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Cell and Protein Compatibility of Parylene-C Surfaces

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Parylene-C, which is traditionally used to coat implantable devices, has emerged as a promising material to generate miniaturized devices due to its unique mechanical properties and inertness. In this paper we compared the surface properties and cell and protein compatibility of parylene-C relative to other commonly used BioMEMS materials. We evaluated the surface hydrophobicity and roughness of parylene-C and compared these results to those of tissue culture-treated polystyrene, poly(dimethylsiloxane) (PDMS), and glass. We also treated parylene-C and PDMS with air plasma, and coated the surfaces with fibronectin to demonstrate that biochemical treatments modify the surface properties of parylene-C. Although plasma treatment caused both parylene-C and PDMS to become hydrophilic, only parylene-C substrates retained their hydrophilic properties over time. Furthermore, parylene-C substrates display a higher degree of nanoscale surface roughness (>20 nm) than the other substrates. We also examined the level of BSA and IgG protein adsorption on various surfaces and found that surface plasma treatment decreased the degree of protein adsorption on both PDMS and parylene-C substrates. After testing the degree of cell adhesion and spreading of two mammalian cell types, NIH-3T3 fibroblasts and AML-12 hepatocytes, we found that the adhesion of both cell types to surface-treated parylene-C variants were comparable to standard tissue culture substrates, such as polystyrene. Overall, these results indicate that parylene-C, along with its surface-treated variants, could potentially be a useful material for fabricating cell-based microdevices.

1. Introduction

Polymeric biomaterials are widely used in therapeutics^{1,2} and diagnostics^{3,4} as micro- and nanobiosensors for cell-based assays, drug delivery, and tissue-engineering applications.⁵ Polymeric microdevices are capable for analyzing cells and proteins,^{6–8}

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generating tissue-engineering scaffolds, $^{9-11}$ and miniaturizing 35bioassays for high-throughput experimentation.¹² With the recent 36 emergence of soft lithography, elastomers, such as poly-37 (dimethylsiloxane) (PDMS), have become enabling materials 38 for the widespread fabrication and the use of microfabricated 39 systems. PDMS offers numerous advantages over traditional 40 biomaterials. It is relatively inexpensive, inert, nontoxic, and 41 can be easily molded to form microstructures.¹³ Despite these 42 desirable characteristics, PDMS has a number of shortcomings. 43For example, although PDMS has been shown to be compatible 44 for short-term culturing of cells,¹⁴ little is known of its long-term 45stability in tissue-engineering applications and in vivo diagnostics. 46 Therefore, it may be important to explore alternative biomaterials 47 that can be used to fabricate biomedical microdevices. Poly-48

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49 (chloro-*p*-xylylene), also referred to as parylene-C, is one such
 50 potential candidate for fabricating biomedical devices.

Parylene-C is a thermoplastic, crystalline, and transparent 51polymer that is extensively used as a coating for insulating 52implantable biomedical devices.¹⁵ In addition, parylene-C is 53chemically inert and nonbiodegradable. Parylene-C is synthesized 54from a low-molecular weight (MW) dimer, dichloro-di(p-5556 xylylene), using a process that involves the decomposition of 57*p*-xylylene to yield chloro-*p*-xylylene, followed by the polymerization of chloro-p-xylylene to parylene-C.¹⁶ Parylene-C can 58 be vapor-deposited onto substrates to generate uniform, pinhole-59 free membranes that can be subsequently dry-etched using oxygen 60 plasma to yield microscale features and patterns that are ideal 61 for culturing cells.¹⁷ The all-carbon structural backbone, high-62 MW, and nonpolar entities make parylene-C highly resistant to 63 most chemicals, as well as to fungal and bacterial growth. In 64 addition to having conducive biochemical properties, parylene-C 65 has a Young's modulus of ~ 4 GPa¹⁸ (compared to 0.75 MPa for 66 PDMS¹⁴)-making it mechanically robust and highly suitable 67 for fabricating stable and reusable microfluidic devices or 68 stencils.¹⁷⁻²² Recent studies have shown parylene-C to be more 69 70 hemocompatible and less thrombogenic than silicon.²³ Parylene-C 71 has also demonstrated high stability in vivo for a variety of applications, such as cardiovascular implants.^{24,25} Furthermore, 72parylene-C is a potentially useful material for in vitro cell culture 73 studies. For example, we have developed the use of parylene-C 74stencils for patterning cells and proteins and for generation of 75cocultures with control over the degree of homotypic and 76 heterotypic cell-cell interactions.^{26,27} Another recent study 77 provides the methodology for making nanoscale sculptured thin 78 film (STF) out of parylene-C.²⁸ Due to the high surface area to 79 volume ratio of the STF, the parylene-C STF supports high level 80 of cell adhesion.²⁸ However, despite the apparent biocompatibility 81 of parylene-C, there has been no direct comparison of parylene-C 82 83 to PDMS and other materials commonly used in BioMEMS.

