Single PEDOT Catalyst Boosts CO2 Photoreduction Efficiency

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ABSTRACT: Atmospheric pollution demands the development of solar-driven photocatalytic technologies for the conversion of CO2 into a fuel; state-of-the-art cocatalyst systems demonstrate conversion efficiencies currently unattainable by a single catalyst. Here, we upend the status quo demonstrating that the nanofibrillar conducting polymer poly(3,4-ethylendioxythiophene) (PEDOT) is a record-breaking single catalyst for the photoreduction of CO2 to CO. This high catalytic efficiency stems from a highly conductive nanofibrillar structure that significantly enhances surface area, CO2 adsorption and light absorption. Moreover, the polymer’s band gap is optimized via chemical doping/dedoping treatments using hydrochloric acid, ammonia hydroxide, and hydrazine. The hydrazine-treated PEDOT catalyst exhibits 100% CO yield under a stable regime (>10 h) with a maximum rate of CO evolution (3000 μmol gcat\(^{-1}\) h\(^{-1}\)) that is 2 orders of magnitude higher than the top performing single catalyst and surpassed only by three other cocatalyst systems. Nanofibrillar PEDOT provides a new direction for designing the next generation of high-efficiency photoreduction catalysts.

**INTRODUCTION**

Excessive fossil fuel combustion has drastically increased net global carbon dioxide (CO2) influx into Earth’s atmosphere, inducing climate change and worsening air quality, human health, and energy security.1,2 The critical need for mankind to counteract increasing atmospheric greenhouse gas concentrations demands intensive research to develop processes that attenuate CO2 emissions.3 Photoreduction of CO2 is one promising approach where abundant and sustainable solar energy is utilized, with the aid of semiconductor photocatalysts, to reduce CO2 into chemical fuels.4,5 In a photocatalytic system, electrons/holes generated by light absorption migrate toward catalytically active sites where they react with adsorbed CO2 molecules. Catalytic efficiency relies on generation and separation of photogenerated charge carriers while minimizing recombination, as well as on efficient CO2 adsorption and light absorption.6,7 Numerous efforts have optimized structure and composition of photocatalysts by increasing surface area, creating surface defects on graphene, introducing nanocrystals to enhance visible light absorption and applying metal/metal oxide cocatalysts, such as Ni, TiO2, and ZnO, to facilitate charge separation. However, these cocatalysts suffer from recombination of charge carriers and band gap mismatch leading to poor conversion efficiencies.8,9 As opposed to a single catalyst, delivery of photogenerated charge carriers between cocatalysts of different band gaps must overcome the high energy barrier of out-of-plane Ohmic or Schottky contact during an intramolecular cascade.10 Although researchers have modified cocatalyst systems by tuning the ratio of components to precisely match the band gap, costly fabrication techniques and low yields highlight the importance of developing a more favorable single catalyst alternative.

Among all single catalysts, conducting polymers (CPs) are a new class of organic semiconductors primed to become the next generation of multifunctional photocatalysts; this is because the π-conjugation backbone in CPs enhances CO2 absorption via π−π interactions with delocalized p-conjugated electrons from CO2 molecules. As semiconductors, CPs possess a wide light absorption range and photostability in the ultraviolet–visible–near-IR (UV–vis–NIR) region that is easily controlled by doping/dedoping treatments and further enhanced by wide-ranging nanostructures.11 Growing interest in CPs has prompted recent studies on CO2 photoreduction using polyaniline (PANI) and polypyrrole (PPy),12,13 unfortunately the low electrical conductivity of these polymers results in low photoreduction efficiency. Poly(3,4-ethylenedioxythiophene) (PEDOT) exhibits high mobility of photogenerated electrons/holes14 and is widely recognized as the most stable conductive CP (7600 S/cm) thus potentially serving as an ideal CO2 photoreduction catalyst. To the best of our knowledge, there is an absence of reports in the current literature regarding its CO2 photoreduction catalytic properties.

