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1	Additive Manufactured Graphene-based electrodes exhibit beneficial
2	performances in <i>Pseudomonas aeruginosa</i> microbial fuel cells
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33 Abstract

A commercial polylactic acid/graphene (8 wt.%) composite filament was used to 34 35 additive manufacture (AM) graphene macroelectrodes (AM-G_Ms). The electrode surfaces were 36 characterised and *Pseudomonas aeruginosa* was selected as the exoelectrogen. The MFC was 37 optimised using growth kinetic assays, biofilm formation, and quantification of pyocyanin 38 production (via liquid chromatography-mass spectrometry) in conditions that were 39 representative of the batch-fed MFC configuration utilised. Cell potential and bacterial viability 40 was recorded at 0h, 24h, 48h, 72h, 96h and 120h, power density and current density were 41 calculated. There was no significant difference between P. aeruginosa cell proliferation in 42 either media tested, interestingly, no accumulation of pyocyanin was evident. Additively 43 manufactured electrodes comprised of graphene (AM-G_Ms) were successfully applied in a P. *aeruginosa* MFC configuration and power outputs (110.74 \pm 14.63 μ Wm⁻²) produced were 44 comparable to that of the 'benchmark' electrode, carbon cloth ($93.49 \pm 5.17 \ \mu Wm^{-2}$). The AM-45 46 G_Ms demonstrated power/current outputs similar to that of the carbon cloth electrodes in both 47 anaerobic LB and glucose-based media over 120 h; the AM-G_Ms had no significant detrimental 48 effect on *P. aeruginosa* viability. This study highlights the potential application of additive 49 manufactured electrodes with the incorporation of nanomaterials (e.g. graphene) as one 50 approach to enhance power outputs.

51

52 Keywords: Microbial Fuel Cell; Additive Manufacturing; Graphene; *Pseudomonas* 53 *aeruginosa*; Pyocyanin; Biotechnology.

54

55 Word Count: 7220

57 Abbreviations

Description	Identifier
Additive Manufactured Polylactic Acid	AM-PLA
Additive Manufacturing	AM
Additively Manufactured Graphene Electrodes	AM-G _M s
Average Roughness	S_a
Cationic Exchange Membrane	CEM
Closed Circuit Voltage	CCV
Colony Forming Units per mL	CFU mL ⁻¹
Crystal Violet Biofilm Assay	CVBA
E cell	Cell Potential
Fusion Deposition Modelling	FDM
Heterogeneous Rate Transfer Constants	k^0_{obs}
Liquid Chromatography Mass-Spectroscopy	LC-MS
Luria Bertani	LB
Microbial Fuel Cells	MFCs
Optical Density	OD
Polylactic Acid	PLA
Scanning Electron Microscopy	SEM

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72 **1.0. Introduction**

Microbial fuel cells (MFCs) are systems in which microorganisms convert chemical energy to electrical energy, usually by utilising an organic electron donor [1]. Microbial fuel cells (MFCs) are an area of vast interest in biotechnology due to their remarkable versatility which includes ambient operating conditions, they can also be utilised as a dual-approach to both produce renewable energy production and treat wastewater [2, 3]. However, the application of MFCs at an industrially relevant scale is yet to be realised, due to issues with scalability, therefore it is imperative to enhance the overall system efficiency [4-6].

80 The manufacturing and assembly process of MFCs can be significantly improved via 81 the generation of monolithic designs [7]. One possible route to achieve such designs and 82 improve MFC architectures, electrodes and ion exchange membranes is additive manufacturing 83 (AM), more commonly known as 3D-Printing. The application of additive manufacturing 84 provides a unique platform for both the rapid design and fabrication of complex structures with 85 novel materials [8-10]. Previously, such architectures were not feasible using more traditional 86 manufacturing routes [11-13]. In an MFC, the anode must be conductive and also exhibit good 87 biocompatibility to sustain bacterial growth, furthermore, a high degree of surface area has 88 been shown to be advantageous [9]. In 2017, You et al. 3D-Printed the first polymer based 89 electrode for MFC application, where the conductive polylactic acid (PLA) filament 90 (ProtoPlant, USA) was printed as a rectangular mesh to provide greater surface area [7]. Whilst, 91 the 3D-Printed electrodes obtained lower outputs than the conventional carbon veil electrodes 92 tested, the results were promising [7]. In another study, porous carbon was fabricated via a 93 controlled carbonation process. The 3D-Printed anodes (pore size: 300 µm) produced a 94 maximum power density of 233.5 \pm 11.6 mW m⁻², which represented a significant 95 improvement in electrochemical performance when compared to the 2D control (carbon cloth) which produced a maximum power density of 69.0 ± 4.7 mW m⁻² [14]. The application of 96 additive manufacturing clearly has unexploited potential which could enhance MFC power 97 98 outputs.

99 Carbon-based materials are the most widely used electrode type in MFC configurations 100 as microbial growth on the surface of metals can accelerate corrosion of the surface; metals 101 submerged in aqueous solutions can produce metal ions [15-18]. Carbon cloth is one of the 102 most widely utilised electrode materials and is often used as a benchmark in MFC studies [15, 103 19, 20]. Carbon cloth is an ideal electrode for MFC configurations as it demonstrates excellent 104 electrical conductivity, good biocompatibility and a large surface area (due to enhance 105 porosity). Carbon cloth is however excessively expensive for application in MFC

configurations (ca. USD\$ 1000 m⁻²) [21]. Therefore, a substantially cheaper (per 100 g) and 106 107 more versatile electrode material was developed (Additive manufactured graphene 108 macroelectrodes; AM-G_Ms) and tested herein. Additive manufacturing equipment and raw 109 materials (*i.e.* PLA) are relatively inexpensive and cost will continue to decrease as research 110 and development into AM increases [22, 23]. Furthermore, Lanzotti et al., (2019) demonstrated 111 the use of recycled PLA as a viable option for AM, this would undoubtedly reduce material 112 expenditure and provide a more economically sustainable, alternative electrode material for 113 MFC configurations [24]. The incorporation of graphene into PLA filaments and other 114 feedstocks for AM is one potential option for improving MFC electrode materials. Graphene, which is defined as a 2D monolayer lattice of sp² hybridised carbon atoms, has unique 115 properties that make it an ideal electrode material. Such properties include: high electron 116 117 conductivity, a large theoretical surface area and the ability to sustain high current densities 118 [25-27]. The cost of the raw PLA/graphene composite filament is 100 USD /100 g, this is 119 capable of producing ca. 13, 333 of the electrodes utilised throughout this study, offering a 120 significant advantage over more expensive, traditional electrode material types [28].

