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¹ Rectangular Bunched Rutile TiO₂ Nanorod Arrays Grown on Carbon ₂ Fiber for Dye-Sensitized Solar Cells

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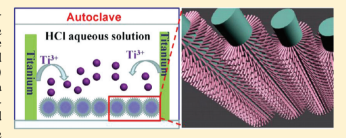
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Supporting Information

ABSTRACT: Because of their special application in photovoltaics, the growth of one-dimensional single-crystalline TiO₂ nanostructures on a flexible substrate is receiving intensive attention. Here we present a study of rectangular bunched TiO₂ nanorod (NR) arrays grown on carbon fibers (CFs) from titanium by a "dissolve and grow" method. After a corrosion process in a strong acid solution, every single nanorod is etched into a number of small nanowires. Tube-shaped dye-sensitized solar cells are fabricated by using etched TiO₂ NR-coated CFs as the photoanode. An absolute energy



conversion efficiency of 1.28% has been demonstrated under 100 mW cm⁻² AM 1.5 illumination. This work demonstrates an innovative method for growing bunched TiO₂ NRs on flexible substrates that can be applied in flexible devices for energy harvesting and storage.

22 INTRODUCTION

23 The insufficient fossil-fuel-based energy supplies and excessive 24 CO₂ emissions are the two major issues for the current global 25 energy strategies. Searching for "green" energy resources is one 26 of the most urgent challenges for the sustainable development 27 of human civilization, which could be potentially solved 28 by renewable energy technology. 1-8 As one of the most pro-29 mising photovoltaic technologies, dye-sensitized solar cells $_{
m 30}$ (DSSCs) have received intensive attention. $^{9-12}$ A conversion 31 efficiency of more than 11% has been obtained by adopting a 32 photoanode that consists of a TiO₂ nanocrystal thin film 33 covered by a monolayer of dye molecules. 13,14 However, this 34 kind of cell is usually based on a rigid fluorine-doped tin oxide 35 (FTO) glass subtrate that is not suitable for transportation, 36 installation, and remote application. To strengthen the 37 adaptability of DSSCs, polymer substrates, 15,16 metal sheets, 17 38 or metal wires 18,19 have been used as substrates to fabricate 39 DSSCs. At the same time, in practical applications, poly-40 mer substrates are limited by their poor thermostability, while 41 metallic substrates are not very stable in the electrolyte 42 that contains iodine (I^{3+}/I^{+}) . To solve this problem, a 43 substrate that could be widely used is highly desirable for this 44 technology.

Here we introduce a fiber-shaped solar cell based on carbon 46 fibers (CFs). Relative to the traditional photoanode, CFs are 47 flexible, conductive, and stable in liquid electrolyte, and they 48 can supply a large surface area, which is critical for 49 nanostructure-based photovoltaic technology. They also have 50 good heat resistance and fatigue. Moreover, CFs could be a

promising candidate to be built into weavelike solar cells that 51 can be fabricated into clothes. Although this material shows the 52 possibility of application for DSSCs, a method for effective 53 growth of commonly used anode materials on the CFs for 54 DSSCs is still an issue, $^{20-22}$ especially for single-crystal TiO₂ 55 nanorods (NRs) or nanowires (NWs). To date, most of the 56 TiO₂ nanostructures on CFs have been limited to amorphous 57 TiO₂ nanoparticles obtained by sol-gel methods. ^{23,24} However, ₅₈ the TiO2 nanoparticles obtained by sol-gel methods are not 59 crystalline, and high temperature with protected gas is required 60 for calcination to form the crystalline phase. Moreover, the 61 material quality is not very good because of the uneven surface 62 and the existence of cracks when increasing the thickness. An 63 ideal solution for preparing TiO2 nanostructures or films on 64 CFs is direct growth of orderly one-dimensional and single- 65 crystalline TiO2 NWs on the CFs. A direct connection of TiO2 66 NWs with the substrate may improve the DSSC performance. 67 Specifically, oriented single-crystalline TiO2 NRs or NWs on a 68 conductive substrate without annealing would be most 69 desirable, 25,26 but achieving these structures has been limited 70 by the availability of synthetic techniques. Herein we introduce 71 a new method that includes "dissolve and grow" and "etch and 72 grow" processes to prepare ultrafine and uniform, single-crystal 73 TiO2 NWs on CFs.