84 In this study, we compared the biocompatibility of parylene-C membranes with PDMS, glass, and optically clear virgin 85 86 polystyrene by analyzing protein adsorption, cell adhesion, and 87 cell morphology characteristics on each of these surfaces. In addition, we treated parylene-C and PDMS with air plasma and 88 89 coated the surfaces of these substrates with fibronectin to study 90 the effects of surface treatments on protein adsorption, cell adhesion, and spreading. Protein adsorption was studied using 91

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bovine serum albumin (BSA) and immunoglobulin G (IgG), and 92 cell adhesion and spreading were studied using NIH-3T3 93 fibroblast and AML-12 hepatocyte cell lines. 94

2. Methods and Materials

2.1. Fabrication of Parylene-C and PDMS. Three inch silicon 96 wafers were first cleaned for ~ 10 min using a 1:1 piranha solution 97 (equal volume mixture of H_2SO_4 and H_2O_2), sufficiently rinsed with 98 deionized water, nitrogen-dried, and then coated with hexameth-99 yldisilazane (HMDS). Following pretreatment, the silicon wafers 100 were deposited with dichloro-di(p-xylylene) by utilizing a Labcoater 101 2 PDS 2010 chemical deposition system (Specialty Coating Systems, 102 Indianapolis). Inside the deposition system, dichloro-di(*p*-xylylene) 103 is first vaporized at 150 °C and 1 torr and then pyrolyzed at 690 °C 104 and 0.5 torr to form chloro-p-xylylene-the monomer of parylene-105 C. A reduction in the chamber temperature causes chloro-p-xylylene 106 to condense onto the wafer surfaces to form parylene-C membranes. 107 Initial loading of dichloro-di(p-xylylene) onto the silicon wafers 108 determines the thickness of the parylene-C membrane at a rate of 109 $0.5 \,\mu\text{m/g}$. With the use of the aforementioned protocol, $10 \,\mu\text{m}$ thick 110 parylene-C membranes were fabricated on silicon substrates. 111

The PDMS substrates were fabricated by directly curing a Sylgard112184 (Essex Chemical) elastomer in the wells of a Costar 24-well113TC-treated cell culture microplate for nearly 2 h at 70 °C, using a11410:1 weight ratio of elastomer to curing agent.115

2.2. Preparation of Surfaces. A total of eight types of surfaces 116 were used in this study. Costar 24-well TC-treated cell culture 117 microplates were utilized as optically clear virgin polystyrene 118 substrates. Parylene-C experimental samples were prepared by 119 carefully cutting the 10 μ m thick parylene-C membranes (section 120 2.1) to form square-shaped pieces of \sim 5 mm \times 5 mm. Each cut-out 121 piece of parylene-C was placed and sealed reversibly onto a PDMS-122 coated well in the microplate. Platinum glass coverslips, $18 \text{ mm} \times$ 123 18 mm in size, were used as the glass samples. Plasma-treated PDMS 124and parylene-C were obtained by treating the two polymers with air 125plasma in a Harrick PDC-001 plasma treatment chamber for 2 min. 126 Protein coating to parylene-C and PDMS surfaces was performed 127 by simply incubating a 5 μ g/mL fibronectin solution on the surfaces 128for 1 h. 129

Each substrate was sterilized prior to the experiments. The 130 sterilization of plain and plasma-treated surfaces consisted of UV 131 irradiation for 30 s, followed by successive washes with 70% ethanol 132 and sterile PBS, respectively. The sterilization of fibronectin-coated 133 surfaces consists of a 30 min of incubation of a sterile solution of 134 fibronectin (5 μ g/mL) on sterilized samples of plain PDMS and 135 parylene-C. 136

