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Research Article

¹ Supported Oxygen Evolution Catalysts by Design: Toward Lower ² Precious Metal Loading and Improved Conductivity in Proton ³ Exchange Membrane Water Electrolyzers

⁴ Yagya N. Regmi, Eden Tzanetopoulos, Guosong Zeng, Xiong Peng, Douglas I. Kushner, ⁵ Tobias A. Kistler, Laurie A. King, and Nemanja Danilovic*



6 ABSTRACT: Reducing the precious metal content of water oxidation catalysts for proton-7 exchange-membrane water electrolyzers remains a critical barrier to their large-scale 8 deployment. Herein, we present an engineered architecture for supported iridium catalysts, 9 which enables decreased precious metal content and improved activity and conductivity. The 10 improvement in performance at lower precious metal loading is realized by the deposition of a 11 conformal layer of platinum nanoparticles on titanium dioxide (TiO₂) using a facile 12 photoreduction method to prepare conductive layer coated supports (CCSs). Platinum 13 nanoparticles are homogeneously dispersed on TiO₂, and the conductivity of the subsequent 14 catalysts with 39 wt % precious group metal loadings is significantly higher than the commercial 15 75 wt % loaded IrO_2 -TiO₂ catalysts. The conformal conductive layer also maintains an 16 enhanced conductivity and electrochemical activity upon thermal annealing when compared to 17 catalysts without the conductive layer and nonconformal heterogeneous conductive layer. The



18 iridium mass activity from half-cell studies shows a 141% improvement for CCS supported catalysts at 42% lower loadings compared 19 to the commercial catalysts. The conductive layer also improves the single cell electrolyzer performance at a similar catalyst loading 20 in comparison to a commercial state-of-the-art catalyst. We correlate the physical properties of the engineered catalysts with their 21 electrochemical performance in electrolyzers to understand structure—activity relationships, and we anticipate further performance 22 improvements upon synthesis and materials optimizations.

23 **KEYWORDS**: proton exchange membrane electrolyzers, water splitting, oxygen evolution reaction, supported electrocatalysts, 24 iridium catalysts, photoreduction, membrane electrode assembly

25 INTRODUCTION

26 Electrochemical hydrogen generation has the potential to 27 decarbonize the energy sector, enabling high renewable energy 28 penetration through long-term energy storage and conver-29 sion.^{1,2} Additionally, deep decarbonization of industrial 30 processes such as food production (ammonia synthesis), oil 31 refining, and steel manufacturing can be realized if the "green" 32 hydrogen from water splitting replaces hydrogen generated via 33 routes that release carbon dioxide and other harmful 34 byproducts.^{3,4} Annually, more than 95% of the 70 million 35 tons of hydrogen is produced globally from hydrocarbons via 36 steam methane reforming (SMR), water-gas shift, and coal 37 gasification, among other processes.⁵ Electrolysis currently 38 accounts for only 4% of hydrogen production. Proton-39 exchange-membrane (PEM)-based water splitting is one 40 promising technology to displace SMR, however several cost-41 cutting measures are necessary to achieve the goal. One such 42 measure is either to use curtailed renewable electricity or to 43 reach power purchase agreements to reduce the feedstock cost 44 (electricity). Eventual widespread deployment of PEM water 45 electrolyzers (PEMWEs) is projected to render the electrochemical stack as the dominant cost, and the catalyst layers 46 (CLs) as the most expensive components in the stack.⁶ Today, 47 the CLs account for about 5% of the total cost of the stack, but 48 this value is projected to increase as electrochemical hydrogen 49 production is scaled up from kW to GW.⁷ Thus, by 50 simultaneously lowering the feedstock costs and either 51 replacing or lowering the platinum group metal (PGM) 52 loadings in catalyst layers (CLs), it will be possible to lower the 53 price of green H₂ to the SMR range.⁵ The techno-economic 54 barrier emanating from high PGM loadings needs to be 55 surmounted to enable large-scale (TW) electrochemically 56 generated hydrogen to become economically competitive with 57 hydrogen generation from SMR.⁷ The balance of plant and 58 balance of cell costs are anticipated to decrease with high-59

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60 throughput manufacturing. Conversely, PGM costs are 61 expected to remain high due to their limited abundance in 62 the earth's crust, of which Ir is far rarer than Pt.⁸ To reduce the 63 precious metal loading in PEM electrolyzers, various strategies 64 including the use of non-PGM catalysts and engineering CLs 65 with lower loadings of PGM catalysts supported on low-cost 66 substrates have been explored.⁹⁻¹³ For example, King et al. 67 recently reported the use of CoP as the cathode catalyst on 68 PEMWE,¹⁴ and Alia et al. have shown the promising stability 69 of various low Ir-loaded anode CLs.¹⁵ While a diverse library of 70 non-PGM materials for cathodic hydrogen evolution reaction 71 (HER) in PEMWE has been developed,¹⁶ analogous materials 72 for anodic oxygen evolution reaction (OER) for PEMWE are 73 yet to be identified.^{17,18} While the ultimate goal is to mitigate 74 the usage of PGM catalysts, the need for short and midterm 75 strategies to lower the PGM loading represents a critical 76 development. To this end, one of the most promising strategies 77 is to develop stable and conductive electrocatalyst supports 78 akin to the carbon supports currently deployed in the CLs of 79 PEM fuel cells.

Supports employed in PEMWE anodes must withstand the highly corrosive acidic (pH < 1) and oxidizing (>2 V) conditions.¹⁹ Furthermore, the support needs to be conductive. Under these conditions, most metals will readily oxidize and form insulating layers, while the few that can withstand the harsh conditions are just as expensive as Ir. While some metal oxides can withstand the corrosive conditions, most are poor conductors.^{20,21} Strategies to boost conductivity of metal oxides such as doping titanium oxide conductivity of metal oxide have shown improved conductivity over undoped oxides; however, their conductivity is lower than that of metals and the dopants are also known to leach out over time, thereby further reducing conductivity and stability.^{22,23}

As depicted in Figure 1a, for unsupported iridium, the bulk 95 of the catalyst particle functions as the conductive support with



Figure 1. Schematics of: (a) bulk Ir catalyst, (b) traditional >50 wt % Ir shell coated on TiO_2 core, (c) proposed catalyst with Pt or Au intermediate layer between Ir and TiO_2 , (d) nanostructuring strategy to reduce Ir loading without losing overall catalyst surface area, and (e) nanostructuring of both the catalyst and conductive intermediate layer to lower PGM loading without losing the catalyst surface area or conductivity.

