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5	The Interaction of Wood Nanocellulose Dressings and the Wound Pathogen P.	
6	aeruginosa	
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25 26 27	<b>Keywords</b> : Nanocellulose, Biofilms, <i>Pseudomonas aeruginosa</i> , COMSTAT, Characterisation	

#### 29 ABSTRACT

30 Chronic wounds pose an increasingly significant worldwide economic burden (over 31 £1 billion per annum in the UK alone). With the escalation in global obesity and 32 diabetes, chronic wounds will increasingly be a significant cause of morbidity and 33 mortality. Cellulose nanofibrils (CNF) are highly versatile and can be tailored with 34 specific physical properties to produce an assortment of three-dimensional structures 35 (hydrogels, aerogels or films), for subsequent utilization as wound dressing materials. 36 Growth curves using CNF (diameter <20 nm) in suspension demonstrated an interesting dose-dependent inhibition of bacterial growth. In addition, analysis of 37 biofilm formation (Pseudomonas aeruginosa PAO1) on nanocellulose aerogels (20 38 39  $g/m^2$ ) revealed significantly less biofilm biomass with decreasing aerogel porosity and 40 surface roughness. Importantly, virulence factor production by *P. aeruginosa* in the 41 presence of nanocellulose materials, quantified for the first time, was unaffected 42 (p>0.05) over 24 h. These data demonstrate the potential of nanocellulose materials 43 in the development of novel dressings that may afford significant clinical potential. 44

45

#### 47 **1. Introduction**

Normal wound healing follows a defined process involving; coagulation, 48 49 inflammation, cell proliferation, matrix repair, epithelialization and remodeling 50 (Bjarnsholt et al., 2008). In contrast, chronic wounds are typically characterized by 51 elevated inflammatory responses and tissue breakdown, and are slow or fail to heal, 52 causing a considerable reduction in patient 'quality-of-life'. With a globally ageing 53 population, the biomedical and socioeconomic burdens of chronic wounds are 54 worsening, with annual costs in the US estimated at > \$50 billion per annum and 55 affecting 6.5 million people (2% of the population) (Jung et al., 2016). Chronic 56 wounds are always colonized with bacteria, and the pathogen Pseudomonas 57 aeruginosa is implicated in non-healing wounds (Davies et al., 2004). The ability of 58 these opportunistic pathogens to produce a thick, mucoid wound biofilm is thought to 59 prevent an effective response by host immune defenses, thereby impairing wound 60 healing (Percival et al., 2012).

61 The most abundant naturally-occurring polymer is cellulose, which can be 62 obtained from a variety of sources including: wood, non-woody plants, agricultural 63 residues, algae and bacteria. In addition to bacterial cellulose (Czaja, Krystynowicz, 64 Bielecki & Brown, 2006; Petersen & Gatenholm, 2011; Portal, Clark & Levinson, 65 2009), nanocellulose from wood pulp has been proposed as an appropriate material 66 for wound dressing applications (Chinga-Carrasco & Syverud, 2014; Powell et al., 67 2016; Rees et al., 2015). Nanocellulose can be produced in large quantities using 68 effective chemical pre-treatments, which facilitate deconstruction of the wood fiber 69 wall into cellulose nanofibrils (CNF) (Chinga-Carrasco & Syverud, 2014; Saito, 70 Nishiyama, Putaux, Vignon & Isogai, 2006; Wågberg et al., 2008). TEMPO-71 mediated oxidation is one of the most applied procedures for CNF production, leading 72 to the introduction of carboxyl groups in the C6 position and small numbers of 73 aldehyde groups (Saito & Isogai, 2004; Saito et al., 2006). TEMPO-mediated 74 oxidation produces morphologically homogeneous CNF, having diameters less than 20 nm and lengths in the micrometer scale (Chinga-Carrasco, Yu & Diserud, 2011; 75 76 Saito & Isogai, 2004). Furthermore, this material can be used to produce an 77 assortment of three-dimensional structures e.g. strong, dense and smooth films 78 (Fukuzumi, Saito, Iwata, Kumamoto & Isogai, 2009) and structured hydrogels and 79 aerogels with high porosity, with capacity to absorb large quantities of moisture 80 (Chinga-Carrasco & Syverud, 2014; Syverud, Kirsebom, Hajizadeh & Chinga81 Carrasco, 2011). We recently developed a method to produce CNF from *P. radiata* 82 fibers, with low endotoxin levels (<50 endotoxin units/g cellulose) (Nordli, Chinga-83 Carrasco, Rokstad & Pukstad, 2016) and demonstrated that TEMPO CNF exhibits no 84 toxicity towards 3T3 cells (Alexandrescu, Syverud, Gatti & Chinga-Carrasco, 2013) 85 and primary human skin cells i.e. fibroblasts or keratinocytes (Nordli et al., 2016; 86 Tehrani, Nordli, Pukstad, Gethin & Chinga-Carrasco, 2016). These studies suggested 87 the potential of these materials for wound dressing applications (Nordli et al., 2016).