2.3. Surface Property Characterization. 2.3.1. Contact Angle 137 Measurements. Contact angles were measured on static drops of 138 water on different substrates by using a contact angle measurement 139 system (Phoenix 300 plus, SEO) to provide information about 140 hydrophobicity of the surfaces (See Table 1). The substrates were 141 measured as-received or as-deposited (plain), and additional 142measurements were performed with a subset of these substrates 143(PDMS and parylene-C) that were treated with oxygen plasma and 144 were coated with fibronectin. The contact angle measurements were 145performed by dispensing deionized water drops $(5-10 \,\mu\text{L})$ on each 146 substrate with a micropipette (Ted Pella Inc.). Each data point 147 represents an average of >10 independent measurements. 148

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Table 1. Contact Angle Measurements

		-		
		contact angle (deg)		
substrates	untreated	plasma-treated	fibronectin-coated	
glass polystyrene PDMS	36.3 ± 2.6 79.1 ± 5.9 105.9 ± 4.5	9.9 ± 1.1^{a} 73.7 + 3 ^b	99.0 ± 6.7	
parylene-C	97.2 ± 4.2	$4.4 \pm 2.4^{\circ}$	105.0 ± 10.4	

^{*a*} Measurements made immediately after PDMS was treated in oxygen plasma. ^{*b*} Measurements made after 40 min following treatment in oxygen plasma. ^{*c*} No significant change observed in measurements made immediately after and following 40 min after plasma treatment.

Table 2. Surface Roughness Measurements

substrate	roughness (nm)
glass	1.6 ± 0.6
polystyrene	1.2 ± 0.2
parylene-C	19.3 ± 6.3
plasma-treated parylene-C	19.3 ± 5.4
fibronectin-coated parylene-C	29.0 ± 11.5
PDMS	2.2 ± 0.6
plasma-treated PDMS	0.4 ± 0.1
fibronectin-coated PDMS	3.2 ± 0.6

149 2.3.2. Surface Roughness Measurements. Surface roughness values of four different substrates (glass, polystyrene, PDMS, and parylene-150C) as received were measured with atomic force microscopy (AFM) 151152(Q-Scope 250, Quesant Instrument Corporation) using noncontact mode with a cantilever tip (NCS 16, Quesant). Scan areas of 50 μ m 153 \times 50 µm were randomly selected on the substrates. To obtain the 154 155 surface roughness values from an as-deposited thin (10 μ m) parylene 156membrane, we first peeled the parylene off the silicon wafer and then placed it on top of a robust substrate (1 mm thick PDMS slab). 157Afterward, we performed the AFM measurements. To obtain the 158 159 surface roughness of the surface-treated parylene-C, we applied 160 surface treatments (O2 plasma treatment or fibronectin coating) on 161the parylene surface mounted on a PDMS slab and performed AFM surface roughness measurements. Three independent measurements 162 from 5 μ m \times 5 μ m squares of each surface were performed and 163 averaged. Roughness values (mean) acquired from various samples 164 corresponding to the variations in surface heights are summarized 165 in Table 2. 166

2.4. Protein Adsorption Measurements. Protein adsorption was 167 168 characterized by incubating 50 μ g/mL of fluorescein isothiocyanate (FITC)-conjugated BSA (Sigma-Aldrich) and 100 µg/mL of FITC-169 170conjugated IgG (Sigma-Aldrich) on each substrate for 1 h. The substrates were encased in aluminum foil to prevent photodegradation 171 of the FITC. Following incubation, the substrates were rinsed with 172deionized water and imaged using a fluorescent microscope (Nikon 173174TE 2000) with a constant exposure time of 500 ms. Emitted fluorescence was then measured using ImageJ pixel brightness 175176 analysis tool (National Institutes of Health, U.S.A.). The average 177 pixel brightness of each image is an indirect measurement of the protein adsorption onto the substrates. Control substrates were also 178 179used to eliminate the effect of autofluorescence from the substrates.