⁹⁶ only the surface Ir atoms participating in catalysis. Currently, ⁹⁷ PEMWE anodes typically utilize unsupported Ir resulting in ⁹⁸ relatively large PGM catalyst particles and clusters on the order ⁹⁹ of 80 nm and larger that are applied at high loading (>1 mg_{Ir}/ ¹⁰⁰ cm²) to maintain adequate CL thickness, connectivity, and porosity.^{24,25} A strategy to reduce PGM loading on PEMWE 101 anodes has been the use of non-PGM cores such as TiO_2 with 102 a shell of Ir catalysts (Ir-TiO₂) as illustrated in Figure 1b. 103 However, in order to maintain conductivity of the shell, Ir 104 loadings in excess of 50 wt % Ir are necessary,²⁶ as 105 demonstrated for half-cell OER and electrolyzer perform- 106 ances.^{2,11,19,27,28} 107

We propose modifying the core—shell architecture by adding 108 a very thin layer of conductive and electrochemically stable 109 interfacial material, namely Pt or Au, between the TiO_2 core 110 and the Ir catalyst (Figure 1c-e). Such conductive layers can 111 also provide the possibility to further nanostructure the Ir 112 catalyst and the conductive layer itself to reduce overall PGM 113 loading without compromising specific activity, as depicted in 114 Figure 1d,e. This strategy represents a new pathway to reduce 115 Ir loading without compromising the activity and conductivity. 116 Replacing Ir with Pt or Au has economic advantages as well in 117 the short and medium terms, that is, until earth-abundant 118 elements replace PGMs as OER catalysts. Pt and Au are a 119 magnitude more abundant than Ir on the earth's crust.²⁹ 120

Our inspiration comes from the use of Pt as a corrosion- 121 resistant protective coating in PEMWE anode porous transport 122 layers and bipolar plates, where they are known to withstand 123 over 50,000 h of service. $^{30-33}$ These precious metal-coated 124 TiO₂ conductive catalyst supports (CCSs) also offer a pathway 125 to enhance mass transport in the CLs by changing the TiO₂ 126 particle size and thus improving the CL porosity. Akin to the 127 roles carbon supports play in Pt/C CLs,³⁴ we advocate that the 128 use of a substrate has the potential to increase the CL 129 thickness, improve the distribution of Ir at low loadings, and 130 renders an extra degree of flexibility in design by allowing for 131 different Ir wt % on the support to be used. Generation of 132 targeted materials interfaces at the nanoscale presents 133 additional opportunities to engineer highly efficient and stable 134 active sites.³⁵

In this paper, we synthesized CCS particles with thin layers 136 of either Pt or Au nanoparticles prepared by a photochemical 137 (PC) reduction deposition method (Table S1). As a control, 138 CCSs were also synthesized using an incipient wetness 139 impregnation (WI) method. In both systems, Ir was 140 subsequently deposited on the CCSs using the WI method. 141 Conductivities were measured using an in-house assembled 142 DC conductivity setup,³⁶ and catalyst activities toward the 143 oxygen evolution reaction (OER) were evaluated in a rotating 144 disk electrode (RDE) and in a PEMWE. Throughout, we 145 compare the performance with state-of-the-art commercial 146 catalysts and previous investigations using supported Ir 147 catalysts from the literature. We also correlate the activities 148 with the physical characteristics of the catalysts using scanning 149 and transmission electron microscopy (SEM and TEM), 150 energy dispersive X-ray spectroscopy (EDS), X-ray photo- 151 electron spectroscopy (XPS), and X-ray diffraction (XRD). 152 Our findings provide rational pathways toward engineering 153 more efficient and stable active sites at the support, conductive 154 layer, and catalyst surfaces and interfaces. 155

EXPERIMENTAL SECTION

156

Photochemical Synthesis of CCS. A TiO₂ nanoparticles ¹⁵⁷ (<100 nm, 42–52 wt % in xylene, Sigma-Aldrich) slurry (5 ¹⁵⁸ mL) is added to 15 mL of deionized water (15 mL) in a ¹⁵⁹ centrifuge vial and vortexed. The suspension is centrifuged at ¹⁶⁰ 3500 rpm (700g) for 10 min before the supernatant is ¹⁶¹ decanted and the precipitate is resuspended in deionized water ¹⁶²

f1

163 (20 mL). The centrifugation and washing are repeated 4 times. 164 The final TiO₂ precipitate is dried under vacuum at 60 °C for 165 12 h. Dried TiO₂ nanoparticles (200 mg) are added to 166 deionized water (50 mL) and combined with an appropriate 167 mass of chloroplatinic or chloroauric acid (Fischer Scientific) 168 to achieve 25 wt % metal on TiO2. The combined materials are 169 stirred for 20 min to prepare a homogeneous mixture. The 170 mixture is placed under a solar simulator equipped with a Xe 171 lamp and AM1.5G filter, while stirring for 4 h to allow the 172 deposition of gold or platinum nanoparticles onto the TiO₂ 173 nanoparticle substrates (Supporting Video S1).³⁷ The solution 174 is then centrifuged three times at 9000 rpm (7000g) for 15 min 175 each. After each cycle, the supernatant is disposed of, and the 176 50 mL centrifuge vial was filled completely with deionized water and sonicated for 5 min before returning to the 177 178 centrifuge.

Incipient Wetness Impregnation and Thermal Re-179 180 duction Synthesis. An appropriate mass of dried TiO₂ 181 nanoparticles is placed in a mortar and pestle. The exact 182 mass of chlorometallic salts of Pt and Au is dissolved in 183 acetone (2 mL, VWR Chemicals). Then, the chlorometallic 184 salts are added dropwise to the TiO₂ powder while 185 continuously grinding with the pestle. The resulting mixture 186 is transferred to a quartz boat and placed in a tube furnace. 187 The temperature is ramped to 400 °C at 10 °C/min under a reducing atmosphere (5° H₂ in Ar). After dwelling at 400 °C 188 189 for 2 h, the temperature is allowed to naturally cool down to 190 room temperature. On both the Au- and Pt-coated CCSs, 25 191 wt % Ir is deposited using the incipient wetness impregnation 192 method as well as to prepare Ir-Pt/Au-TiO₂-PC/WI catalysts. 193 Therefore, PC and WI indicate the synthesis pathway for the 194 deposition of the Pt and Au conductive layer onto TiO₂, with 195 Ir always deposited using WI on CCS.