121 Pseudomonas aeruginosa has previously demonstrated exoelectrogenic properties and 122 thus this bacterium has been utilised in a number of MFC configurations [6, 29-31]. The power 123 outputs obtained by P. aeruginosa MFCs can vary depending on a range of factors, which 124 include, phenazine (e.g. pyocyanin) production, biofilm formation and flagella/pili (i.e., 125 nanowire) efficiency [31, 32]. The exoelectrogenic properties of *P. aeruginosa* is due to the 126 production of soluble redox mediators, such as phenazines [33]. Phenazines exhibit electron 127 transfer mediating properties by acting as electron shuttles, which can transfer liberated 128 electrons to the anode of a MFC [34]. The most predominant and well-characterised natural 129 phenazine secreted by *P. aeruginosa* is pyocyanin [29, 35]. The production rate of pyocyanin 130 by *P. aeruginosa* is affected by a range of factors including, nutrient availability, physiological 131 status and growth phase [29, 36]. The pyocyanin concentration of a MFC has a direct 132 correlation with enhanced power generation, due to the ability of the phenazine to transport electrons through the bacterial cell membrane [30]. Interestingly, the addition of pyocyanin 133 134 produced by *P. aeruginosa* can be utilised by other microorganisms (which may be incapable 135 of pyocyanin production) in MFCs, thus resulting in enhanced current production [34, 37].

In this paper we report the fabrication and characterisation of the AM-G_Ms, for the first time, produced from a commercially available polylactic acid (PLA)-graphene feedstock. These electrodes were then trialled, for the first time, in a MFC configuration and are shown to be comparable to (more expensive (when manufactured per meter)) carbon cloth electrodes. 140

141 **2.0. Methods**

142 2.1 Fabrication of additive manufactured graphene macroelectrodes (AM-G_{MS})

143 The AM designs utilised throughout this study were produced using a MakerBot 144 Replicator + (Makerbot Industries, USA) fused deposition modelling (FDM) 3D-printer with 145 a direct drive extruder at a temperature of 210 °C. A conductive polylactic acid (PLA) / 146 graphene (8 wt%) composite filament (Black Magic 3D, USA) was 3D-printed into electrodes, 147 known henceforth throughout this manuscript as additive manufactured graphene 148 macroelectrodes (AM-G_Ms). An unfilled (pure) PLA filament (AM-PLA), was used as the 149 control throughout all of the experiments presented in this study. This AM-G_Ms has a reported conductivity of 2.13 S/cm [13]. The AM-G_Ms designs were drawn via Fusion 360 software 150 151 (Autodesk, USA) to create a circular disc architecture with a diameter of 6.0 mm, thickness of 1.0 mm and theoretical surface area of 7.5 cm². A radial connecting strip (1.0 mm x 1.0 mm 152 and 10.0 mm long) was also incorporated into the design to allow the simple connection of the 153 154 electrode to the electrochemical system.

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156 2.2 Bacterial Cell Culture

157 Prior to the start of the experiment, P. aeruginosa strain ATCC 9027 was cultured onto 158 Luria Bertani (LB; BD Difco[™], UK) agar. Inoculated plates were incubated at 37 °C for 24 h 159 in aerobic conditions. Pseudomonas aeruginosa strain ATCC 9027 was inoculated into 10 mL 160 LB or glucose-based broth (10 g glucose, 5 g yeast extract, 6.8 g sodium bicarbonate and 8.5 161 g sodium phosphate monobasic per litre; [38]) and incubated in either aerobic or anaerobic conditions at 37 °C, with agitation for 18 h. For anaerobic conditions, cultures were purged 162 163 with nitrogen for 30 min to generate an anaerobic environment using a silicon rubber septa 164 (Suba Seal, Merck, UK). Cultures were vortexed (5 s) and centrifuged (2,342 g for 10 min). 165 Bacterial pellets were re-suspended and cells were washed with 10 mL of sterile distilled water, 166 followed by centrifugation (2,342 g for 10 min). The final pellet was resuspended in 10 mL of 167 LB or glucose-based broth, whilst maintaining the required anaerobic conditions. Cultures were adjusted to an OD_{600 nm} of 1 (\pm 0.1) which equated to *ca*. 5.0 × 10⁸ CFU mL⁻¹. 168

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170 *2.3. Surface Topography*

171 The AM-G_Ms surface topography was measured using optical profilometry [39]. 172 Briefly, the surface roughness was quantitatively defined through the measurement of average 173 roughness (S_a), using a Zemetrics, Zegage 3D optical profiler (Zygo, USA) at × 50 174 magnification. The image analysis software used was Zemaps (Version 1.14.38). To increase 175 accuracy, only biological repeats where > 90 % of the surface was characterised by the software 176 were included in the surface analysis conducted (n = 9).

177

178 2.4. Scanning Electron Microscopy (SEM)

179 Scanning electron microscopy (SEM) was used to visualise the AM- G_M s and P. 180 aeruginosa strain ATCC 9027 retention and subsequent biofilm formation. Following the 181 termination of the MFC experimentation (after 120 h), the anode, cathode and cationic 182 exchange membrane (CEM) were isolated and dried at room temperature in a Class 2 biosafety 183 cabinet (Atlas Clean Air, UK). Samples were then prepared for SEM as described in [40] and 184 [41]. Briefly, the samples were fixed in 4 % gluteraldehyde for 24 h at 4 °C. The samples were then rinsed with sterile deionised water and were subjected to an ethanol gradient (10 % to 30 185 %, 50 %, 70 %, 90 % and 100 % v/v ethanol), samples were dried by desiccation over 24 h and 186 were sputter coated with gold (Polaron, UK) for 30 s (parameters: power 5 mA, 800 V, 30 s, 187 188 vacuum 0.09 mbar, argon gas). Scanning electron microscopy (SEM) was then performed 189 using a JEOL JSM-5600LV model SEM.