88 With the increasing economic burden chronic wounds are placing on 89 worldwide healthcare systems, the use of non-toxic, biodegradable biopolymers from 90 an abundant, sustainable source, such as wood CNF, for wound healing applications 91 would be a distinct advantage in our current 'throw-away' culture. Many 92 commercially available wound dressings possess a high capacity to absorb moisture 93 due to their porous, three-dimensional structure. However, little attention has been 94 paid of the effect of CNF materials (also with high moisture absorption capabilities) 95 on biofilm formation of typical wound pathogens.

This study assessed (i) the effect of CNF hydrogels, at varying concentrations of CNF dispersion, on *Pseudomonas aeruginosa* (PAO1) growth in suspension, (ii) the effect of surface and bulk structure of CNF dressings on biofilm formation of PAO1 and, (iii) for the first time, the effects of these porous structures on pseudomonal virulence factor production.

101

#### 102 **2. Materials and Methods**

103 2.1. CNF production

104 Pinus radiata pulp fibers were used as the raw material for CNF production. 105 The carbohydrate composition of the pulp fibers has previously been reported 106 (Chinga-Carrasco et al., 2012). TEMPO (2,2,6,6-tetramethylpiperidinyl-1-oxyl) 107 mediated oxidation was applied as a pretreatment, using 3.8 mM NaClO/g cellulose, 108 pH 10.5 (Saito et al., 2006). Oxidized fibers (1% w/v) were fibrillated following 109 three passes through a Rannie 15 type 12.56X homogenizer. The degree of 110 polymerization (DP 709), aldehyde groups (71 µM/g) and carboxyl groups (855 111  $\mu$ M/g) have previously been quantified (Rees et al., 2015).

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- 113

#### 114 2.2. Sample Preparation

115 CNF samples were air- or freeze-dried to produce tight (films) or porous 116 structures (aerogels), respectively. Freeze-dried samples were prepared at 0.2, 0.4 or 0.6% (w/v) CNF dispersions, frozen in petri dishes at -20°C and freeze-dried for 48 h. 117 118 Although differing CNF concentrations were used in the liquid dispersions, the actual 119 grammage for all the resulting dried films and aerogels was 20 g/m<sup>2</sup>. All the samples were cut into 1 or 2 cm<sup>2</sup> sections and sterilized by  $\gamma$ -irradiation (15 kGy). 120 Commercially-available wound dressings AquaCel<sup>®</sup> and AquaCel Ag<sup>®</sup> (ConvaTec 121 122 Ltd, Deeside) were used as controls.

123

#### 124 2.3. Absorption Measurements

Fluid absorption of the test and control materials was assessed as previously described (Fulton et al., 2012). Dressing materials were cut into 1 cm<sup>2</sup> squares and weighed to the nearest 0.001 g. Samples were submerged in phosphate buffer saline, pH 7.4 (PBS) for 24 h, covered to prevent evaporation. Samples were then removed from the PBS and allowed to drip for 30 s and a wet weight obtained. Fluid absorption (g/g) = weight of the fluid absorbed/dressing dry weight.