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75 EXPERIMENTAL SECTION

76 A schematic illustration of the growth of NRs on the CFs is shown in 77 Figure 1A. In a typical synthesis process, 0.025-0.1 g of Ti foil and 78 18 mL of 0.05-0.1 M HCl solution were put into a Teflon-lined 79 stainless steel autoclave with a total volume of 25 mL. The CFs were 80 then immersed into the solution after ultrasonic cleaning for 30 min in 81 a 1:1:1 (v/v/v) mixture of acetone, ethanol, and deionized water. The 82 hydrothermal synthesis was conducted at 423-463 K for 2-18 h in an 83 electric oven. The autoclave was cooled to room temperature with 84 flowing water for 10 min after the growth process. Next, the CFs were 85 ultrasonically cleaned for 3 min in a 2:1 (v/v) mixture of isopropanol 86 and water. The thus-cleaned CFs with TiO2 NRs around them were 87 transferred into Teflon-lined stainless steel autoclave again and treated 88 in the 9-27% HCl solution at 433-463 K for 2-10 h for the etching 89 process. The resulting CFs with TiO2 NRs around them were 90 immersed in 100 mL of 0.2 M TiCl₄ aqueous solution for 6-8 h and 91 then annealed in argon at 723 K for 30 min to increase crystallinity. The resulting CFs with TiO2 NRs around them were sensitized in a 93 0.3 mM solution of cis-bis(isothiocyanato)bis(2,2'-bipyridyl-4, 94 4'-dicarboxylato)ruthenium(II) bis(tetrabutylammonium) dye (N-719 95 as received from Solaronix) in dry ethanol for 24 h. Tube-shaped solar 96 cells were assembled according to the method introduced by Zou and 97 co-workers:²⁷ first, the N719-sensitized CFs were wound uniformly 98 and spirally onto a platinum-coated optical fiber, which was then 99 inserted into the sealing capillary ($\Phi = 0.5$ mm). The internal space of 100 the device was filled with a liquid electrolyte [0.5 M LiI, 50 mM I₂, and 101 0.5 M 4-tert-butylpyridine in 3-methoxypropionitrile (Fluka)] by the 102 capillary effect.

The morphology and microstructure of the TiO₂ nanostuctures were examined by scanning electron microscopy (SEM) using a LEO 105 1530 scanning electron microscope and by transmission electron microscopy (TEM) using Hitachi HF2000 and JEOL 4000EX 107 transmission electron microscopes. Phase identification of TiO₂ was conducted by X-ray diffraction (XRD) using a PANalytical X'Pert 109 PRO diffractometer. A SoLux solar simulator was used as calibrated with a Daystar meter to simulate sunlight for an illumination intensity 111 of 100 mW cm⁻². The solar cell was irradiated using a solar simulator 112 (500 W model 91160, Newport) with an AM 1.5 spectrum distribution calibrated against an NREL reference cell to simulate accurately 114 a full-sun intensity (100 mW cm⁻²). The irradiated length of this kind 115 of tube-shaped three-dimensional (3D) DSSC from the side of the 116 working electrode was 1–3 cm.

117 RESULTS AND DISCUSSION

118 Figure 1 shows typical SEM and TEM images of TiO₂ NRs 119 grown on the CFs after hydrothermal reaction at 463 K for 3 h. 120 As can be seen from Figure 1B,C, the entire surfaces of the CFs 121 are covered very uniformly by TiO2 NRs that are tetragonal in 122 shape with square top facets. A cross-sectional view of the TiO₂ 123 NRs is shown in Figure 1D; the diameter and length of the NRs 124 were found to be \sim 150 nm and \sim 3 μ m, respectively. The phase 125 and crystal structure of the TiO₂ NRs were confirmed by the 126 lattice image of Figure 1E. The distances between lattice 127 fringes, 0.32 and 0.29 nm, can be assigned to (110) and (001) 128 of the rutile TiO₂ phase, respectively, suggesting that the TiO₂ 129 NRs grew along the [001] axis. The corresponding selected-130 area electron diffraction (SAED) pattern (Figure 1F) displays 131 the single-crystalline nature and could be indexed to the pure 132 rutile TiO₂ phase. The XRD pattern of the CFs covered by 133 TiO₂ NRs is shown in Figure S1A in the Supporting 134 Information (SI). All of the reflection peaks can be readily 135 indexed to pure rutile TiO₂.