190

191 2.5. Biofilm Quantification

192 To quantify biofilm growth on the electrode surface in the MFC conditions utilised, a 193 crystal violet biofilm assay (CVBA) was conducted. These conditions consisted of LB broth, 194 incubated at 37 °C under static anaerobic conditions in an anaerobic cabinet (80% N₂ 10% H₂ 10% CO₂; Ruskin, Wales). Biofilm growth was determined at set time-points, namely 0 h, 1 h, 195 196 24 h, 48 h, 72 h, 96 h and 120 h. For P. aeruginosa biofilm formation, 1 mL of inoculated LB, with an OD_{600 nm} adjusted to 1.0 ± 0.1 was added to a 12-well, flat-bottomed culture plate, 197 198 under anaerobic conditions (n = 3). The AM-G_Ms were sterilised using 70 % ethanol (for 10 199 min) prior to experimentation and were added individually into the wells of a 12-well culture 200 plate, with the inoculated broth, the lid was added and sealed using Parafilm® to prevent 201 evaporation during incubation. The inoculated well plates were incubated at 37 °C under static 202 anaerobic conditions for the aforementioned time-points.

Following the incubation period, the LB broth was removed and discarded, the electrodes were dried at room temperature for 1 h in a Class 2 biosafety cabinet (Atlas Clean Air, UK). The CVBA experiments were conducted as described previously [18].

207 2.6. Pyocyanin Quantification – Liquid Chromatography Mass-Spectroscopy (LC-MS)

208 Pseudomonas aeruginosa strain ATCC 9027 growth kinetic assays were conducted 209 over 120 h in LB and glucose-based broth, in anaerobic and aerobic-to-anaerobic conditions. 210 At each time point (0 h, 24 h, 48 h, 72 h, 96 h and 120 h), 6.1 mL aliquots were removed, 5 211 mL was used for pyocyanin quantification, 1 mL for optical density measurements and 100 µL 212 for bacterial cell viability measurements. The 5 mL suspensions were centrifuged at 1,721 x g 213 for 10 min. A volume of 1 mL of the supernatant was added to 300 µL of methanol (with 0.1 214 M formic acid). Samples underwent further centrifugation (10,625 x g for 20 min) and 400 µL 215 of the supernatant was removed and added to 800 µL of sterile deionised water. Once the 216 samples were vortexed, 1 mL of the solution was filter sterilised using 0.22 µm filters 217 (Millex®, Ireland) and placed into 0.3 mL glass vials (Agilent, USA) for storage at -80 °C.

218 To quantify the production of the pyocyanin, standards of different pyocyanin 219 concentrations were prepared from High Performance Liquid Chromatography (HPLC) grade 220 pyocyanin (Merck, UK). One millilitre of methanol with 0.1 M formic acid was added to the 221 ampule of pyocyanin. This was vortexed (5 s), giving an initial concentration of 5 parts per 222 million (ppm). For the LC-MS experiments, 5 ppb and 50 ppb concentrations were prepared 223 from the above stock solution. The 5 ppb and 50 ppb were analysed alongside the LC-MS 224 experiments to provide a known standard baseline at two different concentrations, thus 225 enabling comparison of pyocyanin production between microbial samples.

226 An Agilent LC-MS 6540 UHD Q-TOF (Agilent, USA) with an ACE UltraCore 2.5 µm 227 SuperC18TM 100 mm x 2.1 mm column (Advanced Chromatography Technologies, Scotland) 228 was used for LC-MS analysis. The polar solvent used was LC-MS grade H₂O with 0.1 % formic 229 acid, whilst, acetonitrile with 0.1 % formic acid was the mobile phase solvent. The LC-MS 230 analysis was conducted in positive ion mode at 45 °C with a MS range of 100 - 3,000, which 231 was calibrated using four reference masses. A constant flow rate of 0.3 mL/min was maintained 232 within the column for 30 min and 10 µL samples were taken from each pre-prepared sample 233 vial.

The data generated from the LC-MS was multiple Agilent .D files. All the .D output files from the LC-MS were converted to an open source format (.mzXML) to facilitate further analysis, utilising the open source converter ProteoWizard MSConvert [42]. The peak-picking filter was chosen as this uses centroidisation, which facilitated the use of the feature detection algorithm, *centWave* to detect and compare close-by and partially overlapping features [43]. The .mzXML files were imported into MzMine2 for peak detection and analysis [44]. The m/z

240 peak for pyocyanin was confirmed in both the 5 ppm and 50 ppm reference samples as 211.086,

which corresponded to the monoisotopic peak of pyocyanin (210.079) with the addition of 241 242 [M+H]⁺ peak [45]. This reference mass was selected as the targeted peak to be used with the 243 peak detection function with the corresponding reported retention time. An intensity deviation tolerance of 50 %, noise level of 1.0×10^3 , m/z deviation of 5 ppm and retention time deviation 244 245 of 10 % were selected. To quantify the amount of pyocyanin per sample, the response factor 246 was used as described in [46], with the concentration for each sample determined by using the 247 average calculated response factor for the two standards. The relative response factor was 248 multiplied by 1000 to convert from mg mL⁻¹ to ppm. This value was divided by the calculated sample dilution factor for LC-MS preparation (which was 0.256) to determine the actual 249 pyocyanin concentration at different time-points, per sample. Pyocyanin production per CFU 250 251 mL⁻¹ was calculated, in order to determine if the growth media utilised resulted in a significant difference in pyocyanin production. 252

Datasets generated were analysed and the pyocyanin present quantified. Graphs were generated using the average sample concentration *via* GraphPad Prism (Version 8) and the standard error of the mean was notated using error bars.

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258 2.7. Microbial Fuel Cell Configuration

259 260 *2.7.1. MFC*

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0 2.7.1. MFC Construction

262 A classic H-shaped MFC device was utilised (Figure 1), which consisted of two microbial fuel cell glassware components (Adams & Chittenden Scientific Glass, USA), each 263 264 with a total volume of 120 mL. The two compartments were separated by a cationic exchange 265 membrane (CEM; CMI-7000, Membrane International, USA). Prior to use, all components 266 were sterilised with 70 % ethanol. The anodic chamber consisted of 120 mL LB or glucose-267 based broth, which was degassed with nitrogen to remove dissolve oxygen to generate 268 anaerobic conditions, prior to inoculation. An aliquot of 1 mL of prepared P. aeruginosa culture was added to the anodic compartment, resulting in a starting OD_{600 nm} of 0.01 (this 269 equated to ca. 5.0×10^6 CFU mL⁻¹). The cathodic compartment consisted of 120 mL, 50 mM 270 271 potassium hexacyanoferrate and 0.1 M potassium chloride (as the supporting electrolyte).