2.5. Cell Culture. NIH-3T3 fibroblasts were cultured in Dul-180 181 becco's modification of Eagle medium (DMEM) (Invitrogen) supplemented with 10% fetal bovine serum (Atlanta Biologicals) 182and 1% penicillin-streptomycin (MediaTech). AML-12 hepatocytes 183 were preserved in 44.5% DMEM and 44.5% Ham's F12 media 184 (Invitrogen) supplemented with 10% FBS and 1% penicillin-185 186 streptomycin (Sigma-Aldrich). The cells were maintained under 187 humid conditions, at 37 °C, and in a 5% CO₂ atmosphere.

188**2.6. Preparation of Samples for Cell Adhesion.** The NIH-3T3189fibroblasts and the AML-12 hepatocytes were trypsinized and190resuspended in their respective media to form a 5×10^4 cells/mL191stock solution. A volume of 2 mL of this stock solution was then192incubated on each of the substrates for 6 h. Each well of the Costar19324-well microplate has a base area of 283.5 mm². This corresponds194to a loading density of ~353 cells/mm². Next, the substrates were

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rinsed twice with $1 \times$ phosphate-buffered saline (Invitrogen), and 195 cells adhering to the substrates were then fixed using 4% paraformaldehyde (Sigma-Aldrich) and permeabilized with 0.2% Triton X-100 197 (Sigma) for 10 and 5 min, respectively. 198

2.7. Visualization and Imaging of Adhered Cells. To count the number of adhered cells on each surface, images of fluorescently labeled nuclei were collected using a fluorescent microscope (Nikon Eclipse TE 2000). Three pictures per well and three wells per substrate were analyzed and counted using ImageJ software.

2.8. Visualization and Imaging of Cells for Estimating Shape204Factors. To analyze cell spreading on various surfaces, data was205collected from at least 70 adhered cells per sample. To effectively206analyze cell shape, the dimensionless shape factor, S, was used to207compare the spreading of cells. It is computed as208

 $S = 4\pi A/P^2$

where A is the area occupied by the cell and P is the perimeter of
the cell. A shape factor of 1 corresponds to a perfect circle, whereas
a shape factor of 0 represents a line. Cell shape factors were computed
utilizing the calibration and measurement features of the SPOT
Imaging Software.219
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3. Results and Discussion

We evaluated the surface properties of parylene-C stencils in 215comparison with other commonly used biomedical materials, 216such as PDMS, glass, and polystyrene. In addition, we analyzed 217the effect of two common surface treatments, oxygen plasma 218and protein coating on these substrates. The surfaces were 219 characterized for their hydrophobicity and roughness as well as 220 for protein adsorption, cell adhesion, and cell morphology. 221Particular attention was paid to the differences between parylene-C 222 and PDMS, due to their emerging applications in biomedical 223microfabrication. 224

3.1. Surface Analysis. Surface hydrophobicity and surface 225roughness are important factors in cell adhesion and the resulting 226 cellular morphology.^{14,29-31} In addition, hydrophobicity has also 227 been shown to affect protein adsorption. $^{32-35}$ Therefore, it is 228 important to evaluate these properties in parylene-C membranes 229 to understand the interaction of mammalian cells with these 230 substrates. To assess the hydrophobicity of the surfaces, we 231measured contact angles of as-deposited and treated parylene-C 232surfaces and compared the values to control surfaces (Table 1). 233The substrates varied greatly in their water contact angles, from 234 \sim 36° for glass to \sim 111° for PDMS. As-deposited parylene-C 235and plain PDMS were both hydrophobic as they exhibited contact 236 angles of $\sim 100^{\circ}$, which is consistent with our previous study.³⁶ 237Furthermore, fibronectin-coated parylene-C and PDMS were also 238hydrophobic (contact angles of $\sim 100^{\circ}$). This hydrophobic 239

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Figure 1. Adsorption of FITC–BSA (A) and FITC–IgG (B) onto each of the substrates. (A) Adsorption of FITC–BSA onto parylene-C and PDMS exceeds that on glass and polystyrene. Surface modifications of parylene-C and PDMS show a negative effect on FITC–BSA adsorption. (B) FITC–IgG adsorbs onto glass, parylene-C, and PDMS in a comparable manner. Adsorption onto polystyrene is the highest. Surface modification of parylene-C and PDMS reduces their affinity for FITC–IgG. In general, plasma treatment and fibronectin coating the two polymers reduces their ability to adsorb proteins. The * indicates p < 0.05.

