]. 1998;136(1–2):231–4. Available from:
16		http://www.scopus.com/inward/record.url?eid=2-s2.0-0032580422&partnerID=tZOtx3y1
17	93.	Bowen WR, Lovitt RW, Wright CJ. Atomic force microscopy study of the adhesion of
18		Saccharomyces cerevisiae . J Colloid Interface Sci. 2001;237(1):54-61.
19	94.	Le DTL, Guérardel Y, Loubière P, Mercier-Bonin M, Dague E. Measuring Kinetic
20		Dissociation/Association Constants Between Lactococcus lactis Bacteria and Mucins
21		Using Living Cell Probes. Biophys J [Internet]. 2011 Dec;101(11):2843–53. Available
22		from: http://linkinghub.elsevier.com/retrieve/pii/S0006349511012562
23	95.	Ovchinnikova ES, Krom BP, van der Mei HC, Busscher HJ. Force microscopic and
24		thermodynamic analysis of the adhesion between Pseudomonas aeruginosa and
25		Candida albicans. Soft Matter [Internet]. 2012;8(24):6454. Available from:
26		http://xlink.rsc.org/?DOI=c2sm25100k
27	96.	Emerson IV RJ, Bergstrom TS, Liu Y, Soto ER, Brown CA, McGimpsey WG, et al.
28		Microscale correlation between surface chemistry, texture, and the adhesive strength of
29		staphylococcus epidermidis. Langmuir. 2006;22(26):11311-21.

1	97.	Razatos A, Ong YL, Sharma MM, Georgiou G. Molecular determinants of bacterial	
2		adhesion monitored by atomic force microscopy. Proc Natl Acad Sci U S A.	
3		1998;95(19):11059–64.	
4	98.	Kang S, Elimelech M. Bioinspired single bacterial cell force spectroscopy. Langmuir.	
5		2009;25(17):9656–9.	
6	99.	Beaussart A, El-Kirat-Chatel S, Herman P, Alsteens D, Mahillon J, Hols P, et al. Single-	
7		cell force spectroscopy of probiotic bacteria. Biophys J. 2013;104(9):1886-92.	
8	100.	Diao M, Taran E, Mahler S, Nguyen TAH, Nguyen A V. Quantifying adhesion of	
9		acidophilic bioleaching bacteria to silica and pyrite by atomic force microscopy with a	
10		bacterial probe. Colloids Surfaces B Biointerfaces. 2014;115:229-36.	
11	101.	Beaussart A, El-Kirat-Chatel S. Quantifying the forces guiding microbial cell adhesion	
12		using single-cell force spectroscopy. Nat Protoc [Internet]. 2014;9(5):1049-55. Available	
13		from:	
14		http://www.ncbi.nlm.nih.gov/pubmed/24722404%5Cnhttp://www.nature.com/nprot/journa	
15		l/v9/n5/abs/nprot.2014.066.html	
16	102.	Rodriguez-Emmenegger C, Janel S, de los Santos Pereira A, Bruns M, Lafont F.	
17		Quantifying bacterial adhesion on antifouling polymer brushes via single-cell force	
18		spectroscopy. Polym Chem [Internet]. 2015;6(31):5740–51. Available from:	
19		http://xlink.rsc.org/?DOI=C5PY00197H	
20	103.	Aguayo S, Strange A, Gadegaard N, Dalby MJ, Bozec L. Influence of biomaterial	
21		nanotopography on the adhesive and elastic properties of Staphylococcus aureus cells.	
22		RSC Adv [Internet]. 2016;6(92):89347–55. Available from:	
23		http://xlink.rsc.org/?DOI=C6RA12504B	
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26	Figure	e legends	
27	Figure	a 1. Schematic representation of the Atomic Force Microscope.	
28 29 30	Figure 2. AFM images of bioprocess engineering surfaces (A) Cyclopore microfiltration membrane (3.2µm ²) (B) Humic acid layer fouling an ultrafiltration membrane (3.5µm ²) (C) Air bubbles at a membrane surface (0.75µm ²) (D) Stainless steel process plant surface.		

(10µm²).(E) Lawn of *Saccharomyces cerevisiae* cells (brewing yeast NCYC 1681) showing

- 2 budding scars (15 µm²) (F) Shewanella oneidensis cell showing protein clusters at surface
- formed during anaerobic respiration and microbial fuel cell operation (1.2µm²)

5 Figure 3. AFM Force spectroscopy curve of a single *S.cerevisiae* cell (stationary phase) with freshly cleaved mica in 10-2 M NaCl pH 5.0.

Figure 4. SEM image of a cell probe A) Saccharomyces cerevisiae B) Aspergillus niger