The AM- G_M s, AM-PLA and carbon cloth electrodes all have a geometric surface area of 7.5 cm² and were used (independently) for both the anode and cathode in each study. The electrodes were prepared by attaching the electrodes to 0.75 mm² stranded, tinned copper wire (Farnell, UK) *via* heat shrink tubing. The exposed surface between the electrodes and the heat shrink tubing was the sealed with electrical tape and epoxy resin (Araldite[®], Huntsman Advanced Materials, Switzerland). This ensured the isolation of the wire to prevent direct electron transfer by *P. aeruginosa* and minimised the risk of antimicrobial copper ions being produced. Therefore, only the calculated surface area was available to partake in electron transfer. Each MFC configuration tested was carried out in biological triplicates (n = 3).

281

282 2.7.2 Measurement and Analysis

The MFC configuration was connected to a high-resolution ADC-24 (Pico technology, UK) data acquisition system *via* an Arduino UNO (Arduino, Italy), which was coupled to a 10 K Ω external resistor. The closed circuit voltage (CCV) reading was recorded every 24 h, which resulted in voltage readings at 0 h, 24 h, 48 h, 72 h, 96 h and 120 h, over the 120 h incubation period, using Picolog software (version 6.1.10). The CCV recorded at each time-point was used as the Ecell (cell potential (V)) and by using Ohm's law, power [47] and current [48, 49] densities were calculated:

- 290
- 291
- 292 Power density:
- 293

Power Density (W m⁻²) =
$$\frac{(\text{E Cell (V)})^2}{\text{Resistance }(\Omega) \times \text{Surface Area }(m^{-2})}$$
 [1]

Current Density (A m⁻²) = $\frac{\text{E Cell (V)}}{\text{Resistance }(\Omega) \times \text{Surface Area }(m^{-2})}$

- 294
- 295 Current density:
- 296
- 297

298 2.7.3 Bacterial Enumeration

Bacterial viability was determined at each time point (0 h, 24 h, 48 h, 72 h, 96 h and 120 h) by Miles and Misra assays [50]. Colonies were counted and viability (CFU mL⁻¹) was determined to generate growth curves that could then be compared against the power/current densities generated. Enumeration of bacterial cells was carried out in biological triplicates (n = 3).

304

305 2.8. Statistical Analysis

Statistical analysis was conducted by performing two-way ANOVA coupled with
 Tukey's multiple comparison tests for post hoc analysis using GraphPad Prism (version 8.4.2;
 GraphPad Software, USA) to determine significant differences at a confidence level of 95 %

[2]

309 (p < 0.05). Error bars represent the standard error of the mean. Asterisks denote significance, 310 $*(p \le 0.05), **(p \le 0.01), ***(p \le 0.001)$ and $****(p \le 0.0001)$.

311

312 **3.0. Results**

313 3.1. Surface Topography Characterisation

314 The surface morphology of the AM-G_Ms was examined using SEM (Figure 2.A.). At a 315 relatively lower magnification (100×) the surface topography of AM-G_Ms was revealed, a 316 highly heterogenous topography was observed due to the striations created during the 317 manufacturing process. In light of this, optical profilometry was used to further characterise 318 the topographical features of both sides of the AM-G_Ms (Figure 2.B.). Average surface 319 roughness of the additive manufactured electrodes revealed a difference in surface topography 320 when comparing the top and the underside of the AM-G_Ms and the control, AM-PLA. The 321 topside of the AM-G_Ms surface produced an average roughness of 4.40 µm, compared to the 322 underside of the same electrode (2.65 μ m). This was due to the manufacturing process as the 323 underside of the electrode was extruded on to the smooth flat bed of the 3D printer. Both sides 324 of the AM-G_Ms electrode exhibited a significantly greater surface roughness, than AM-PLA 325 (containing no graphene). Only samples that had > 90 % of the surface analysed were included 326 in this study to enhance accuracy and reliaility.

327

328 3.2. Biofilm Quantification

329 In order to quantify biofilm growth, biofilm studies were run simultaneously with the 330 MFC experimentation, which allowed biofilm growth on the electrodes to be studied without 331 disturbing the potential difference produced in the MFC experiments (Figure 3). To assess 332 biofilm formation P. aeruginosa strain ATCC 9027 was inoculated in anaerobic LB broth, as 333 per the MFC experiment protocol. Both the AM-G_Ms and AM-PLA electrodes were evaluated 334 after the following time-points, 0 h, 1 h, 24 h, 48 h, 72 h, 96 h and 120 h. At each time-point, 335 greater bacterial biomass was observed on the AM-G_{Ms}, which was significant at 24 h ($p \le$ 0.0001), 48 h ($p \le 0.0001$), 96 h ($p \le 0.001$) and 120 h ($p \le 0.01$) compared with the AM-PLA. 336 337 A maximum biomass was recorded on the AM-G_Ms at 48 h. However, after 48 h, a decrease in 338 biomass was observed on both the AM-G_Ms and AM-PLA electrodes respectively. The presence 339 of graphene in the AM-G_Ms enhanced *P. aeruginosa* biofilm formation when cultured in LB 340 broth, under anaerobic conditions.



343 Pyocyanin production was quantified in parallel with the bacterial growth kinetics 344 experiment, and samples were prepared after each time-point for LC-MS analysis (Figure 4). 345 Overall, the results demonstrated little difference in the quantity of pyocyanin present when P. 346 aeruginosa strain ATCC 9027 was grown in different growth media, namely, glucose-based 347 and LB broth P. aeruginosa strain ATCC 9027 in anaerobic conditions (Figure 4.A.). The maximum amount of pyocyanin produced by P. aeruginosa strain ATCC 9027 in LB broth 348 349 was 43.50 ppb after 24 h, and the maximum amount of pyocyanin produced by *P. aeruginosa* 350 strain ATCC 9027 in glucose-based broth was 74.20 ppb after 0 h. However, no significant 351 difference in pyocyanin production was observed when growth was conducted in glucose-352 based and LB broth.

353 Bacterial viability was calculated in colony forming units per mL (CFU mL⁻¹) (Figure 354 4.B.). To quantify the amount of *P. aeruginosa* strain ATCC 9027 present at set time-points 355 bacteria were grown in the different growth media, namely, glucose-based and LB broth in 356 anaerobic conditions. Anaerobic growth in both glucose-based broth and LB broth, resulted in 357 traditional growth phase models, including the presence of lag phase, exponential growth (log 358 phase), stationary phase and death phase. In the presence of LB medium, *P. aeruginosa* grew 359 to a maximum of 2.94×10^7 CFU mL⁻¹, whilst, the highest recorded viability for an anaerobic *P. aeruginosa* culture in glucose-based medium was 1.3×10^7 CFU mL⁻¹. 360

The concentration of pyocyanin was then determined as a result of pyocyanin produced per CFU mL⁻¹ (Figure 4.C.). A significant increase in pyocyanin production was recorded in glucose-based broth at 2 h ($p \le 0.01$) and 120 h ($p \le 0.0001$) when compared to LB broth at the respective incubation time points.