131

## 132 2.4. Taxonomy of the samples

133 The varying percentages of CNF (wet weight) accounted for varying degrees of 134 porosity in the dried materials. Once dried, there was no difference in the 135 concentration of CNF incorporated in each of the prototype samples ( $20 \text{ g/m}^2$ ; Table 136 1).

137

#### 138 **Table 1**

139 Characteristics of the nanocellulose (CNF) materials used in this study.

Sample name	Concentration of CNF dispersion (w/v)	Sample preparation (resulting material) <sup>a</sup>
A0.2	0.2%	Freeze-dried (aerogel)
A0.4	0.4%	Freeze-dried (aerogel)
A0.6	0.6%	Freeze-dried (aerogel)
Film	0.2%	Air-dried (film)

## 142 2.5. Screening for the ability of CNF to support bacterial growth.

143 Bacterial growth and carbon utilization studies were undertaken using CNF 144 dispersions. CNF dispersions were sterilized by  $\gamma$ -irradiation, and then diluted in 145 deionized sterilized water to a final CNF concentration of 0.2, 0.4, 0.6 and 0.8%. The 146 ability of CNF to inhibit or promote growth of *P. aeruginosa* PAO1 in planktonic 147 culture was examined. Overnight cultures of PAO1 were diluted to OD<sub>600</sub> 0.08 148 (colony forming units =  $1 \times 10^8$ ) in either MH broth, PBS or deionized water, and 149 mixed 1:2 (v/v) with CNF dispersion or water in a 24 well plate. Plates were 150 incubated at 37°C aerobically for 24 h measuring  $OD_{600}$  every hour at 600 nm ( $OD_{600}$ ) 151 in a FLUOstar Optima plate reader (BMG LABTECH).

- 152
- 153 2.6. Log<sub>10</sub> reduction assay

154 Time kill assays adapted from Ong, Wu, Moochhala, Tan, & Lu (2008) were 155 used to evaluate the antimicrobial efficacy of aerogel and film materials in comparison to a commercially available wound dressing (AquaCel®) control. CNF 156 samples or AquaCel  $(Ag)^{\text{@}}$  were added to a 6-well plate;  $2 \times 2 \text{ cm}^2$  per well in total. 157 158 MH-broth (6 ml) and 60 µl of PAO1 overnight culture adjusted to OD<sub>600</sub> 1.0 were then added to each well. Plates were incubated at 37°C, 24 h (aerobically), with 159 shaking. To enumerate bacterial growth, 10 µl drops (3 x 10 µl per dilution) of 160 bacterial suspension was removed from each dilution and dropped onto the surface of 161 162 an MH-agar plate. The drops were air-dried before being incubated overnight at 37°C to enumerate colony forming units (CFU/ml). Log reduction was calculated as log10 163 164 CFU/ml (initial bacteria upon challenge) minus log10 CFU/ml (surviving bacteria at 165 time point after challenge). Bactericidal activity was defined as a  $\geq 3 \log 10 \text{ CFU/ml}$ 166 (equivalent to  $\geq$ 99.9%) reduction in bacterial numbers.

167

# 168 2.7. Assessment of virulence factors

169 Overnight cultures of *P. aeruginosa* PAO1 were adjusted to  $OD_{600}$  1.0, and 60 µl 170 were added to MH-broth (6 ml) in a 6-well plate and grown for 24 h ± CNF samples 171 or AquaCel (Ag)<sup>®</sup> (2×2 cm<sup>2</sup>). Bacterial cultures were then centrifuged (10000 g) for 172 10 min to produce a cell-free culture supernatant, used for extraction of virulence 173 factors. 174 Pyocyanin pigment was extracted from the cell-free supernatant using chloroform 175 (3:2; v/v). Pyocyanin (in the chloroform-phase) was re-extracted with 0.2 M HCl 176 (2:1; v/v) and the absorbance read at 540 nm (Sarabhai, Sharma & Capalash, 2013). 177 Rhamnolipids were extracted from the cell-free supernatant using ethyl acetate 178 (1:1, v/v), vortexed for 15 sec and centrifuged (10000 g, 4°C, 5 min). The top layer 179 was removed and the extraction repeated (x2) before allowing the ethyl acetate to 180 evaporate overnight. Deionized water was used to dissolve the precipitate and orcinol 181 reagent (0.19% orcinol in 53% H<sub>2</sub>SO<sub>4</sub>; 1:9, v/v) added. The sample was incubated at 182 80°C for 30 min before reading the absorbance at 421 nm (Smyth, Perfumo, McClean, 183 Marchant & Banat, 2010).