Figure 1A shows the schematic diagram of the formation mechanism of TiO₂ NRs by the "dissolve and grow" method, and it may be described by the following chemical reactions:²⁸

$$2\text{Ti} + 6\text{HCl} \rightarrow 2\text{TiCl}_3 + 3\text{H}_2(g)$$

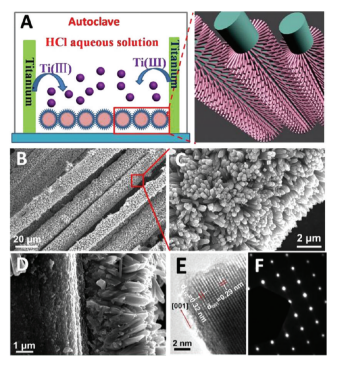


Figure 1. (A) Schematic representation of the growth of TiO_2 nanorod (NR) arrays on carbon fibers (CFs) by the "dissolve and grow" method. (B, C) SEM images of the top view of TiO_2 NR arrays on CFs. (D) Cross-sectional view of the well-aligned TiO_2 NR arrays. (E) HRTEM image of a single TiO_2 NR. (F) SAED pattern of the same TiO_2 NR.

$$Ti^{3+} + H_2O \rightarrow TiOH^{2+} + H^+$$

$$TiOH^{2+} + O_2 \rightarrow Ti(IV)$$
-oxo species + $O_2 \rightarrow TiO_2$

At the very beginning, in the presence of HCl, Ti foil reacts 139 with H⁺ at high temperature and pressure and gradually 140 dissolves, continually releasing the Ti(III) precursors into the 141 reaction solution. Because Ti(III) is not stable in aqueous 142 solution, TiOH²⁺ is produced by hydrolysis of Ti(III). 143 According to the suggestion by Fujihara and co-workers, 28 144 TiOH²⁺ is oxidized to Ti(IV) by reaction with dissolved 145 oxygen. The Ti(IV) complex ions are thus used as the growth 146 units, and the formation mechanism of the rutile TiO2 NRs 147 may be described as follows: For rutile TiO2, a TiO6 148 octahedron forms first by bonding of a Ti atom and six oxygen 149 atoms. The TiO₆ octahedron then shares a pair of opposite 150 edges with the next octahedron, forming a chainlike structure. 151 Because the growth rate of the different crystal faces depends 152 on the numbers of corners and edges of the coordination 153 polyhedra available, the growth of rutile NRs follows the $_{154}$ sequence (110) < (100) < (101) < (001). 29,30 Thus, rutile $_{150}$ NRs growing along the [001] direction are formed.

According to the XRD pattern shown in Figure S1B in the SI, $_{157}$ the CFs have an amorphous structure. Apparently, there is no $_{158}$ lattice match between rutile $\mathrm{TiO_2}$ and the CF substrate. $_{159}$ Therefore, it is impossible for $\mathrm{TiO_2}$ NRs to grow on the fiber $_{160}$ surface through epitaxial growth. To provide a better under- $_{161}$ standing of the growth mechanism of this process, controlled $_{162}$ experiments were carried out on a CF substrate by varying the $_{163}$ acid concentration. In the reaction process, the acid not only $_{164}$ acts as a reactant to dissolve Ti but also plays an important role $_{165}$

166 in restricting the hydrolysis rate of Ti(III) by providing an 167 acidic environment.

When the concentration of HCl is low, the Ti foil dissolves 169 relatively slowly, and the concentration of the Ti(IV) precursor 170 is also very low at the beginning. However, when most of the Ti 171 is dissolved, the growth rate of TiO2 nanorods becomes much 172 faster than before as a result of not only the increased 173 concentration of the Ti(IV) precursor but also the 174 consumption of HCl, which obviously increases the pH of 175 the solution. More and more Ti(III) produced in the solution 176 leads to a high degree of supersaturation of Ti(IV), and thus, a 177 large number of polycrystalline clusters form in the solution by 178 homogeneous nucleation. Next, these clusters are deposited on 179 the surface of the CF and act as the nucleating centers for 180 further growth of nanorods. Because there is no crystal plane 181 trend for these clusters, nanorods can grow along any direction and then self-assemble into microspheres or tufted nanoflowers, 183 as shown in Figure 2A,B. The microsphere/microflower film