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Here, for the first time, we demonstrate that a nanofibrillar PEDOT film is an ideal single catalyst for CO$_2$ photoreduction. This film is deposited via our previously reported rust-based vapor-phase polymerization 15 resulting in a homogeneous PEDOT coating of high surface area possessing a nanofibrillar morphology and characterized by high electronic conductivity (1200 S/cm). The CO$_2$ adsorption and light absorption properties are controlled by the polymer’s band gap via chemical doping and dedoping treatments using an acid (HCl) and base (NH$_4$OH or hydrazine), respectively. Our single catalyst results in a state-of-the-art CO yield rate (3000 μmol g$^{-1}$ h$^{-1}$) with 100% CO yield under stable regime (>10 h) representing 2 orders of magnitude higher than the best single catalyst reported and surpassed by only three other cocatalyst systems.

**RESULTS AND DISCUSSION**

A conformal PEDOT nanofibrillar coating is produced via rust-based vapor-phase polymerization 15 by initially sputtering a 60 nm coating of Fe$_2$O$_3$ oxidant precursor onto a robust polytetrafluoroethylene (PTFE) membrane (Figure 1A). A conformally packed oxide coating exhibits a yellow color in the solid state (Figure 1B) and when dissolved, this oxide serves as an oxidizing agent promoting the deposition of a homogeneous PEDOT coating. Polymer deposition is initiated by dissolving Fe$_2$O$_3$ at 140 °C using HCl vapor; dissolution liberates Fe$^{3+}$ ions that promote oxidative polymerization of EDOT vapor. Homogeneous PEDOT nanofibrillar coating supported on a PTFE membrane (E) Optical image of a PEDOT-coated PTFE membrane. The molecular and solid-state structure of a blue PEDOT-coated PTFE film (2 cm × 5 cm) (Figure 1E) is characterized via microscopy and spectroscopy. The polymer is uniformly integrated within a superhydrophobic substrate.

Photocatalyst morphology, surface area, and intrinsic chemical properties are vital parameters for CO$_2$ adsorption, light absorption, and CO$_2$ reduction efficiency. The molecular and solid-state structure of a blue PEDOT-coated PTFE film (2 cm × 5 cm) (Figure 1E) is characterized via microscopy and spectroscopy. The polymer is uniformly integrated within the PTFE membrane.

Figure 1. Schematic diagram of deposition of a nanofibrillar PEDOT film on a PTFE membrane and microscopic/spectroscopic characterization of a nanofibrillar PEDOT film. (A) Pristine porous PTFE membrane is (B) sputtered with a 60 nm Fe$_2$O$_3$ layer. (C) Rust-based vapor phase polymerization produces PEDOT nanofibers by (a) liberating Fe$^{3+}$ from Fe$_2$O$_3$ and triggering oxidative polymerization of (b) EDOT vapor. (D) Homogeneous PEDOT nanofibrillar coating supported on a PTFE membrane. (E) Optical image of a PEDOT-coated PTFE membrane. (F) Scanning electron micrograph (SEM) shows bulk morphology of a homogeneous PEDOT coating. (G) Close-up SEM reveals the free volume architecture between nanofibers. (H) Transmission electron micrograph (TEM) of a single nanofiber (aspect ratio ~ 50). (I) EDX maps of a nanofiber show an elemental composition consisting of C, S, O, and Fe. (J) PXRD confirms PEDOT’s polycrystalline structure with three characteristic sharp peaks.
the PTFE membrane (Figure S1) and scanning electron micrographs (SEM) of low (Figures 1F and S2) and high magnification (Figure 1G) reveal a free volume web-like architecture composed of spherical aggregates interconnected by nanofibers. This open nanofibrillar structure possesses a high surface area that enhances absorption of both light and CO2; a transmission electron micrograph (TEM) shows a 100 nm mean nanofiber diameter with an aspect ratio of 50 (Figure 1H). Elemental mapping via energy-dispersive X-ray spectroscopy (Figure 1I) shows a uniform distribution and strong signals for C, O, and S corresponding to PEDOT’s ethylenedioxy and thiophene ring whereas Fe signal disappears once PEDOT is purified using 6 M HCl, suggesting no Fe impurities.