196 Thermal Oxidation of CCS Supported Ir Catalysts. 197 The catalysts are loaded on quartz boats and placed inside a 198 furnace. The temperature is ramped at 10 $^{\circ}$ C/min to 400 $^{\circ}$ C 199 under ambient conditions. The furnace temperature is kept at 200 400 $^{\circ}$ C for 2 h and then naturally allowed to cool down to 201 room temperature.

Physical Measurements. XRD plots are recorded in 202 203 Bragg-Brentano optics geometry with a Cu K α source on a 204 Rigaku SmartLab instrument. SEM micrographs are obtained 205 using a SEM FEI Quanta 250 FEG equipped with a Bruker 206 Quantax 200 EDX detector. XRF measurements are carried 207 out using a Bruker M4 Tornado Micro-XRF. STEM micrographs were acquired using a 200 kV FEI monochromated F20 208 209 UT Tecnai instrument. The TEM-EDX maps are collected on 210 a FEI TitanX 60-300 microscope equipped with a Bruker 211 windowless EDS detector with a solid angle of 0.7 steradians. 212 XPS data are collected on a Kratos Axis Ultra DLD system at a 213 takeoff angle of 0° relative to the surface normal at room 214 temperature. A monochromatic Al K α source ($h\nu$ = 1486.6 215 eV) is used to excite the core level electrons of the material. Pt 216 4f, Ir 4f, Ti 2p, C 1s, and O 1s core levels are collected, with a 217 pass energy of 20 eV, step size of 0.05 eV, and 8 sweeps each to 218 obtain a good signal-to-noise ratio. The survey spectra (Figure 219 S1) are acquired with a pass energy of 160 eV, step size of 1 220 eV, and 3 sweeps. The measurements are performed at ²²¹ ultrahigh-vacuum conditions (7.5×10^{-9} Torr).

Conductivity Measurements. A homemade DC conductivity setup was used to measure the particle conductivity.³⁶ Priefly, the catalyst powder was sandwiched between copper electrodes and compressed to 200 lb/in², the thickness was measured, and an I-V curve was generated from -50 to 50 226 mV using a potentiostat (SP300, BioLogic). The catalyst 227 conductivity was extracted from the slope of the curve and the 228 catalyst thickness. 229

Electrode Preparation. The electrochemical measure- 230 ment system and methods are adapted from earlier reports.³⁸ 231 The gold disk working electrode is prepared by polishing with 232 a 0.05 μ m alumina slurry. The disk is rinsed and bath sonicated 233 for 30 s to remove residual slurry and impurities. The disk is 234 rinsed again and dried via a compressed nitrogen air gun. The 235 glassware is cleaned by rinsing in ultrapure water (18.2 M Ω , 236 Milli-Q), boiling for 1 h, and rinsing again. Catalyst inks are 237 coated as a thin film layer on a polished gold tip working 238 electrode (5 mm diameter). To prepare the inks, catalyst 239 powder (3.5 mg) is combined with water (7.6 mL), 2-propanol 240 (2.4 mL, VWR), and Nafion solution (40 μ L, 5 wt % in 241 alcohol, Ion Power). The catalyst suspension was bath 242 sonicated for 20 min. Ten μ L of the well-dispersed catalyst 243 ink was pipetted onto the gold disk surface of the working 244 electrode as it rotated on an inverted RDE at 100 rpm to 245 distribute the ink evenly. The RDE speed is increased to 700 246 rpm and allowed to dry for 15-20 min until it was completely 247 dry. 248

Electrochemical Measurements (RDE). Electrochemical 249 measurements are performed using a Pine Instrument RDE 250 and SP-300 potentiostat from BioLogic Science Instruments. 251 The activities are recorded in 0.1 M HClO₄ electrolyte at 25 252 °C. A gold wire and dynamic hydrogen electrode (DHE) are 253 used as the counter and reference electrodes, respectively. All 254 potentials are adjusted to RHE. Once set up, the system is 255 purged with ultrahigh-purity Ar for 15 min before the working 256 electrode is placed in the electrolyte, and the electrochemical 257 measurements are recorded. Commercial supported baseline 258 catalyst, Elyst Ir75 0480, IrO₂-TiO₂ (75 wt % Ir) on TiO₂, is 259 purchased from Umicore. 260

The 50 preliminary cyclic voltammetry (CV) scans are 261 recorded at a 500 mV/s between 0.025 and 1.5 V. An 262 additional 3 CV scans are run at a rate of 20 mV/s between 263 0.025 and 1.0 V. The 50 conditioning CV cycles are then 264 collected from 1.2 to 1.8 V at 100 mV/s and 2500 rpm. Five 265 OER activity CVs are then recorded at 20 mV/s. The short- 266 term stability of selected samples is evaluated at 1.8 V and 267 2500 rpm. 268

Membrane Electrode Assembly. The fabrication of the 269 membrane electrode assembly (MEA), the cell integration and 270 testing have been described elsewhere.³⁹ In brief, cathode half- 271 cell catalyst-coated membranes using Nafion N117 (Ion 272 Power) were prepared as described earlier on a Sono-Tek 273 Exactcoat spray coater to achieve 0.3 mg/cm² Pt loading (45.6 274 wt % Pt/C, Tanaka). Carbon gas-diffusion layers with a 275 microporous layer (29 BC, Sigracet) were used on the cathode. 276 The spray coating is not suitable for TiO₂ supported anode 277 catalysts, since the supported catalyst particles precipitate 278 readily on the syringe, delivery tubes, and nozzle head.¹¹ Thus, 279 a modified slot-die method was utilized.⁴⁰ In brief, 1 mL of 280 water, catalyst, and Nafion (11.6 wt % of the catalyst) were 281 added in a centrifuge tube and sonicated using a probe 282 sonicator. The ink was then immediately cast on the N117 283 membrane secured in place over the vacuum hot plate of the 284 Sono-Tek Exactcoat at 50 °C. The catalyst loading was 285 determined using an in-house prepared calibration curve based 286 on XRF intensity counts. Titanium porous transport layers 287 (Proton OnSite/NEL) were used on the anode after wet 288