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366

367 *3.4. Microbial Fuel Cell Analysis*

368 Throughout MFC experimentation, the closed circuit voltage of the three electrode types (AM-G_Ms, AM-PLA and carbon cloth) was recorded at 0 h, 24 h, 48 h, 72 h, 96 h and 120 369 370 h in two media types, anaerobic glucose-based and anaerobic LB (Figure 5.A.). The AM-PLA 371 confirmed that the base material (with no graphene incorporated) demonstrated little conductivity. Both the AM-G_Ms and carbon cloth electrodes demonstrated conductivity from 0 372 373 h to 24 h, which was monitored as an increase in cell potential. This was followed by a 374 stationary period and finally from 72 h to 120 h a decrease in cell potential was observed. 375 Interestingly, this correlated with the kinetics of bacterial viability, which were calculated at 376 the same time-points (Figure 5.B.). The highest potential recorded was when P. aeruginosa 377strain ATCC 9027 was inoculated with carbon cloth in anaerobic glucose-based medium after37896 h ($324.00 \pm 10.06 \text{ mV}$). The highest potential recorded from the AM-G_{MS} was in LB medium379after 48 h ($287.00 \pm 18.52 \text{ mV}$). However, there was no statistical difference between the AM-380G_{MS} in varying media.

381 During MFC experimentation, growth kinetic experiments were used to determine the 382 viability of P. aeruginosa strain ATCC 9027 from MFCs over defined time-points (0 h, 24 h, 383 48 h, 72 h, 96 h and 120 h) (Figure 5.B.). All six conditions (three electrode materials × two 384 media sources) resulted in conventional bacterial growth phase models, including the presence 385 of lag phase, exponential growth (log phase), stationary phase and death phase. The maximum viability was observed when P. aeruginosa strain ATCC 9027 was inoculated in anaerobic LB 386 medium with carbon cloth electrodes after 72 h (6.94×10^7 CFU mL⁻¹). The maximum viability 387 388 of P. aeruginosa strain ATCC 9027 in the presence of the AM-G_Ms throughout this study was reported in anaerobic glucose-based broth after 96 h incubation (4.50×10^7 CFU mL⁻¹). There 389 390 was no significant difference (p > 0.05) in cell viability when *P. aeruginosa* strain ATCC 9027 was grown in the presence of AM-G_Ms, AM-PLA and carbon cloth electrodes in both glucose-391 392 based and LB broth, which indicated that the incorporation of graphene in the AM-G_Ms resulted 393 in no significant detrimental antimicrobial effect.

394 Power density was calculated from the cell potential values obtained during MFC 395 experimentation using equation [1] (Figure 5.C). In the presence of *P. aeruginosa* strain ATCC 396 9027, the AM-PLA (no graphene) provided low power densities in both anaerobic glucosebased and LB broth (< 1 μ W m⁻²). The AM-G_Ms and the carbon cloth electrodes recorded an 397 increase in power density until 24 h, followed by a stationary period and then a decrease in 398 399 power density from 96 h. The highest recorded power density from the electrodes used in this study was $110.74 \pm 14.63 \ \mu\text{W} \text{ m}^{-2}$, which was obtained from the AM-G_Ms in LB medium after 400 48 h incubation. The highest recorded power density from the carbon cloth electrode was 93.49 401 \pm 5.17 µW m⁻² after 96 h incubation in anaerobic glucose-based broth. Statistical analysis 402 revealed at 48 h that the AM-G_Ms in LB broth (110.74 \pm 14.63 μ W m⁻²) outperformed (P < 403 404 0.05) the carbon cloth electrode in anaerobic glucose-based broth (58.86 \pm 10.84 μ W m⁻²). 405 However, after 72 h ($p \le 0.05$), 96 h ($p \le 0.05$) and 120 h ($p \le 0.01$) incubation, the carbon 406 cloth electrode gave a greater current density than AM-G_{Ms} in anaerobic glucose-based 407 medium. This demonstrated that the carbon cloth electrodes in glucose-based broth were the 408 most efficient electrode after 72 h incubation. There was no significant difference in power 409 density between the AM-G_Ms and carbon cloth electrodes when *P. aeruginosa* was incubated 410 for 120 h in anaerobic LB broth.

411 Current density was also calculated from the MFCs, via the recorded potential values 412 and equation [2] (Figure 5.D.). The highest current density achieved in this study was 382.67 \pm 24.69 µA m⁻²; this was calculated from the AM-G_Ms in anaerobic LB broth after 48 h 413 414 incubation with P. aeruginosa strain ATCC 9027. The highest current density achieved by the same electrode in anaerobic glucose-based broth was $284.00 \pm 10.36 \ \mu A \ m^{-2}$ after 24 h. The 415 highest current density achieved by the carbon cloth electrode was $343.11 \pm 6.94 \ \mu A \ m^{-2}$ after 416 417 120 h incubation with P. aeruginosa stain ATCC 9027 in anaerobic LB medium. No significant statistical difference in current density was observed when P. aeruginosa was incubated over 418 419 120 h, in anaerobic LB broth and anaerobic glucose-based medium, in the presence of the AM-420 G_{Ms} and carbon cloth electrodes. Therefore, both of these electrode materials showed 421 comparable current and current density generation in a MFC configuration.

422

423 3.5. Bacterial Visualisation via SEM

424 Following 120 h incubation, the anode, cathode and the CEM surface, which was in 425 contact the anolyte, were isolated and visualised via SEM (Figure 6). Visualisation of P. 426 *aeruginosa* on the surface of the AM-G_Ms in LB medium showed a sparsely populated surface 427 (Figure 6.A.) when compared with the AM-PLA, which showed denser bacterial retention 428 (Figure 6.D.). The cathode (AM-G_{MS}) in anaerobic LB broth showed no bacterial retention, 429 indicating no bacterial transfer between the two chambers of the MFC (Figure 6.B.). The 430 surface of the CEM isolated from the anodic chamber showed an abundance of *P. aeruginosa* 431 retention, which was increased around surface defects (Figure 6.C.). The AM-G_Ms in glucose-432 based medium (Figure 6.E.) demonstrated a thin and evenly distributed bacterial retention 433 across the surface of the electrode. The AM-PLA electrode in glucose-based broth (Figure 6.F.) 434 demonstrated enhanced bacterial retention around surface defects after 120 h of anaerobic 435 incubation. This variation in bacterial retention may be due to the presence of graphene on the 436 surface of the AM-G_Ms.