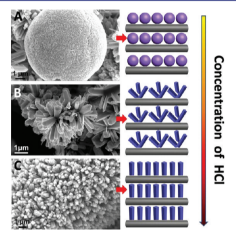


Figure 2. SEM images of the ${\rm TiO_2}$ nanostructures obtained at different concentrations of HCl on CFs: (A) 0.5 M; (B) 1 M; (C) 1.67 M. The corresponding schematic diagrams represent the growth processes of ${\rm TiO_2}$ nanostructures on CFs.

184 does not firmly adhere to the CF substrate and can easily be 185 peeled off.

When the concentration of HCl is moderate, the reaction 187 proceeds more smoothly because of the "double buffer" effect 188 of HCl. For the "double buffer" effect, first, HCl strictly 189 controls the dissolution rate of Ti, which guarantees that the 190 concentration of Ti(III) is not too high or too low. Second, 191 the consumption of HCl does not obviously decrease the con-192 centration of H⁺ when the concentration of HCl is moderate, 193 so a stable acidic environment is maintained, effectly restricting 194 the hydrolysis of Ti(IV). Therefore, the hydrothemal solution 195 maintains a low degree of supersaturation. Because of the 196 "double buffer" effect of HCl, it is difficult for TiO2 poly-197 crystalline clusters to form in the solution by homogeneous 198 nucleation; instead, the low degree of supersaturation favors 199 heterogeneous nucleation on the surface of CFs. Figure S2 in 200 the SI shows the SEM images of a rutile TiO2 NR film grown 201 on the CF substrate at 463 K for different times. When the 202 hydrothermal treatment time was 1 h, we can observe that lots of 203 very tiny TiO₂ nucleations form on the CF surface; some formed 204 TiO₂ nanorods are also shown in Figure S2B. When the 205 hydrothermal treatment time is increased, the growth of NRs is 206 very fast, and the substrate is completely covered by NRs after

2.5 h of hydrothermal treatment (Figure S2D). Microspheres 207 and microflowers do not form under these conditions.

When the concentration of HCl is high, nothing is grown on 209 the CFs, and the solution is clear even after reaction for over 210 16 h at 463 K. This may be due to the fact that the high acidity 211 seriously restricts the hydrolysis of Ti(III) and restrains both 212 homogeneous and heterogeneous nucleations.

The diameters of the obtained TiO_2 nanorods on CFs are 214 150–500 nm, which are too large for applications in many 215 fields. To increase the surface area of the TiO_2 NRs further, we 216 treated them in hydrochloric acid solution at 453 K for 3–5 h. 217 The obtained NRs were characterized by SEM and TEM, as 218 shown in Figure 3. It can be obviously seen that a different 219

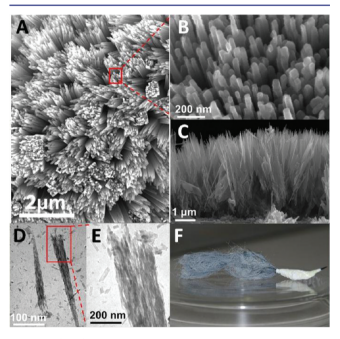


Figure 3. (A, B) Top-view SEM images of bunched TiO_2 NR arrays after hydrothermal treatment in HCl solution at 453 K for 4 h at (A) low and (B) high magnification. (C) SEM cross-sectional view of the bunched TiO_2 NR arrays. (D, E) TEM images of the the corresponding bunched TiO_2 nanorod. (F) Optical photo of the carbon fibers coated by bunched TiO_2 nanorod arrays.

morphology was obtained after 4 h of hydrothermal treatment 220 at 453 K. The top-view SEM images of the TiO_2 NRs in Figure 221 3A,B show that every single NR is etched into many smaller 222 TiO_2 NWs ranging from 10 to 30 nm in diameter and oriented 223 perpendicular to the (001) facet, while all of the the original 224 NRs keep the rectangular outline. The cross-sectional view of 225 the etched TiO_2 NRs in Figure 3C shows obvious etching 226 traces on the top of the NRs. From the TEM images of the 227 etched TiO_2 NRs in Figure 3D,E, it can be observed that a 228 single NR is etched into many smaller NWs, which reveals that 229 the NR can be etched from top to bottom before the formation 230 of the nanotube. We call this kind of structure "rectangular 231 bunched TiO_2 NRs". The formation mechanism of the 232 orientation-aligned NRs can be explained as follows.