Figure 2. SEM image (a) and corresponding EDS maps (b and c) of Pt-TiO₂-PC. SEM image (d) and EDS maps (e and f) of Au-TiO₂-PC. TEM image (g), high-resolution TEM (h) and Pt particle size distribution of 100 Pt nanoparticles (i) of Pt-TiO₂-PC. High-angle annular dark-field (HAADF) image (j) and EDS maps (k-m) of Ir-Pt-TiO₂-PC-ann. All scale bars in (a-f) are 0.5 μ m and scale bars in (j-m) are 40 nm. EDS spectra corresponding to EDS maps above are in Figures S5 and S6.

289 proofing with 3.5% PTFE (DISP 30, Ion Power). Ten mil and
290 7 mil PTFE sheets (McMaster-Carr) were used to seal around
291 the porous-transport and gas-diffusion layers, respectively.

Cell Testing. For the cell testing, liquid water at 80 °C was pumped on the anode side at 100 mL/min and N_2 at 200 sccm and 100% relative humidity on the cathode side. The cell temperature was monitored on the cathode end plate and the water temperature at the anode outlet. Chronopotentiometric (CA) steps were collected for 3 min from 25 to 5000 mA total current at various intervals after conditioning the cell at 80 °C generate polarization curves. A multichannel potentiostat (VSP300) from BioLogic equipped with electrochemical impedance spectroscopy (EIS) and a 10 A booster was used for all electrochemical cell tests. EIS data were collected from 304 10 mHz to 200 kHz.

305 **RESULTS AND DISCUSSIONS**

³⁰⁶ Photoreductions of chlorometallic salts were used to prepare ³⁰⁷ Au-TiO₂-PC and Pt-TiO₂-PC CCSs (Figure S2). Analogous ³⁰⁸ materials were also prepared using the WI method followed by ³⁰⁹ thermochemical reduction (M-TiO₂-WI) using the same ³¹⁰ precursor chemicals.⁴¹ Ir was deposited on all the M-TiO₂ ³¹¹ materials using the WI method to generate Ir-(Au/Pt)-TiO₂-³¹² (PC/WI). The target deposition of each PGM metal on TiO₂ ³¹³ is 25 wt % except for the control Ir-TiO₂-WI, where we instead ³¹⁴ prepared 50 wt % Ir without the intermediate conductive layer ³¹⁵ (Table S1). The relative ratios of each element post-synthesis ³¹⁶ were determined using XRF (Figures S3 and S4). Irrespective ³¹⁷ of the precursor ratio or the irradiation duration, the Pt loading ³¹⁸ did not exceed 18 wt %, indicating that the photoreduction synthesis method utilized here is self-terminating. To 319 investigate the effects of thermal annealing on physical 320 properties as well as electrocatalysis, the materials were 321 annealed at 400 °C for 2 h. The thermally treated samples 322 are designated as "ann" hereafter (Table S1). 323

To probe the uniformity and morphology of the prepared 324 catalysts, SEM, TEM, and the corresponding EDS elemental 325 maps were collected. The SEM (Figure 2a), corresponding 326 f2 EDS maps (Figure 2b,c), and EDS spectrum (Figure S5) show 327 that Pt is uniformly deposited on the TiO₂ surface when the 328 conductive Pt layer is deposited using the PC method. As 329 apparent from the TEM image in Figure 2g,h, Pt nanoparticles, 330 roughly 2-3 nm in size (Figure 2i), deposit evenly on the TiO₂ 331 surface when Pt is deposited on TiO₂ using the PC method. 332 Furthermore, the Pt nanoparticle distribution is maintained 333 after Ir deposition and thermal annealing (Figure 2j-m and 334 Figure S6). Conversely, a non-uniform Au distribution in Au- 335 TiO₂-PC is observed (Figure S7), with some areas of 336 homogeneous coverage (Figure 2d-f). The EDS spectra for 337 Au-TiO₂-PC in Figures S7 and S8 indicate lower loadings of 338 Au when compared to the Pt EDS spectrum in Figure S5 even 339 though the starting precursor ratio for both Au and Pt was 25 340 wt %. 341

Half-Cell Characterization. Using RDE, we probed the 342 electrochemical activity of the CCS electrocatalysts with and 343 without thermal treatment compared to the commercial 344 baseline Umicore (IrO₂-TiO₂). The *iR* corrected OER activity 345 in 0.1 M HClO₄ (Figure 3), gauged by onset potentials, was 346 f3 highest for Ir-Pt-TiO₂-PC > Ir-Au-TiO₂-PC > Umicore > Ir- 347 TiO₂-WI, while the geometric current density at 1.8 V was 348 highest for Ir-Au-TiO₂-PC > Ir-Pt-TiO₂-PC ~ Umicore > Ir- 349



Figure 3. OER activities (*iR* corrected) of Ir catalysts supported on TiO₂. (a) Activities normalized to a geometric surface area of the working electrode. (b) OER activities at 1.8 V normalized to mg of Ir. (c) CA plots at 1.8 V. (d) Tafel plots (dotted lines) with calculated Tafel slopes (mV/ dec). The electrolyte is 0.1 M HClO₄, the working electrode disc and counter electrode wire are gold, and the reference electrode is a dihydrogen electrode. The working electrode is rotated at 2500 rpm, and the scan rate is 20 mV/s. Total catalyst loadings on the working electrodes are 3.49 μ g_{Ir}/cm².

350 TiO₂-WI. Overall, annealing was shown to have a deleterious 351 effect on Ir-Au-TiO₂-PC and Ir-TiO₂-WI; however, Ir-Pt-TiO₂-352 PC-ann retained a substantial amount of its activity even 353 slightly exceeding the commercial Umicore IrO_2 -TiO₂ catalyst. 354 Retention of excellent activity after oxidation is critical for 355 OER catalysts intended for PEMWE because they are 356 employed in highly oxidizing electrochemical environments. 357 CCSs without Ir deposition do not show significant OER 358 activities as expected (Figure S9).