437

441

438 **4.0. Discussion**

- 439 *4.1.AM-G_Ms Characterisation*
- 440 4.1.1. Electrochemical Analysis of AM-G_{MS}

442 The AM-G_Ms utilised throughout this study have previously been electrochemically 443 characterised [13]. The heterogeneous rate transfer constants (k^0_{obs}), were found to correspond 444 to 1.00×10^{-3} cm s⁻¹ and 4.58×10^{-4} cm s⁻¹ for the PLA / graphene filaments (prior to AM) and the AM-G_Ms, respectively, within a 1 mM hexaammineruthenium (III) chloride/0.1 M KCl solution [13]. The PLA / graphene composite filament exhibited faster electron kinetics than AM-G_Ms, this could be due to improved percolation of agglomeration of graphene platelets in the filament prior to printing, once printed the dispersion of the graphene platelets improved, resulting in reduced percolation [13].

450

451 *4.1.2. Surface Characterisation*

452 A vital parameter to be considered when attempting to understand bacterial-surface 453 retention is surface topography [51]. The size and shape of morphological features of a surface 454 can directly influence bacterial retention [52]. Previous studies indicate that an increase in 455 surface topography results in enhance bacterial retention and may also facilitate biofilm 456 formation [53, 54]. Scanning electron microscopy of the topside of the AM-G_Ms at \times 100 457 magnification revealed a striated surface morphology due to the FDM process; the circular 458 cross-section of the extrudate from the print head, combined with relatively high melt viscosity 459 and solidification on cooling, prevents melding into a microscopically flat surface. Instead a 460 "log cabin texture" is produced, resulting in pits and grooves at a macro-scale level and 461 enhanced bacterial retention, resulting in enhanced power outputs from the electrodes in an 462 MFC configuration [55, 56]. The graphene aggregates in the PLA based composite led to local 463 variations in solidification shrinkage and increased roughness of the AM-G_{MS} surface relative 464 to the AM-PLA surfaces (which in comparison demonstrated a smoother surface topography). The 465 micro-roughnesss on the striations in Figure 2 A is due to the graphene aggregates.

466 To further explore surface topography, both sides of the AM-G_Ms and the AM-PLA 467 electrodes were evaluated *via* optical profilometry and S_a (average surface roughness) values 468 were recorded. Due to the fibrous nature of carbon cloth, surface characterisation by optical 469 profilometry was not viable on this electrode material. The S_a values recorded revealed that 470 that the topside of the AM-G_Ms and AM-PLA electrodes resulted in a greater surface roughness 471 when compared with the underside of the electrodes. The average surface roughness (S_a) of the AM-G_Ms topside was 4.40 µm and for the underside it was 2.65 µm. The cell dimensions of *P*. 472 473 aeruginosa are in the region of 0.5 µm - 1.0 µm (diameter) and 1.5 µm - 5.0 µm (length), 474 therefore this difference in surface roughness could have a significant effect on bacterial 475 retention [57, 58]. The optical profilometry results demonstrated that the additive 476 manufacturing process did not yield homogeneous surfaces, the observed smoother underside 477 of the electrodes was due to this surface being the first layer of the "print", extruded onto the 478 smooth flat bed of the 3D-printer [59], the extruded filament melt will mould to the smooth bed surface before solidification, resulting in a smoother surface overall. The roughness valuesobtained in this study are in line with previously published literature [60].

481 The AM-G_Ms electrodes have previously been characterised using Raman spectroscopy 482 by [61]. The Raman spectra of the AM-G_Ms feature the characteristic carbonaceous peaks, 483 commonly referred to as the G and 2D (G') bands at ca. 1580 cm⁻¹ and 2690 cm⁻¹, respectively 484 [62-64]. The graphene content of the AM-G_Ms comprised of multi-layer graphene sheets, 485 determined from the highly symmetrical 2D (G') band peak. Furthermore, there was a distinct 486 lack of a characteristic shoulder on the 2D(G') band, which is commonly observed in graphitic 487 materials [63, 65]. Whilst, the large D band was indicative of a high number of edge plane-like 488 sites [66] which give rise to beneficial electron transfer sites, making it a useful electrode 489 material.

490

491 4.2. Biofilm Quantification

492 Biofilm quantification was evaluated using predetermined MFC conditions (P. 493 aeruginosa ATCC 9027 inoculated into LB broth at 37 °C under static, anaerobic conditions). 494 *Pseudomonas aeruginosa* biofilm formation on the AM- G_M s was evaluated over 0 h – 120 h in 495 anaerobic LB broth to replicate MFC conditions. Over the 120 h incubation period, greater 496 biofilm formation was consistently evident on the AM-G_{MS} when compared to the AM-PLA 497 counterparts. A peak in biomass was observed at 48 h and 72 h for the AM-G_Ms and AM-PLA 498 electrodes, respectively. The peaks were followed by a reduction in biomass, this could be due 499 to biofilm detachment mechanisms which can be induced by environmental cues such as 500 nutrient, pH and oxygen availability [67]. Therefore, the reduction in biofilm formation during 501 the latter stages of the incubation period could be due to the anaerobic, nutrient-limited 502 conditions that were utilised.

503

504 4.3. Pyocyanin Quantification – Liquid Chromatography-Mass Spectroscopy (LC-MS)

505 Pyocyanin production was quantified via LC-MS in parallel with each time-point of the growth dynamic experiment. The utilisation of different growth media can result in the 506 507 production of different metabolites at varying concentrations [68], therefore LC-MS analysis 508 was conducted on *P. aeruginosa* strain ATCC 9027 samples over 120 h in anaerobic LB broth 509 and glucose-based broth. P. aeruginosa exhibited a significantly greater viability when 510 inoculated in LB medium, compared to glucose-based broth under anaerobic conditions. Under 511 anaerobic conditions, P. aeruginosa survival and proliferation is supported by mechanisms 512 including denitrification, pyruvate fermentation and arginine fermentation [69, 70]. As P.