First of all, from the SEM image shown in Figure 4A, we can 234 observe that the obtained $^{71}O_2$ NRs are aggregates of much 235 smaller $^{71}O_2$ square crystals. Upon treatment with HCl 236 solution, the etching rate in the grain boundaries of the crystals 237 is much faster than in other places because of the greater 238 number of defects and higher reactivity in the grain boundary. 239

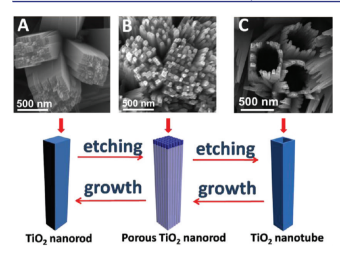


Figure 4. Formation mechanism of the rectangular bunched TiO₂ NR arrays and NT arrays. SEM images of (A) TiO₂ NR arrays, (B) bunched TiO₂ NR arrays, and (C) TiO₂ NT arrays are shown, and the corresponding schematic diagrams represent the growth mechanism of TiO₂ nanostructures on CFs.

240 With the continuous corrosion on the crystal boundaries, the 241 densely packed ${\rm TiO_2}$ square crystals are separated from each 242 other, and the original ${\rm TiO_2}$ NRs become many thinner NWs, 243 as shown in Figures 3A and 4B. Moreover, these thin NWs are 244 still not stable in the acidic solution, and when the 245 hydrothermal treatment time is further increased, all of the 246 NWs dissolve, producing rectangular nanotubes, as shown in 247 Figure 4C and further confirmed by the TEM images in Figure 248 S3 in the SI. According to the work of Liu et al., 31 the formation 249 mechanism of the ${\rm TiO_2}$ nanotube (NT) arrays can be explained 250 as follows.

When the NRs are treated with hydrochloric acid during the 252 hydrothermal process, the HCl preferentially etches the TiO₂ 253 NRs in the [001] direction because the (001) facet below is far 254 more reactive than the crystalline facet of the side wall, causing 255 the dissolution rate in the long-axis direction to be faster than 256 that in the side-wall direction. This can explain why the formed 257 NWs dissolve first and then leave the rectangular-shaped NTs. 258 To provide further comfirmation of the mechanism for the 259 formation of orientation-aligned NRs and NTs, we again used 260 0.025-0.1 g of Ti foil and 18 mL of 0.05-0.1 M HCl solution 261 as growth solution, with the resulting rectangular TiO2 NTs as 262 the growth substrate. After different growth times, the bunched 263 TiO2 NRs and original TiO2 NRs were obtained in succession 264 again, revealing that the etching and growth process is 265 reversible and providing further evidence that the more reactive 266 crystalline facet is more reactive in both growth and corrosion. It is worth mentioning that the TiO2 NR film grown on the 268 FTO substrate can easily be peeled off upon etching in the HCl

273 surface tension between the CF surface and TiO_2 NRs. 274 To assemble the DSSCs, CFs covered with \sim 5 μ m long TiO_2 275 NRs were used as photoanodes, and all of the photoanodes 276 were treated with 0.2 M $TiCl_4$ for 6–8 h. The design and 277 principle of the fiber-based solar cell is shown in Figure 5A. An 278 optical fiber coated with platinum was used as the counter 279 electrode. When sunlight irradiates the photoanode, which is 280 sensitized with N719, the excited electrons are rapidly injected

269 solution. However, for the CF substrate, the film remains

270 adhered to the CF surface and cannot be peeled off even when

271 the outline of NTs begins to dissolve (Figure S4 in the SI).