Protease activity was determined using 2% azocasein solution in 50 mM PBS, pH 7. The azocasein solution was incubated with the cell-free supernatant (1:1; v/v) in a total reaction volume of 400  $\mu$ l, for 1 h at 37 °C. The reaction was stopped by the addition of 500  $\mu$ l 10% trichloroacetic acid, and residual azocasein removed by centrifugation (8000 g, 5 min). Protease absorbance was measured at 400 nm (Adonizio, Kong & Mathee, 2008).

For elastase determination, cell-free supernatant was mixed (3:1; v/v) with elastin-congo-red solution (5 mg/ml in 0.1 M Tris-HCl pH 8; 1 mM CaCl<sub>2</sub>). Samples were incubated at 37 °C, for up to 16 h, at 200 rpm, centrifuged at 3000 g (10 min) before absorbance was read at 490 nm (Sarabhai et al., 2013).

194

#### 195 2.8. Microscopy characterization of CNF structures

Copper grids were immersed in a 0.01% suspension of the CNF sample and
stained with uranyl acetate. Scanning Transmission Electron Microscopy (STEM)
using a Hitachi S-5500 electron microscope was used to acquire images in bright field
mode (x150000 magnification), using an acceleration voltage of 30 kV.

AFM imaging (tapping mode) of the film sample was performed with a Multimode AFM with Nanoscope V controller, (Digital Instruments) and images were acquired in ScanAsyst mode at room temperature. The AFM tips of spring constant value ~0.4 N/m were purchased from Bruker AFM probes. The image size was 2  $\mu$ m x 2  $\mu$ m, with a resolution of 1.95 nm/pixel.

The aerogels and films were freeze-dried for SEM analysis. Cross-sections of the films and aerogels were prepared by ion milling (using an IM4000 system), where the milling time was 5 hours at 2.5 kV. In addition, samples were cut from the aerogels and films using an 8 mm punch biopsy for surface analysis. All samples
were sputter-coated with gold and images acquired using a Hitachi scanning electron
microscope (SEM, SU3500), in secondary electron imaging mode.

211

212 2.9. Confocal laser scanning microcopy (CLSM) and scanning electron microscopy
213 (SEM) of biofilm growth on CNF materials

CNF samples  $(1 \text{ cm}^2)$  were added to a 12-well plate to which contained MH broth 214 215 (3 ml) and 60 µl of PAO1 overnight culture adjusted to OD<sub>600</sub> 0.4. Plates were 216 incubated at 37°C in a static aerobic environment for 24 or 48 h before removing the 217 supernatant and then washing the biofilms once with deionised water/PBS. Biofilms 218 were stained with LIVE/DEAD<sup>®</sup> BacLight<sup>TM</sup> bacterial viability kit (Invitrogen, Paisley, UK) containing SYTO 9 dye (staining LIVE cells, green) and propidium 219 220 iodide (staining DEAD cells, red) and incubated in the dark (10 min) before mounting 221 on microscope slides with spacers, being set in Vectorshield (Vector Laboratories, 222 UK) and having a coverslip on top, sealed with nail varnish. Biofilms were imaged 223 with a Leica TCS SP2 confocal system (x63). Bacterial growth was quantified using 224 COMSTAT image-analysis software (Heydorn et al., 2000).

Biofilm samples were prepared for SEM analyses by immersion in glutaraldehyde (2.5%) for 24 h, before being washed with deionised water (x4) and immersed in fresh deionised water, and then frozen (24 h) and freeze-dried for 24 h. The biofilms were then imaged on the Tescan Vega SEM at 5 kV. Additionally, the A0.6 aerogel with PAO1 biofilm growth was also ion-milled (as described above) and SEM images acquired in secondary electron imaging mode.