513 aeruginosa has no known nitrogen fixing ability, denitrification is not a viable survival 514 pathway in this instance, however, it is potentially the fermentation of amino acids such as 515 arginine that support proliferation in anaerobic LB medium [71]. To metabolise glucose under 516 anaerobic conditions the Entner-Doudoroff pathway is used by *P. aeruginosa*, which converts 517 one glucose molecule into two pyruvate molecules, resulting in the production of one ATP 518 molecule and two excess reducing equivalents [72, 73]. The pyruvate can be further converted 519 to acetate, producing additional ATP [73]. The results showed little variation in the 520 concentration of pyocyanin produced by P. aeruginosa strain ATCC 9027 in the presence of 521 anaerobic LB broth and glucose-based broth. The maximum concentration of pyocyanin in 522 anaerobic glucose-based broth was 74.20 ppb at 0 h, whilst in anaerobic LB broth it was 43.50 ppb after 48 h incubation. Pyocyanin production was then quantified in regards to CFU mL⁻¹ 523 524 obtained at each timepoint. A significant increase in pyocyanin production was recorded in 525 glucose-based broth at 2 h and 120 h when compared to LB broth. This indicated that the 526 glucose-based medium promoted the production of the electron shuttle, pyocyanin, when 527 compared against LB broth, which could lead to further biological-based optimisation of MFC 528 configurations in the future [6, 68, 74]. Throughout this study, pyocyanin accumulation was 529 not evident over time. The fluctuations observed in the concentrations of pyocyanin may be a 530 consequence of redox cycling, which can generate reactive oxygen species (ROS), such as 531 H_2O_2 , which can degrade pyocyanin [75, 76].

532

533 4.4. Microbial Fuel Cell Analysis

534 The AM-G_Ms was applied in a MFC configuration, as PLA has excellent electrical insulation properties at < 70 °C [77, 78], another conductive control electrode, carbon cloth, 535 536 was utilised for appropriate comparisons of the AM-G_Ms to be made. Carbon cloth is widely 537 applied as an electrode in MFC configurations due to its high conductivity, good 538 biocompatibility and relatively large surface area [21]. However, it is excessively expensive as 539 an electrode material in MFC configurations [21]. When inoculated with P. aeruginosa strain 540 ATCC 9027, the AM-G_Ms demonstrated excellent conductivity, producing a maximum 541 potential (287 ± 18.52 mV). Carbon cloth produced a maximum potential in anaerobic-based 542 glucose-based broth (324 \pm 10.06 mV) whilst, the AM-PLA, which demonstrated limited 543 conductivity, produced a maximum potential of $(32 \pm 1.01 \text{ mV})$.

In parallel to the cell potential recording every 24 h, a sample of the anolyte was taken (immediately after the cell potential recording, maintaining anaerobic conditions) and used to calculate cell viability. In all cases (three electrode materials in two types of media), *P*. 547 aeruginosa demonstrated typical growth kinetics. Interestingly, throughout this study there was 548 no significant difference in cell viability between bacteria exposed to the AM-G_{MS}, AM-PLA 549 and carbon cloth electrodes. This indicated that the presence of graphene within the AM-G_{MS} 550 did not have a detrimental effect on *P. aeruginosa* viability. This appears to be contradictory 551 to the reported controversial antimicrobial activity of graphene which is yet to be fully 552 elucidated [79, 80]. Potential antimicrobial mechanisms of the graphene such as membrane 553 perturbation, ROS generation and cellular wrapping are thought to be involved [79, 81, 82]. 554 However, in this study, the AM-G_Ms containing 8 wt.% graphene demonstrated no significant 555 detrimental activity on P. aeruginosa viability.

556 Power and current densities were calculated using the recorded cell potential values 557 from the MFCs. The highest current and power densities generated were $110.74 \pm 14.63 \ \mu W$ m⁻² and $382.67 \pm 24.69 \,\mu\text{A}$ m⁻², respectively. These values were calculated from *P. aeruginosa* 558 strain ATCC 9027 in the presence of the AM-G_Ms after 48 h anaerobic incubation in LB 559 560 medium, under a fixed external resistor of 10 k Ω . The highest recorded power density was demonstrated by the carbon cloth electrode (93.49 \pm 5.17 μ W m⁻² after 96 h) in anaerobic 561 glucose-based broth. The highest recorded current density was $343.11 \pm 6.94 \ \mu A \ m^{-2}$ after 120 562 563 h incubation with P. aeruginosa stain ATCC 9027 in anaerobic LB medium. In a previous 564 study that used a comparable MFC configuration, Jayapriya and Ramamurthy (2014), 565 inoculated the anodic chamber with P. aeruginosa in a glucose-based medium (20 g peptone, 566 13.3 g glucose, 10 g potassium sulphate, 1.4 g magnesium chloride, per litre (pH 7.0)). 567 Potassium ferricyanide was used as the catholyte and the resistor was fixed in the ohmic 568 polarisation region. When carbon cloth was applied as both the anode and cathode in this configuration, a power density of 132.9 μ W m⁻² was recorded [83]. This indicates that the novel 569 570 AM-G_{MS} utilised throughout this study demonstrated results that were comparable of that to the 571 current benchmark electrode utilised in MFC configurations. As research in AM continues to 572 expand, electrodes produced by this manufacturing technique will continue to become more 573 efficient and less expensive [84]. One route to optimise the AM-G_Ms to enhance power outputs in 574 MFC configurations could be to increase the graphene concentration present. In a previous study 575 Foster et al., (2019) developed a bespoke PLA / graphene composite FDM filament with graphene 576 content varying from 1 - 40 wt.% [85].

577 There was no significant difference in recorded power and current densities between 578 the MFCs which utilised the AM- G_{MS} , compared with the more widely used carbon cloth 579 electrode type, when used in the same medium over 120 h. The main restriction of MFCs for 580 industrial applications is their low power densities (compared to other energy sources). However, these results demonstrated that the novel utilisation of additive manufactured electrodes coupled with the addition of graphene can produce comparative outputs to the current 'gold' standard and with future research could enhance power outputs of MFC technologies [86].