While geometric OER activity of Ir-Pt-TiO₂-PC and 359 360 commercial Umicore catalyst are similar at 1.8 V (Figure 361 3a), the benefit of employing a Pt conductive layer is 362 demonstrated when the activities are normalized to an Ir 363 mass loading (Figure 3b). At 1.8 V applied potential, Ir-Pt-TiO₂-PC generates 1.02 A/mg_{Ir} compared to 0.42 A/mg_{Ir} for 364 the Umicore catalyst. This represents a 141% improvement in 365 366 mass-normalized OER activity with a 42% lower PGM content (75 wt % Ir in Umicore catalyst vs 25 wt % Ir and 18 wt % Pt 367 in Ir-Pt-TiO₂-PC). Even after thermal oxidation, the Ir-Pt-368 TiO_2 -PC-ann catalyst generates 0.59 A/mg_{Ir} activity, that is, a 369 370 39% improvement in Ir mass-normalized OER activity. The 371 utility of a uniform and thermally stable conductive layer is also 372 highlighted when Ir-Pt-TiO₂-PC is compared to Ir-Au-TiO₂-373 PC and Ir-TiO₂ catalysts; the OER activities of those two 374 systems drop drastically upon thermal annealing. The mass 375 normalized activities in Figure 3b are comparable to recently 376 reported Ir catalysts supported on TiO₂ supports (Table S2). 377 The mass normalized activities are still a magnitude lower than 378 the state-of-the-art unsupported³⁸ and antimony-doped tin 379 oxide supported Ir catalysts.⁹ However, activities of even the

commercial catalysts can vary widely depending on the ink 380 preparation methods and electrochemical conditions. Each 381 catalyst needs to be optimized for an ionomer to catalyst ratio, 382 sonication/stirring/shearing, and other catalyst ink preparation 383 parameters.³⁸ 384

When the commercial catalysts and as-synthesized Pt-CCS 385 supported materials are subjected to chronoamperometry 386 (CA) at 1.8 V to assess initial performance, it is observed 387 that the commercial Umicore catalyst initially generates higher 388 geometric OER activities, but the activities converge within 5 389 h. Critically, the synthesized Ir-Pt-TiO₂-PC catalysts generate a 390 similar activity after 5 h of CA to the commercial catalyst while 391 containing 33% less PGM loading. When the as-prepared and 392 annealed Ir-Pt-TiO₂-PC catalysts are compared (Figure 3c), 393 the activity of as-prepared catalyst is significantly higher 394 initially but drops precipitously to reflect that of the annealed 395 catalyst within 30 min. The Ir-Au-TiO₂-PC catalyst, initially, 396 also shows the highest activity at 1.8 V applied potential, but 397 becomes inactive within minutes. Longer-term stability studies 398 including online catalyst dissolution measurements and 399 operando characterization of electrocatalysts are deemed 400 beyond the scope of the current investigations. 401

From the Tafel slopes in Figure 3d, it is apparent that the 402 commercial catalyst (Umicore) and as-synthesized Ir-TiO₂-WI 403 and Ir-Au-TiO₂-PC exhibit faster OER kinetics (65 mV/dec) 404 in comparison to Pt-containing CCS supported catalysts (81–405 190 mV/dec). Annealed Ir-TiO₂ and Ir-Au-TiO₂ are not 406 included in the Tafel analysis since the activity is negligible in 407 the relevant potential window (Figure 3a). The Tafel slopes of 408 Pt-containing CCS supported catalysts intriguingly show two 409

410 distinct regions. At low overpotentials (<250 mV), both as-411 synthesized and annealed Ir-Pt-TiO₂ catalysts show high Tafel 412 slopes (>148 mV/dec), indicating slow kinetics but low onset 413 potentials. At higher overpotentials (>250 mV), faster kinetics 414 (81 mv/dec) predominate OER activity. Previous investiga-415 tions on Ir and Ru via experimental and theoretical analysis 416 have ascribed the "kink in Tafel slopes" to either different 417 active sites or catalyst surface reorganization at different 418 applied potentials.^{42,43} These results may also indicate that the 419 presence of the interfacial layer modulates the catalyst active 420 site. Generally, lower Tafel slopes indicate faster kinetics and 421 better electrochemical performance. Our observations here 422 indicate that other effects may be affecting overall performance 423 more dominantly than those encompassed in Tafel slopes. 424 Further optimizations to enhance the OER kinetics coupled 425 with the low onset potentials hold significant promise to boost 426 overall activity of low Ir-loading CLs using supported catalysts. Structure-Activity Relationships. In order to rationalize 427 428 the trends in electrochemical activity and to understand the 429 structure-activity relationships, we thoroughly characterized 430 the catalysts for electronic conductivity, crystallinity, and 431 surface composition. Conductivities of the prepared catalysts 432 (Figure 4) show that the presence of Pt conductive layer is



Figure 4. Electronic conductivities of various catalysts supported on TiO_2 . Umicore is the commercial IrO_2 - TiO_2 catalyst, and "ann" represents thermally annealed materials.

433 responsible for maintaining a relatively high conductivity 434 during thermal annealing. The conductivities are similar to 435 previous reports on various supported IrO_x catalysts prepared 436 using the Adams fusion method (Table S4).^{26,44,45} As 437 synthesized, the Ir-TiO₂-WI, Ir-Pt-TiO₂-PC, and Ir-Au-TiO₂-438 PC have similar conductivities (37–44 S/cm). However, upon 439 annealing, the conductivities of Ir-TiO₂-WI-ann and Ir-Au-440 TiO₂-PC-ann decrease dramatically to 6 and 18 S/cm, 441 respectively, but the conductivity of Ir-Pt-TiO₂-PC-ann slightly 442 improves to 44 S/cm. The observations may indicate that Ir 443 and Au agglomerate rapidly under thermal annealing 444 conditions (Figure S7), and they are weakly adsorbed on the 445 TiO₂ surface. This leads to the dramatic OER activity 446 reduction observed in Figure 3a,b. Pt however is strongly 447 bound to the TiO₂ surface, retains conformal homogeneous 448 distribution (Figure 2), and maintains a relatively high OER 449 activity post-annealing.