231

232 2.10. Statistical analysis

GraphPad Prism 3 was used to perform statistical analysis (GraphPad software Inc, La
Jolla, USA) including one-way ANOVA using Tukey-Kramer post-test (growth curve
data and COMSTAT data) and Dunnet's multiple comparison tests (log<sub>10</sub> reduction
and virulence data). P<0.05 was considered significant.</li>

237

# **3. Results**

## 239 3.1. Growth of P. aeruginosa PAO1 in CNF hydrogel suspensions

240 Irradiated CNF samples at  $\geq 0.4\%$  concentrations indicated dose-dependent 241 inhibition of *P. aeruginosa* PAO1 (Fig. 1), possibly reflecting increased viscosity of 242 these dispersions or direct inhibition by CNF. No change in optical density was

243 observed for CNF in either PBS or water after 24 h, indicating *P. aeruginosa* did not

244 grow in, nor could it utilize CNF as a carbon source; growth only being observed in

the presence of MH broth (Fig. S1).



246

**Fig. 1.** Growth characteristics for *P. aeruginosa* PAO1 in CNF. *P. aeruginosa* PAO1 grown in either MH-broth/dH<sub>2</sub>O or MH-broth/ $\gamma$ -irradiated CNF (adjusted to give final concentrations of CNF at 0.2, 0.4, 0.6 or 0.8%). (n=3).

250

# 251 *3.2. Characterization of the Cellulose Materials*

The CNF material was highly fibrillated and structurally homogenous (Rees et al., 2015), containing CNF with diameters in the nanoscale (<20 nm) and with a high aspect ratio, as exemplified in this study (Fig. 2).

255 The aerogel, A0.2 (manufactured from a 0.2% CNF dispersion) possessed the 256 largest thickness, porosity and surface roughness, as demonstrated by SEM and fluid 257 absorption (Fig. 3A; Table S1). The porosity and surface roughness of the other 258 aerogels decreased with the increasing CNF concentration; A0.6 (manufactured from 259 a 0.6% CNF dispersion) showing the lowest (Figs 3A-C). In contrast, the film was far 260 more dense and extremely thin, with no apparent porosity (Fig. 3D). The commercial dressings AquaCel<sup>®</sup> and AquaCel Ag<sup>®</sup> displayed rough surfaces, which during the 261 262 course of the experiment altered, following fluid absorption. The more porous 263 (thicker) CNF aerogels absorbed more moisture when compared to the film; resembling absorbancy of AquaCel<sup>®</sup> and AquaCel Ag<sup>®</sup>. 264



Fig. 2. Highly fibrillated nanocellulose. (A) Atomic force microscopy imaging
(AFM). (B) Scanning Transmission Electron Microscopy imaging (STEM). Arrows
indicate individual nanofibrils.

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266



271

Fig. 3. Scanning electron microscopy of aerogels and films (cross-sectional and aerial views) made with CNF. (A) A0.2. (B) A0.4. (C) A0.6. (D) Film. Arrows indicate local thickness of the materials. (N.B. A0.2 was too thick to be visualized in its entirety within the maximum field of view).

- 276
- 277 3.3 Log<sub>10</sub> reduction assay

The ability of CNF films and aerogels (with differing porosities) to inhibit the growth of planktonic *P. aeruginosa* PAO1 showed that none of the CNF materials, (both aerogels,  $0.29-0.37\pm0.33-0.37$  and film samples,  $0.25\pm0.21$ ) demonstrated significant change in Log<sub>10</sub> CFU, being comparable to the negative controls AquaCel<sup>®</sup> dressing ( $0.24\pm0.16$ ) and MH ( $0.38\pm0.37$ ). AquaCel Ag<sup>®</sup> (the positive 283 control) containing ionic silver however, demonstrated significant bactericidal activity

284 ( $\geq$ 3 log-fold) reduction in Log<sub>10</sub> CFU compared to the control (6.62±0.94).