585

586 4.5. Bacterial Visualisation via SEM

587 The anode, cathode and CEM were isolated after 120 h incubation and prepared for 588 SEM in order to visualise *P. aeruginosa* retention to the MFC components. When the AM-G_{MS} 589 were applied as the anode, in both anaerobic LB broth and glucose-based broth, the bacterial 590 cell retention was sparse (when compared to the AM-PLA counterparts). This difference in cell 591 retention may be due to the incorporated graphene, potentially due to modifications in surface 592 energy and electrostatic interactions [87]. Biofilm composition, morphology, thickness and 593 other physical properties play an integral role in bioelectricity production [88]. An increase in 594 biofilm formation on the anodic surface has previously been shown to be directly proportional 595 to MFC power outputs [89]. In light of this, in future studies the graphene content and surface 596 properties could be modified (via AM parameters) and this could result in enhanced power and 597 current densities.

598

599

601 **5.0. Conclusions**

602 Additive manufactured PLA/graphene (8 wt.%) composite electrodes (AM-G_{MS}) have 603 been used in MFC applications for the first time. A commercial PLA/graphene (8 wt.%) 604 composite FDM filament was used throughout the study. Raman Spectroscopy revealed that 605 the AM-G_Ms were comprised of multi-layered graphene. Optical profilometry revealed that the 606 electrode surface in contact with the printer bed had a smoother topography than the top 607 surface. The AM-G_Ms had significantly higher surface roughness values when compared to the 608 AM-PLA (control). Growth kinetics were measured to assess Pseudomonas aeruginosa 609 viability in the presence of the electrodes and no significant detrimental activity was observed.

610 For the first time, AM-G_Ms containing 8 wt.% graphene were utilised in a *P. aeruginosa* 611 inoculated MFCs (using the conditions previously characterised), in two media types, 612 anaerobic LB and anaerobic glucose-based broth. Over the 120 h incubation period the AM-613 G_Ms exhibited good electrical conductivity and produced results comparable to that of a widely 614 utilised 'benchmark' electrode material (carbon cloth). When inoculated in LB broth, in a P. 615 aeruginosa MFC, the AM-G_{MS} produced power and current densities comparable to that of the 616 current benchmark electrode of MFCs, carbon cloth, whilst being considerably less expensive and 617 a more sustainable, environmental-friendly electrode material. Following MFC experimentation, 618 SEM was conducted on the electrodes, whilst bacterial retention was sparse this did not affect 619 the power outputs observed, and no detrimental antimicrobial effect due to the presence of 620 graphene on the surface of the AM-G_Ms was observed. In future research the surface profiles / 621 morphologies of the AM-G_Ms electrodes at the nano, micro and macro-scales could be optimised. 622 Additionally, wastewater could be trialled as the anolyte AM-G_Ms to further translate results to an 623 industrial application. This study demonstrated the potential of additive manufacturing, coupled 624 with the incorporation of nanomaterials for the fabrication of electrodes, to ultimately increase 625 power outputs of MFCs.

626

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644

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5.3. Author Contributions

650	K.A.W, C.E.B and A.J.S conceptualised the project. A.J.S., N.A.H., J.A.B. and C.M.L. were
651	involved in data acquisition and analysis. D.W. designed and fabricated the voltammeter
652	configuration used to monitor power outputs over time. K.A.W. and C.E.B. supervised the
653	project and supplied the relevant materials. A.J.S drafted the final manuscript. All authors read
654	and approved the final manuscript.
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658	

660 Figure 1. The two-chambered, classical H-shaped MFC utilised in this study. K₃Fe(CN)₆,

661 Potassium hexacyanoferrate (III), KCl; Potassium chloride, CEM; Cationic Exchange662 Membrane, Polytetrafluoroethylene; PTFE.



- Figure 2. A. SEM image depicting the surface morphology of the AM-G_{MS} at 100 × magnification. B. Average surface roughness (S_a) of both sides of the AM-G_{MS} and the AM-PLA control (n = 9). Asterisks denote statistical significance, * ($p \le 0.05$), ** ($p \le 0.01$), and **** ($p \le 0.0001$).
- 674



Figure 3. Biofilm quantification was determined using CVBA following *P. aeruginosa* strain ATCC 9027 incubation over 120 h in LB broth (deducting the sterile broth controls). Conditions replicated those selected for MFC experimentation (incubation conditions: static, LB broth, 37 °C, anaerobic) (n = 3). Asterisks denote significance, **($p \le 0.01$), ***($p \le 0.001$) and ****p (≤ 0.0001).



Figure 4. A. Growth kinetics were recorded by sampling *P. aeruginosa* strain ATCC 9027 over 120 h and conducting bacterial viability calculations, using two media types, glucosebased and LB broth (incubation conditions; anaerobic; 37 °C; static). **B.** Pyocyanin quantification *via* LC-MS from *P. aeruginosa* strain ATCC 9027 over 120 h **C.** Pyocyanin production normalised using growth kinetic data (CFU mL⁻¹) (n = 3). Where error bars are not visible they are of similar or smaller size compared to the symbol. Asterisks denote significance,*($p \le 0.05$), **($p \le 0.01$) and ****($p \le 0.0001$).

694



697 Figure 5. Microbial fuel cell configurations were trialled with the AM-G_Ms inoculated with anaerobic *P. aeruginosa* strain ATCC 9027 MFCs (n = 3). A. Cell potential (V). B. Growth 698 kinetics (CFU mL⁻¹). C. Power density (μ W m⁻²). D. Current density (μ A m⁻²). For the 699 700 statistical analysis, only results between the AM-G_Ms and carbon cloth electrodes were 701 analysed, the black solid line denoted the different electrodes in glucose-based medium, the 702 black dashed line denoted AM-G_Ms in glucose-based broth against carbon cloth in LB broth, 703 the red dashed line denoted the AM-G_Ms in LB broth against carbon cloth in glucose-based medium. Asterisks denote significance, $*(p \le 0.05)$, $**(p \le 0.01)$, $***(p \le 0.001)$ and $****(p \le 0.001)$ 704 705 \leq 0.0001).



- 709 Figure 6. The electrodes and CEM were isolated post-MFC experimentation and *P. aeruginosa*
- 710 strain ATCC 9027 was visualised using SEM (at 5.00 K × magnification). A. AM-G_Ms anode
- 711 incubation in LB broth. **B.** AM-G_Ms cathode (no bacteria present). **C**. CEM in LB broth (surface
- 712 from the anodic compartment visualised). **D.** AM-PLA anode incubation in LB broth. **E.** AM-
- 713 G_Ms anode incubation in glucose-based medium. **F.** CEM in glucose-based broth (surface from
- 714 the anodic compartment visualised).



717 **6.0. References**

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