450 XRD was used to investigate the crystallinity of the CCSs 451 and catalysts. For the synthesized Pt-TiO₂-PC CCSs (Figure 452 5a), only TiO₂ peaks are visible in the XRD, suggesting that 453 the PC deposition method leads to the synthesis of either an amorphous Pt layer or extremely small Pt nanoparticles. This is 454 in agreement with the observed 2–3 nm crystalline Pt 455 nanoparticles (Figure 2). Conversely, for the Pt-TiO₂-WI, 456 the Pt peaks such as the Pt (111) at 39.6° are apparent. When 457 Ir is deposited to prepare Ir-Pt-TiO₂-PC, peaks corresponding 458 to Ir and TiO₂ are observed, but the Pt peaks remain absent. 459 These observations indicate that Pt deposited on TiO₂ via the 460 PC method does not aggregate under thermochemical 461 reduction conditions during Ir deposition via the WI method 462 to prepare Ir-Pt-TiO₂-PC. Unlike the Pt-TiO₂-WI and Ir-Pt- 463 TiO₂-WI systems where the Pt (111) peak at 39.6° is apparent, 464 the peak is absent for both Pt-TiO₂-PC and Ir-Pt-TiO₂-PC. 465

Conversely to the Pt CCSs, the XRD peaks for Au are 466 prominent irrespective of the synthesis method (Figure 5b). 467 The Au (111) peak at 38.1° is wider and less prominent in Ir- 468 Au-TiO₂-PC than in Ir-Au-TiO₂-WI, indicating that the Au 469 crystallite sizes are smaller from the PC method than from the 470 WI method. However, as determined by electron microscopy 471 (Figure S7) and visual observations, the Au precursor is 472 photosensitive, and some portion of it is readily reduced upon 473 exposure to ambient light without semiconductor-assisted 474 photoreduction. It is likely that the Au peaks in Ir-Au-TiO₂-PC 475 are from the initial precipitation rather than PC reduction on 476 TiO₂ surface. However, akin to Ir-Pt-TiO₂-PC, the Au peaks in 477 Au-TiO₂-PC do not show significant aggregation or crystallite 478 size coarsening during the Ir deposition via the WI method.

Curiously, Ir-Pt-TiO₂-PC does not show Pt or Ir oxidation 480 during thermal treatment, as apparent from Figure 5c, with no 481 significant peaks corresponding to iridium oxides (IrO_r) or 482 platinum oxides (PtO_x). However, the Au and Ir are oxidized 483 to Au₂O₃ and IrO₂ for Ir-Au-TiO₂-PC-ann. The observations 484 indicate that the Pt interlayer may have a stabilizing effect on Ir 485 with respect to oxidation, and the bulk of Pt itself does not 486 oxidize. The oxidation and aggregation lead to both 487 conductivity reduction (Figure 4) and activity loss (Figure 488 3) for Ir-TiO₂-WI and Ir-Au-TiO₂-PC-ann, but Ir-Pt-TiO₂-PC 489 is thermally stable and maintains high conductivity and 490 activity. Particle size analysis shows that crystallite sizes 491 generally increase upon annealing (Table S3). The Pt CCS 492 supported Ir crystallite size increases from 10 to 28 nm, while 493 the crystallite size of the commercial supported Ir catalyst is 9 494 nm. Thus, some of the activity loss can be attributed to 495 agglomeration and loss in the active surface area. 496

Collectively, the XRD (Figure 5), microscopy (Figure 2), 497 and electrochemical activity trends (Figure 3) also point to a 498 subtle difference in how Au and Pt reductions and depositions 499 may proceed. By the eye, the Au precursor decomposes rapidly 500 when exposed to ambient light, forming a precipitate. 501 However, the slow evolution of color (darkening) during PC 502 irradiation indicates that some Au precursor remains in the 503 solution and is reduced at a much slower rate to metallic Au 504 nanoparticles under PC irradiation (Figure S2). These Au 505 nanoparticles agglomerate during the Ir deposition via the WI 506 method to prepare Ir-Au-TiO2-PC and thermal treatment to 507 generate Ir-Au-TiO2-PC-ann. We chose not to employ 508 strategies such as the use of surfactants or dilute precursors 509 to prevent Au nanoparticle formation and aggregation due to 510 the associated complications with post-synthesis purification 511 steps. The Pt deposition in contrast occurs by photoreduction 512 at the TiO₂ surface, that is, semiconductor-assisted photo- 513 driven reduction to form Pt-TiO2-PC. The process thus leads 514 to Pt nanoparticles evenly distributed on a TiO₂ surface, and 515 strong adsorption to the substrate prevents Pt nanoparticle 516



Figure 5. XRD diffraction patterns for: (a) Pt coated CCSs (blue) and Ir supported on CCSs (red). (b) Ir supported on Au coated CCSs and (c) effects of thermal annealing in air for Ir deposited on CCSs synthesized via PC reduction. Diffraction pattern of as-obtained TiO_2 nanoparticles (TiO_2) is included in (a), and all the reference patterns are accompanied by corresponding PDF numbers.



Figure 6. XPS spectra of Ir catalysts supported on CCSs synthesized using a photochemical reduction method. (a-d) Ir-Pt-TiO₂-PC, (e-h) Ir-Pt-TiO₂-PC-ann, (i-l) Ir-Au-TiO₂-PC, and (m-p) Ir-Au-TiO₂-PC-ann. The spectra are as collected without background correction.

 $_{517}$ aggregation during WI reduction to prepare Ir-Pt-TiO₂-PC and $_{518}$ thermal treatment steps to prepare Ir-Pt-TiO₂-PC-ann.