285

#### 286 *3.4. Virulence factor assays*

287 The production of virulence factors (pyocyanin, rhamnolipids, total proteases and elastase) by P. aeruginosa PAO1 was not affected by CNF (Fig. 4). However, for 288 289 elastase (measured at 1, 3 and 8 h) a significant difference to the control was 290 detectable at  $\geq 3$  hours of incubation. A modest decreasing trend in pyocyanin 291 production was noted based on the porosity of the aerogel materials, with A0.2 292 (largest pores, Fig. 3A) producing the greatest amount, whereas the A0.6 (smallest 293 pores, Fig. 3C) and film (no pores at the assessed scale) produced the least, however 294 this was found not to be significant.

295





Fig. 4. The effect of the different CNF film and aerogels on virulence factor production by *Pseudomonas aeruginosa* PAO1 (A) pyocyanin. (B) rhamnolipid. (C) total protease. (D) elastase, when compared to the negative controls MH broth and AquaCel Ag<sup>®</sup>. (\*significantly different as compared to the PAO1 control; n=3; p<0.05).

302

## 304 3.5. Confocal Laser Scanning Microscopy (CLSM)

305 P. aeruginosa PAO1 was able to form established biofilms on all the CNF and commercial (AquaCel<sup>®</sup>-based) dressing materials (Fig. 5, and Fig. S2 in the 306 supporting information) at 24 and 48 h. LIVE/DEAD<sup>®</sup>-staining confirmed bacterial 307 viability in the established biofilms. Little difference in bacterial viability was 308 evident for all the dressings, (apart from AquaCel Ag<sup>®</sup> which showed sparse biofilm 309 growth at 24 h). COMSTAT image analysis was employed to quantify biomass, 310 311 surface roughness (using a roughness coefficient) and mean biofilm thickness (Fig. 5 312 and Fig. S2) in test samples and control Aquacel Ag<sup>®</sup>.



- 314
- 315

Fig. 5. CLSM images and COMSTAT analysis of LIVE/DEAD<sup>®</sup> staining of 24 h biofilms grown on CNF aerogels A0.2, A0.4 and A0.6 (0.2, 0.4 and 0.6% respectively), film and commercial dressings (Aquacel and Aquacel Ag) showing mean: (A) biomass; (B) thickness and (C) roughness coefficient (n=3, \*p>0.001). All materials were significantly different to Aquacel Ag (data not shown on graphs).

321 COMSTAT analysis (n=7 images from n=3 replicates) revealed the viability of 322 the biofilms and that a distinct decrease in viable cell numbers was observed between 323 24 and 48 h, (Fig. 5A and Fig. S2A in the supporting information). In the aerogels, 324 biomass reflected material porosity; the greatest biomass being observed in A0.2 and 325 decreased to A0.6 (the least porous aerogel; p<0.05). AquaCel Ag<sup>®</sup> had very little 326 growth at both 24 and 48 h, reflecting the CLSM.

Biofilm thickness varied between samples tested, although at 24 h these differences were less apparent (Fig. 5B). As expected, mean thickness was higher on all the materials compared to AquaCel Ag<sup>®</sup> (Fig. 5B and Fig. S2B in the supporting information), with differences more apparent at 48 h.

There were significant differences in roughness coefficient between biofilms grown on the different materials, with biofilms grown on A0.2 being the roughest, while biofilms grown on A0.6 and Film were significantly smoother (p<0.05; 24 h); AquaCel Ag<sup>®</sup> being the roughest of all at both 24 and 48 h (Fig. 5C and Fig. S2C in the supporting information). Generally, at 48 h there were less distinguishable differences between surface roughness, probably reflecting the large proportion of non-viable cells present on all the materials at this time.

338



339

Fig. 6. SEM image of cross-section following ion milling of a *P. aeruginosa* biofilmgrowing on the A0.6 aerogel.

342

343 3.6. Scanning electron microscopy (SEM) of biofilm growth on CNF materials.

344 SEM imaging of a cross-section through a CNF aerogel following ion milling 345 confirmed that a large part of the biofilms were formed inside the aerogel structure (Fig. 6), probably due to the relatively large porosity of the samples. SEM imaging of
the surface of the materials can be found in Figs. S3-S6 in the supporting information.