property of fibronectin-coated PDMS is confirmed by results 240obtained by other groups.³² Even though there has not been 241investigation in the past on fibronectin-coated parylene-C, it is 242logical to expect it to be hydrophobic. Because fibronectin 243244 coatings have no electron donor components and have low surface energy,33 materials coated with fibronectin would not form 245246hydrogen bonds with water molecules, so they would become 247hydrophobic. Furthermore, air plasma treatment reduced the One of the main drawbacks of using PDMS for fluidic devices 252is that the plasma-induced hydrophilicity of the PDMS surfaces 253is short term.⁷ In many applications involving fluidics and cells, 254the ability to generate substrates that remain hydrophilic may be 255beneficial. To compare the stability of plasma-treated surfaces, 256we measured the contact angles of plasma-treated parylene-C 257and PDMS surfaces immediately and 40 min after plasma 258treatment. It was observed that although the plasma treatment 259

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Figure 2. Adhesion of (A) NIH-3T3 fibroblasts and (B) AML-12 hepatocytes on the various substrates. (A) The cells do not adhere to as-deposited parylene-C and plain PDMS. Furthermore, plasma treatment and fibronectin coating of the two polymers increase their adhesiveness to NIH-3T3 cells. (B) Similar trends are exhibited by AML-12 adhesion to the various substrates. The * indicates $p \le 0.05$.

initially decreased the contact angle values, the hydrophilicity 260of a PDMS substrate deteriorated rapidly (Table 1). This is due 261 to the viscoelastic properties of PDMS, in which the surface 262molecules "turn over" with time exposing non-plasma-treated 263

molecules of the PDMS on its surface. On the other hand, the 264contact angle for plasma-treated parylene-C did not change 265significantly after 40 min. The fact that the plasma-treated parylene 266 surface stays hydrophilic for longer periods could be advantageous 267

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Figure 3. Dimensionless cell shape factor measurements for (A) NIH-3T3 fibroblasts and (B) AML-12 hepatocytes cultured on various substrates. (A) NIH-3T3 cells exhibit greater spreading on fibronectin-coated parylene-C and PDMS, as compared to the other substrates. (B) Due to nonaxial spreading, shape factor was not an adequate measurement of AML-12 proliferation on the surfaces, and ANOVA was not conducted on this data. The * indicates p < 0.05.

for various biological applications. We next examined the surface 268 roughness of the parylene-C and PDMS substrates by using AFM. 269 As shown in Table 2, as-deposited and treated parylene-C 270substrates were significantly rougher compared to other substrates, 271 including glass, PDMS, and polystyrene. Fibronectin-treated 272273parylene-C surfaces had the highest roughness values of ~ 30 nm. The higher surface roughness values of parylene-C may be 274due to the irregularities in the deposition process, which were 275276further increased with fibronectin molecules adsorbed onto the 277surface. On the other hand, PDMS, glass, and polystyrene were much smoother with surface roughness values of <3 nm. An 278279 increase in surface roughness enhances the protein adsorption 280 level, since there is more available surface area for proteins to attach.²⁹ When there are more proteins adsorbed onto the surface, 281 more integrin receptors on the cells will bind to the proteins and, 282 therefore, mediate the attachment of cells.²⁹ 283

3.2. Protein Adsorption. To generate substrates that are 284favorable for cell adhesion, a routine procedure is to coat a layer 285of adhesive proteins on the substrates. To measure protein 286 adsorption properties of parylene-C relative to other substrates, 287we incubated each sample with fluorescently labeled BSA and 288IgG. First observation we made was that significantly more BSA 289 and IgG adsorbed to polystyrene relative to glass. This finding 290 is consistent with other studies that BSA and IgG adsorb more 291 onto highly hydrophobic surfaces like polystyrene, compared to 292 relatively hydrophilic surfaces like glass.^{14,29,31,33} Similarly, BSA 293 adsorption level on plain PDMS and as-deposited parylene-C 294were 3 times higher relative to glass (Figure 1A). On the other 295 hand, IgG adsorption levels on plain PDMS and as-deposited 296 parylene-C were similar to adsorption on glass. We believe that 297 this discrepancy is caused by the intrinsic difference in the 298 structure of two proteins. 299

Biocompatibility of Parylene-C



Figure 4. Micrographs of NIH-3T3 fibroblasts (A) and AML-12 hepatocytes (B) on various surfaces. The insets contain images which have been cropped and magnified for optimal viewing. Scale bar = $100 \ \mu$ m.