⁵¹⁹ By XPS (Figure 6), it is apparent that the majority of the ⁵²⁰ near surface Ir deposited on Pt CCSs is IrO_2 with some ⁵²¹ metallic (Ir^0) characteristic, apparent from the core Ir 4f ⁵²² binding energies (i.e., Ir 4f_{7/2} at 61.9 eV and Ir 4f_{5/2} at 64.9 eV,

f6

respectively).^{46,47} The XPS spectrum for Ir-Pt-TiO₂-PC does $_{523}$ not change before and after annealing in air, as apparent from $_{524}$ the Ir 4f region (Figure 6a,e). There is, however, an increased $_{525}$ level of Ir oxidation after annealing for Ir-Au-TiO₂-PC, as $_{526}$ apparent from the change in Ir 4f spectra in Figure 6i,m.⁴⁸ For $_{527}$ the Ir deposited on Au CCSs (Figure 6i,m) and bare TiO₂ 528



Figure 7. (a) Electrolyzer polarization curves using annealed Ir-Pt-TiO₂ and commercial Umicore catalysts at 1.0 mg/cm² PGM on the anode, 0.3 mg/cm² Pt (Pt/C) on the cathode, and liquid water feeds at 80 °C and N117 membranes. (b) Tafel plots of *iR* free polarization curves with the corresponding equations in the kinetics region.

529 (Figure S10), the Ir is mostly oxidized (IrO₂) with Ir $4f_{7/2}$ at 530 63 eV and Ir $4f_{5/2}$ at 66 eV. In agreement with the XRD 531 (Figure 5c), XPS confirms that Ir deposited on Pt CCS is more 532 metallic in nature relative to Ir deposited on Au CCS. Furthermore, the Ti 2p peaks are at lower binding energies for 533 534 Ir-Pt-TiO₂-PC before and after annealing (Figure 6b,f), in 535 comparison to, both, Ir-Au-TiO₂-PC (Figure 6j,n), and for Ir- TiO_2 (Figure S10). Similarly, the O 1s peaks for Ir-Pt-TiO₂-PC 536 537 also appear at about 1 eV lower than that of Ir-Au-TiO₂-PC and Ir-TiO₂-WI. However, there is a significant change in the 538 539 Pt 4f spectrum for Ir-Pt-TiO₂-PC. There is an increase in Pt²⁺ $_{540}$ (74 eV) with respect to Pt^{0} (72 eV) after annealing. 49 Au 4f 541 peaks show a slight shift to lower binding energy upon 542 annealing as the full width at half-maximum gets smaller 543 (Figure 6k,o).

The difference in the minima between the Ir 4f peaks in 544 545 Figure 6i for Ir-Au-TiO₂ compared to that of Ir-Pt-TiO₂ 546 (Figure 6a) has previously been rationalized as the interference 547 from the Ti 3s peak by Aguilar-Tapia et al.⁵⁰ From the 548 comparison of the XPS spectra, it is evident that the direct 549 interaction between Ir and TiO₂ leads to higher binding energy shifts for Ir and Ti, while an interlayer of Pt prevents Ir and Ti 550 from influencing each other's binding energies. The observa-551 552 tions from XPS further confirm that Au does not form a 553 uniform layer on TiO₂ to act as an interlayer between Ir and 554 TiO₂. Ir-Au-TiO₂-PC behaves more akin to Ir-TiO₂ than Ir-Pt-555 TiO2-PC during thermal annealing. Additionally, there is 556 evidence of a charge transfer from Ir to Pt, which leads to Pt 557 itself oxidizing during annealing (Figure 6c,g) and con-558 sequently Ir retaining a metallic character.^{46,51} In agreement 559 with previous studies of Au-IrO2 composites, Au on the 560 contrary leads to increased Ir oxidation since the binding energies of Au 4f peaks shift slightly lower.⁵² The stability of 561 the Ir surfaces deposited on Pt-CCSs evidenced from XPS 562 spectra in Figure 6 directly correlate to the retention of high 563 OER activities in Figure 3. Furthermore, the drastic loss in the 564 565 OER activities of Au-CCS and TiO₂ supported Ir upon 566 thermal annealing highlights the utility of using a stable conformal conductive layer between the OER Ir catalysts and 567 568 the non-PGM TiO₂ cores.

569 **PEMWE Performance.** As a preliminary proof-of-concept 570 demonstration for catalyst integration into a MEA, the most 571 promising CCS catalyst (Ir-Pt-TiO₂-PC-ann) was integrated into a PEMWE cell with 1.0 mg/cm² PGM loadings on the 572 anode and 0.3 mg/cm² Pt (Pt/C) on the cathode and 573 compared with the commercial TiO₂ supported Umicore 574 catalyst at the same anode catalyst loadings. The balance of cell 575 components, the membrane, the pretreatment conditions, and 576 the testing conditions are described elsewhere.³⁹ From the 577 polarization curves at 80 °C in Figure 7a, it is apparent that the 578 f7 Ir-Pt-TiO₂-PC-ann catalyst outperforms the commercial 579 Umicore catalyst. It must also be emphasized that the CCS 580 supported catalysts are yet to be optimized for CL integration 581 and MEA performance, which is beyond the scope of this 582 work. Additionally, it is also well established that each OER Ir 583 catalyst requires a unique set of catalyst ink preparation and 584 catalyst layer fabrication methods to perform optimally.⁵³ 585 Nevertheless, the CCS supported catalysts outperform 586 previously reported TiO₂ supported Ir catalysts assessed 587 under similar testing conditions after optimization (Table 588 S4).^{25,54,55} However, as with half-cell activities, the MEA 589 performances are still significantly inferior to state-of-the-art 590 unsupported IrO_x catalysts (Table S5).¹⁵ Ir-Pt-TiO₂-PC-ann 591 achieves 1 A/cm² activity at 1.87 V, while the Umicore catalyst 592 requires 1.89 V.