348

# 349 **4. Discussion**

350 It has become increasingly recognized that chronic wounds are able to sustain 351 a substantial bacterial biofilm which (both directly and indirectly) influences clinical 352 outcome (Hill et al., 2010). The CNF materials did not support bacterial growth of 353 Pseudomonas aeruginosa PAO1, confirming that the CNF could not be used as a 354 bacterial carbon source by the wound pathogen PAO1 (Powell et al., 2016). Materials 355 that significantly impair microbial growth would be a distinct advantage for a wound 356 dressing and, interestingly, a dose-dependent inhibition of bacterial growth by the 357 CNF hydrogel was evident. This inhibition may, in part, reflect the apparent viscosity 358 of the medium, as motile species, such as P. aeruginosa, move through media obtaining available nutrients (Sampedro, Parales, Krell & Hill, 2015). In keeping 359 360 with this notion, the Log<sub>10</sub> reduction assay revealed that CNF aerogels and films 361 failed to inhibit microbial planktonic growth of P. aeruginosa PAO1 in these studies (as did the commercial wound dressing AquaCel<sup>®</sup>). The CNF materials may, 362 363 therefore, exhibit distinct properties in hydrogel, aerogel or film forms.

364 P. aeruginosa PAO1 produces the blue pigment, pyocyanin, which is a 365 secondary metabolite that interrupts cellular function and is also a virulence factor (Lau, Hassett, Ran & Kong, 2004) affecting mammalian cells by inhibiting cell 366 367 respiration and ciliary function (Ran, Hassett & Lau, 2003; Sorensen & Klinger, 368 1987). Rhamnolipids are also virulence factors produced by *P. aeruginosa* which aid 369 colonisation by acting as a surfactant, reducing surface and interfacial tension 370 (Soberón-Chávez, Lépine & Déziel, 2005). In addition, the production of 371 rhamnolipids in *P. aeruginosa* can enable the utilization of alternative carbon sources 372 (such as alkanes) by pseudosolubilisation of insoluble substrates that would not 373 normally be broken-down (Beal & Betts, 2000). P. aeruginosa also secretes proteases 374 and elastase that have been implicated as virulence factors (Caballer et al., 2001), 375 both of which are known to damage dermal matrix proteins and hinder cell migration 376 and wound healing (Oldal & Tranfny, 2005). The lack of effect of CNF wound 377 dressing materials on these virulence factors was characterized for the first time in the 378 present study.

379 Wound dressings represent, not only a physical barrier to infection and 380 trauma, but also incorporate antimicrobials to inhibit bacterial growth and provide an 381 optimal (moist) environment to facilitate wound healing (i.e. cell proliferation, 382 migration and differentiation). In close proximity to damaged skin, for considerable 383 periods of time, they must be biocompatible. A number of natural polymers are 384 already used as dressing materials including polysaccharides e.g. alginates and 385 chitins, and proteins e.g. collagen (Mogoşanu & Grumezescu, 2014). Unlike bacterial 386 nanocellulose, CNF has not traditionally been used for biomedical applications (Lin 387 & Dufresne, 2014). Whilst a number of researchers have investigated CNF 388 interactions in immune/inflammatory responses (Hua et al., 2015; Nordli et al., 2016), 389 as scaffolds for cell culture (Lou et al., 2014; Ninan et al., 2013) and as spray-dried 390 coatings (or stabilisers) for tablet production (Kolakovic et al., 2011), few have 391 studied their interaction with the bacteria and biofilms which characterize human 392 infection. This study clearly advances our previous studies with nanocellulose for 393 wound dressings (Nordli et al., 2016; Powell et al., 2016), and represents novel 394 findings in this area.

395 TEMPO-mediated oxidization of fibers is used to facilitate breakage of native 396 hydrogen bonding of individual cellulose fibrils from wood cellulose fibers without 397 causing extensive damage to the materials which would otherwise occur (Isogai, Saito 398 & Fukuzumi, 2011). This chemical pre-treatment is followed by physical 399 homogenization of the materials; the fibrillation process determining the 400 characteristics of the resulting material, *i.e.* fibril size. This resultant uniform, highly-401 fibrillated material is suitable for a range of medical applications and may be 402 functionalized/modified in individual biomedical applications (Lin & Dufresne, 403 2014). An important consideration for clinical use is sterility. The lack of structural 404 changes in aerogels and films following  $\gamma$ -irradiation reflects its suitability for large-405 scale manufacturing/processing. The recent demonstration confirming that ultrapure 406 CNF (lipopolysaccharide content <50 endotoxin units/gram cellulose) meets the endotoxin limits required in medical devices for wound management (Nordli et al., 407 408 2016), was also reassuring.