In addition, we analyzed the effects of plasma treatment and 300 initial protein coating on IgG and BSA adsorption. Plasma 301 302 treatment is routinely used to increase the surface hydrophilicity of materials, such as PDMS and polystyrene, and can be used 303 to modify the surface of parylene-C substrates (Table 1). In our 304 studies, plasma treatment of parylene-C and PDMS increased 305 the hydrophilicity of the surfaces and reduced the adhesion of 306 307 both BSA (Figure 1A) and IgG (Figure 1B). This is because of increased hydrogen bonding between the surface and water 308 molecules, which displaces the weak electrostatic interaction 309 and hydrophobic interactions between serum proteins and the 310 surface.33 In addition, fibronectin coatings, which improve cellular 311 adhesion on biomaterials, could also be used to minimize the 312 subsequent adsorption of BSA and IgG. This can be explained 313 by the fact that the adsorption of the first layer of protein results 314 315 in the creation of a thermodynamically stable interface of water molecules coupled with the hydrophilic regions of the adsorbed 316 protein layer.34 This phenomenon is commonly used in immu-317 noassays, in which an adsorbed layer of protein is applied to 318319 minimize background adsorption of the antibody to the substrate.³⁴

Thus, our results indicate that as-deposited parylene-C has high BSA and IgG adsorption, while surface treatments on parylene-C can be used to decrease levels of adsorption of these proteins. The ability to modify the level of protein adsorption on the parylene-C substrates is of potential value for various biomedical applications and microfabrication techniques.

3.3. Cell Adhesion and Spreading. To evaluate the cytocompatibility of parylene-C substrates relative to other materials, we analyzed the adhesion of and spreading of fibroblast (NIH-329 3T3) and hepatocyte (AML-12) cell lines. In these experiments, cells were seeded on various surfaces and incubated for 6 h, and

the adherent cells were counted and measured. In Figure 2, parts 331 A and B, varying levels of cell adhesion on the different substrates 332 were displayed. Interestingly, both plain PDMS and as-deposited 333 parylene-C substrates heavily repelled cell adhesion for both 334 NIH-3T3 and AML-12 cells. On both surfaces, cells remained 335 in round shape and could be easily washed away. These substrates 336 were significantly less adhesive to cells than tissue culture 337 polystyrene and glass controls. 338

To investigate how surface treatments influence cell adhesion, 339 we examined the effects of plasma treatment and fibronectin 340 coating on the parylene-C and PDMS substrates. It was found 341that both treatments resulted in an increase in the cell adhesiveness 342 of the substrates. Previously, it has been demonstrated that plasma-343 treated PDMS surfaces display enhanced cell adhesion.14,29,31 344 Also, fibronectin, which is an extracellular matrix component 345that mediates cell adhesion and spreading, has been used 346 extensively to increase cell adhesion.³² As expected, the adhesion 347 levels of both NIH-3T3 and AML-12 cells on surface-modified 348 substrates (via plasma treatment and fibronectin coatings) were 349 significantly enhanced as shown in Figure 2, parts A and B. 350 Interestingly, there was a difference in cell adhesion levels 351between plasma-treated and fibronectin-coated PDMS, whereas 352 no significant change was observed for parylene-C substrates. 353 This phenomenon can be caused by the temporary effect of plasma 354treatment on PDMS compared with the longer lasting effect of 355 plasma treatment on parylene-C (Table 1). Therefore, it may be 356 that as the plasma-induced hydrophilicity of the PDMS substrates 357 is decreased, plasma-treated PDMS was less favorable to cell 358 adhesion. In comparison, plasma-treated parylene-C remained 359 hydrophilic and suitable for cell adhesion. These results 360 demonstrate that even though as-deposited parylene-C substrates 361

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are relatively rough (>20 nm) and hydrophobic, and cells do not
 adhere onto these surfaces. Furthermore, it is possible to increase
 the surface-adhesive properties simply by surface treatment
 approaches such as plasma treatment and fibronectin coating.