High-frequency resistances (HFRs), which account for 594 ohmic losses of the entire cell, were used to generate iR- 595 corrected polarization curves in Figure 7a.⁵⁶ The Tafel analysis 596 in Figure 7b indicates that the OER kinetics of Ir-Pt-TiO₂-PC 597 are superior (108 mV/dec) to the Umicore catalyst (120 mV/ $_{598}$ dec). The trend is in contrast to half-cell observations in Figure 599 3d, which showed lower Tafel slopes for Umicore catalysts (65 600 mV/dec) than for Ir-Pt-TiO₂-PC (81 mV/dec). The $_{601}$ observation re-emphasizes that RDE performances are not 602 always translatable to cell performance. The HFR-corrected 603 polarization curves (Figure 7a) also allow a MEA performance 604 comparison among various membrane thicknesses. From Table 605 S5, it is apparent that the CCS supported catalysts generate 606 comparable activities to other oxide supported IrO_x catalysts at 607 1 A/cm² under similar conditions and cell configuration. As 608 with the half-cell analysis in Figure 3 however, further 609 optimizations in catalyst structure, ink composition, and 610 catalyst integration methods are necessary to match the 611 performances of the state-of-the-art catalysts and catalyst 612 layers. 613

CL thickness and porosity also influence the performance by 614 615 improving transport. Cross-sectional SEM of the MEA 616 illustrates that the CL thickness increased and the porosity 617 was manipulated by employing CCS supports (Figure S11). 618 The anode CL thickness is comparable to previous reports 619 using 2–3 mg/cm² PGM catalyst loadings.^{22,40,54,57–59} Further 620 investigations are underway to understand the contributions of 621 the kinetics, mass transport, catalyst layer fabrication methods, 622 and catalyst ink compositions in electrolyzer performance 623 using CCS integrated CLs.

624 CONCLUSIONS

625 We have demonstrated a facile photoreduction method to 626 generate a conformal layer of platinum nanoparticles on TiO₂, 627 forming conductive layer coated supports (CCSs). The 628 conductive intermediate layer prevents bulk iridium oxidation 629 in the Ir-Pt-TiO₂ catalyst during annealing and improves the 630 composite electrocatalyst conductivity at a significantly lower 631 precious group metal (PGM) loading (39 wt %) in comparison 632 to a TiO₂ supported commercial catalyst with 75 wt % PGM 633 loading. Photoreduction generated a conformal platinum layer 634 that is strongly bound to the semiconductor surface and is 635 stable in thermochemical and electrochemical environments. 636 The presented X-ray photoelectron spectroscopy, X-ray 637 diffraction, and electron microscopy images show significant 638 opportunities to further engineer highly active and stable 639 surfaces, interfaces, and active sites utilizing the conductive 640 interlayer in comparison to bare TiO₂ supports. The half-cell 641 oxygen evolution reaction (OER) activities show that the Pt-642 CCS supported iridium generates a 141% higher OER mass 643 activity than the commercial TiO₂ supported catalyst from 644 Umicore. Furthermore, the single-cell performance of the CCS 645 supported catalyst reveals a superior performance to other 646 supported catalysts reported thus far, even prior to any 647 integration and fabrication optimizations. Our investigations 648 also reaffirm that translating the half-cell catalyst performance 649 to device level performance requires considerable optimization 650 during the catalyst layer fabrication and device integration. We 651 expect future investigations utilizing CCSs to improve the 652 electrolyzer performance significantly from CCS supported 653 materials and device optimization via synthetic modifications 654 as well as targeted catalyst ink preparation for each catalyst and 655 cell fabrication methodologies.

656 ASSOCIATED CONTENT

1 Supporting Information 657

658 The Supporting Information is available free of charge at 659 https://pubs.acs.org/doi/10.1021/acscatal.0c03098.

Synthesis routes, table of catalysts used in the 660 investigation, additional SEM images and EDS maps, 661 EDS spectra, XPS plots, XRF data, OER data 662 corresponding to CCS only materials, and table of 663 MEA performance of TiO₂ supported Ir catalysts from 664 recent investigations (PDF) 665

Video S1: Generation of conductive layer coated catalyst 666 substrates via photoreduction (AVI) 667

AUTHOR INFORMATION 668

669 Corresponding Author

Nemanja Danilovic – Energy Storage and Distributed Resources 670

- Division, Lawrence Berkeley National Laboratory, Berkeley, 671
- 672 California 94720, United States; Email: ndanilovic@lbl.gov

Autions

Authors	673
Yagya N. Regmi – Energy Storage and Distributed Resources	674
Division, Lawrence Berkeley National Laboratory, Berkeley,	675
California 94720, United States; Department of Chemistry,	676
Manchester Metropolitan University, Manchester M1 5GD,	677
United Kingdom; © orcid.org/0000-0001-6588-7683	678
Eden Tzanetopoulos – Energy Storage and Distributed	679
Resources Division, Lawrence Berkeley National Laboratory,	680
Berkeley, California 94720, United States; College of Chemistry,	681
University of California Berkeley, Berkeley, California 94720,	682
United States	683
Guosong Zeng – Chemical Sciences Division and Joint Center	684
for Artificial Photosynthesis, Lawrence Berkeley National	685
Laboratory, Berkeley, California 94720, United States	686
Xiong Peng – Energy Storage and Distributed Resources	687
Division, Lawrence Berkeley National Laboratory, Berkeley,	688
California 94720, United States	689
Douglas I. Kushner – Energy Storage and Distributed Resources	690
Division, Lawrence Berkeley National Laboratory, Berkeley,	691
California 94720, United States; 💿 orcid.org/0000-0002-	692
3020-7737	693
Tobias A. Kistler – Chemical Sciences Division and Joint Center	694
for Artificial Photosynthesis, Lawrence Berkeley National	695
Laboratory, Berkeley, California 94720, United States; Walter	696
Schottky Institute and Physics Department, Technische	697
Universität München, 85748 Garching, Germany;	698
© orcid.org/0000-0001-6458-8024	699
Laurie A. King – Department of Chemistry, Manchester	700
Metropolitan University, Manchester M1 5GD, United	701
<i>Kingdom;</i> orcid.org/0000-0002-0772-2378	702
Complete contact information is available at:	703
https://pubs.acs.org/10.1021/acscatal.0c03098	704

Author Contributions

N.D. and Y.R. conceived the project and secured funding. Y.R., 706 E.T., L.K., and N.D. conceived synthesis routes and/or 707 synthesized the materials. Y.R. and E.T. conducted XRD and 708 SEM/EDS. Y.R., E.T., and X.P. conducted STEM/EDS. Y.R., 709 G.Z., and L.K. collected and analyzed XPS. Y.R., E.T., and 710 N.D. conducted and analyzed electrochemical measurements. 711 T.K. operated the solar simulator lamp and associated camera. 712 D.K. designed the conductivity setup. Y.R. and E.T. composed 713 the manuscript, and all the authors edited the written work 714 under the leadership of N.D. 715

Notes

The authors declare no competing financial interest.

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