Three-dimensional visual assessment of the CNF CLSM images proved difficult and subjective due to the inherently rough surfaces and heterogeneous nature of the biofilms which formed. COMSTAT image analysis was therefore employed to characterize and study the biofilms formed on the materials; a technique that proved 413 useful. However, in terms of biomass and thickness, COMSTAT was only able to 414 quantify changes in bacteria which remained adherent to the material surface and 415 were visualised by CLSM. It was evident that biofilms infiltrate the dressing 416 materials. Ion milling of biofilms growing on the materials showed that this was 417 evident in the aerogels used in this study. Hence, the biomass and thickness data for 418 the aerogels (being extremely porous) will have been substantially underestimated 419 compared to those of the film (possessing no apparent pores for the biofilms to 420 invade).

421 Surface roughness plays a significant role in bacteria adhesion (Hsu, Fang, 422 Borca-Tasciuc, Worobo & Moraru, 2013; Yoda et al., 2014). This was confirmed in 423 our study as the aerogel with the lowest surface roughness and porosity (A0.6) 424 demonstrated the lowest biofilm growth (in terms of both thickness and biofilm 425 height) and this was also markedly less than that observed on the commercial AquaCel<sup>®</sup> dressings. Aerogels are likely to have a clinical advantage over the film for 426 427 wound applications due to their enhanced absorbency; chronic wounds often 428 producing large amounts of exudate. This study showed that, despite having identical 429 grammage, the aerogels (A0.2, A0.4 and A0.6) possessed distinctly different material 430 properties (e.g. surface roughness, fluid absorption). The commercial dressings AquaCel<sup>®</sup> and AquaCel Ag<sup>®</sup> displayed an inherently rough surface, which 431 progressively altered during the course of the experiment to become smoother with 432 433 increased absorption of fluids as noted by Walker, Hobot, Newman & Bowler (2003).

434 Our results indicate an ability to fabricate the desired surface/pore structure of 435 the CNF materials and their absorbance, which will be important in distinct clinical 436 applications. For example, wound dressings should have a smooth surface (to reduce 437 bacterial adhesion), with adequate porosity to enhance absorbency (and retain wound 438 fluid), whilst maintaining a moist environment to promote healing. These 439 microbiological studies show the potential use of CNF as a wound dressing material. 440 Not only was it demonstrated that CNF dispersions did not support bacterial growth 441 but, when the CNF dispersions were challenged with P. aeruginosa PAO1, bacterial 442 growth was inhibited. Using CNF as a potential wound dressing material represents 443 an important clinical use for this sustainable resource, with properties comparable to 444 those of commercially-available dressings.

445

# 446 **5.** Conclusions

447 Our previous studies have importantly shown these materials to be non-toxic to 448 human cells. This study confirms that CNF produced using differing manufacturing 449 processes generates a potential wound dressing material, which allows for absorbency 450 based on porosity to be controlled. Bacterial virulence was unaltered when grown on 451 these materials, showing equivalency with other commercially available wound 452 dressings already on the market. Moreover, these materials are biodegradable and 453 may be renewably sourced; both of which are important for biosustainability. Also as 454 the physical and chemical composition of CNF is highly malleable, the development 455 of nanocellulose dressings offers significant clinical potential.

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## 458 ASSOCIATED CONTENT

# 459 Supporting Information

Growth characteristics for *P. aeruginosa* PAO1 in CNF; Absorbancy of the CNF
materials used in this study; CLSM images and COMSTAT analysis of
LIVE/DEAD<sup>®</sup> staining of 48 h biofilms grown on CNF aerogels; and SEM images of
24 and 48 h *P. aeruginosa* PAO1 biofilms grown on the CNF materials.

464

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