272 This may be attributed to the good bonding force and low

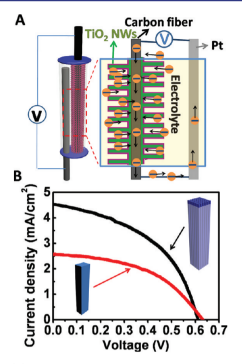


Figure 5. (A) Schematic representation of the configuration of the CF/TiO $_2$ NR array-based tube-shaped 3D DSSC. (B) J-V curves of DSSCs based on TiO $_2$ NR arrays and bunched TiO $_2$ NR arrays on CFs.

into the conduction band of the rutile TiO2 NRs and 281 transported along the CFs, from which they flow to the 282 counter electrode through the external circuit. Current 283 density-voltage (I-V) curves of these kinds of tube-shaped 284 solar cells based on TiO2 NRs and bunched NRs are shown in 285 Figure 5B. Under 100 mW cm⁻² AM 1.5 illumination, different 286 performances were observed. For the NR-based solar cell, the 287 short-circuit current density, open-circuit voltage, fill factor, and 288 efficiency were $J_{\rm sc}$ = 2.57 mA/cm², $V_{\rm oc}$ = 0.63 V, FF = 0.47, and 289 η = 0.76%, respectively, while for the bunched-NR-based solar ₂₉₀ cell, $J_{sc} = 4.58 \text{ mA/cm}^2$, $V_{oc} = 0.61 \text{ V}$, FF = 0.46, and $\eta = 1.28\%$. 291 The current density of the bunched-NR-based solar cell is 292 larger than that in original NR-based solar cell, which can be 293 attributed to the larger surface area of the bunched NRs, which 294 enables more dye molecules to be adsorbed. The relatively low 295 $V_{\rm oc}$ in both cells may be explained by the electron 296 recombination at the interface of the CFs and the loss of 297 light upon passage through the electrolyte.

The CF-based 3D DSSC has several outstanding features. 299 First, from a configuration perspective, this tube-shaped solar 300 cell can capture light from all directions, thus showing the 301 potential of this system for applications under intensively 302 focused sunlight. Second, because of their high electrical 303 conductivity, corrosion resistance toward I₂, high reactivity for 304 triiodide reduction, and low cost, carbonaceous materials are 305 quite attractive as possible replacements for platinum as 306 counter electrodes, 32–34 and therefore, we can introduce CF 307 or other carbon materials as the counter electrodes for this 308 configuration. In this case, we can fabricate DSSCs based on 309 carbon materials and TiO₂ that are promising as low-cost solar 310 cells in the future. Third, CF-based solar cells are suitable to be 311 fabricated into large-area solar cells by growth of the TiO₂ NRs 312 on carbon paper or carbon cloth, which is very promising for 313 integration into clothes in the future. Finally, the success in 314

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 $_{315}$ growing ultrafine $_{71O_2}$ NWs on CFs is significant for their $_{316}$ application in organic solar cells, photocatalysis, and lithium ion $_{317}$ batteries.

In summary, we have developed an innovative and cost-319 effective approach for growing bunched TiO₂ NRs on CFs for 320 enhancing the performance of DSSCs. The first step involves 321 direct transformation of pure Ti into vertically aligned single-322 crystal TiO2 NRs on the CFs using the "dissolve and grow" 323 method. The second is to etch the TiO2 NRs into bunched 324 TiO2 NRs using the hydrothermal method in strong acid 325 condition. On the basis of the bunched TiO₂ NR-covered CFs, 326 tube-shaped 3D DSSCs were assembled found to exhibit an 327 efficiency of 1.28%, which is 68% higher than that of the 328 original NRs. Combining the advantages of TiO₂ NRs and CFs, 329 the hybrid structure can be applied in many fields, such as 330 photocatalysis, gas sensing, organic solar cells, and lithium ion 331 batteries. Furthermore, the growth of single-crystal TiO2 332 nanostructures on conductive flexible substrates without high-333 temperature anealing may be beneficial for application in some 334 flexible devices.

335 **ASSOCIATED CONTENT**

336 Supporting Information

337 XRD patterns of CFs with and without TiO_2 NRs on them, 338 SEM images of TiO_2 NWs grown on CFs for different times, 339 and TEM and SEM images of TiO_2 nanotubes grown on CFs. 340 This material is available free of charge via the Internet at 341 http://pubs.acs.org.

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345 Notes

346 The authors declare no competing financial interest.

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