In addition to analyzing the number of cells adhering onto the 366 various surfaces, we also examined the degree of cell spreading. 367 The level of spreading is important because it influences various 368 369 parameters, such as cellular proliferation and differentiation.³² 370 To determine the degree of cell spreading on different substrates, we quantified cell morphology by calculating a dimensionless 371 372 shape factor with the results illustrated in Figure 3. The dimensionless shape factor ranges from zero (line-for linearly 373 spread cell) to one (circle-for cells that have not elongated) and 374can be used to validate the degree of cell adhesiveness on a 375 surface. It is then only appropriate to use this shape factor to 376quantify cell spreading if the adhered cells display axial spreading. 377 Note that, whereas NIH-3T3 cells spread along a single axis, 378 AML-12 cells spread more uniformly with extended pseudopodia, 379 therefore exhibiting inherently higher shape factors. Phase 380 381 micrographs of both NIH-3T3 and AML-12 cells are displayed in Figure 4. Randomly selected cells from each image were 382 383 chosen and traced using SPOT imaging software to emphasize 384 the differences in cell morphology among the various substrates.

The results indicate that NIH-3T3 cells spread well on 385 fibronectin-coated parylene-C and PDMS as well as tissue culture-386 treated polystyrene (Figure 3A). As for plasma-treated parylene-C 387 and PDMS, a small fraction of NIH-3T3 cells on these substrates 388 exhibited adhered and spread morphologies, whereas the rest 389 maintained their circular phenotype. The majority of NIH-3T3 390 cells on plain substrates of parylene-C, PDMS, and glass remained 391 circular. On the other hand, since all AML-12 cells adhered in 392 a nonelongated manner, their cell shape factor values were higher 393 394 and similar in value (Figure 3B). As a result, cell shape factor is not a conclusive measure of spreading of AML-12 cells. 395

396 The cell adhesion and morphology on as-deposited parylene-C 397 and plain PDMS substrates were not significantly different 398 (Figures 2 and 3), despite a clear difference in surface roughness 399 between the two materials (Table 2). Several studies have concluded that increasing the surface roughness increases the 400 levels of cell adhesion and spreading.^{25,29,32} However, after 401 402 comparing the level of cell adhesion and spreading between 403 as-deposited parylene-C and plain PDMS substrates, we believe

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that the intrinsic differences between materials outweighed the404effect of surface roughness. Therefore, future biocompatibility405studies that utilize different polymeric substrates with varying406levels of surface roughness are required to fully clarify this matter.407

Overall, we have found that plasma-treated and fibronectin-408 coated parylene-C membranes were as compatible for cell culture 409 as commonly used substrates such as glass and polystyrene. The 410 ease with which as-deposited surfaces can be made cell adhesive 411 and the cell-resistant property of as-deposited parylene-C may 412 potentially be useful in biomedical applications. Similarly, these 413 substrates can be engineered to enhance or reduce protein 414 adsorption which is conducive to biological research. Long-term 415studies on the biocompatibility and more comprehensive trials 416 with other proteins and cell types might be necessary to fully 417 understand the benefits and limitations of parylene-C. 418

4. Conclusions

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Although parylene-C has been used as a biologically inert 420 coating on implantable devices for many years, there had not 421 been systematic studies of the biocompatibility of parylene-C 422and its surface-treated variants. In this paper, we have compared 423 parylene-C to other commonly used cell culture substrates and 424 demonstrate that surface-treated parylene-C substrates exhibit 425 adhesion levels comparable to commercially available tissue 426 culture-treated polystyrene. On the other hand, as-deposited 427 parylene-C substrates, which are not cell adhesive, can be used 428 as restrictive coating to minimize cell adhesion. Overall, 429 parylene-C can easily be surface modified into a suitable substrate 430 for culturing mammalian cells. Given that parylene-C has already 431 been shown to be well suited for microfabrication, and that it can 432 be made into flexible and robust devices, the data presented here 433 would be useful for the implementations that tailor to the 434biocompatibility of parylene-C. The new implementations of 435 parylene-C will likely lead to new technologies and devices for 436 biomedical applications. 437

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