Current–voltage (I–V) measurements, carried out to probe the electronic charge transport of a PEDOT film, demonstrate ohmic behavior (linear relationship) and a low resistance (large positive slope) (Figure S3) characteristic of a homogeneous and continuous percolation network that facilitates charge transport. Powder X-ray diffraction patterns (Figure 1J) further elucidate PEDOT’s charge transport as a function of its polycrystalline structure. Three characteristic peaks are observed at 6.4°, 13.1°, and 26.3° corresponding to lattice planes (100), (200), and (020), respectively. The first and second planes stem from lateral chain packing (x-axis) and the third is due to π−π stacking (y-axis). Low peak width values at half-height (0.4, 0.7, and 1.4) indicate a crystalline polymer structure responsible for enhanced charge transport.16

Four-point probe conductivity measurements (Figure S4) show a high conductivity (1200 S/cm) stemming from the homogeneous deposition of an ordered PEDOT structure.

PEDOT’s CO2 adsorption capacity is enhanced by doping/dedoping treatments with acid, base, and reducing agents that alter its surface energy.17 We investigate this property utilizing HCl (37%), NH4OH (30%), and hydrazine to dope/dedope a PEDOT film resulting in PEDOT–HCl (light blue), PEDOT–NH4OH (purple), and PEDOT–hydrazine (black) (Figure 2A) samples. Contact-angle measurements are carried out to study surface energy revealing that all PEDOT coatings decrease PTFE’s surface energy, while the hydrazine-treated PEDOT film exhibits the wetting degree angle at 60° (Figure 2B). PEDOT-coated membranes promote water molecule permeability confirmed by water content using TGA (Figure S5), SEM (Figure S6), and EDX (Figure S7); we suspect this is due to charged dopant effects. We gain insight into the impact of doping on chemical bonding using Raman spectroscopy. PEDOT’s peak (Figure 2C) at 1357 cm−1 corresponds to C=C π bonds, while peaks at 1437 and 1510 cm−1 correspond to symmetric and asymmetric C=C stretching. The symmetric C=C stretching band shifts toward 1426 cm−1 after doping with HCl.18 This shift toward higher wavenumbers corresponds to greater doping levels due to an increased ratio of quinoid vs benzoid structures in PEDOT and is consistent with previously reported behavior for other PEDOT films.19

The CO2 photoreduction properties of PEDOT films are evaluated using a photoreduction analysis system under atmospheric pressure and room temperature (Figure 2D). Three initial control tests are performed to establish the source of reduction products: the first test is conducted with He as the source gas, instead of CO2, the second test is performed in the absence of PEDOT, and the third test is carried out under a lack of illumination. Carbonaceous product is absent from all three control test results demonstrating that catalysis is unfeasible in the absence of CO2, PEDOT film, or light illumination. Notably, our gas–solid interface-based photo-
reduction system produces 100% CO as the main product because CO formation is kinetically favorable, requiring only a minimal number of electrons and protons for CO2 reduction. Electrons and holes generated by UV−vis−NIR light (250−950 nm) arrive at the photoactive PEDOT film interface (active site) (Figure 2E) facilitating simultaneous reduction of CO2 (electrons) and oxidation of H2O (holes). The PEDOT nanofibrillar morphology boosts light-harvesting because internal reflection of light is multiplied within the fibrillar nanostructure. Previous studies demonstrate that introducing Lewis basic sites on the surface of a catalyst enhances CO2 adsorption and activation.29 The enhanced CO2 uptake capacity exhibited by PEDOT−hydrazine catalysts at 80 °C. (B) UV−vis−NIR spectra prove that both a reducing agent and base enhance light absorption in the visible region partially converting polaronic PEDOT to its neutral state. (C) Ultraviolet photoelectron valence band spectra for PEDOT−HCl, PEDOT−NH4OH, and PEDOT−hydrazine samples predict the secondary electron edge and onset of the density of states. (D) Proposed mechanism of charge separation/transfer shows LUMO/HOMO band gaps for PEDOT−HCl, PEDOT−NH4OH, and PEDOT−hydrazine photocatalysts under a range of wavelengths (250−950 nm).

Figure 3. PEDOT mechanism for CO₂ photoreduction. (A) CO₂ adsorption capacity for PEDOT−HCl, PEDOT−NH₄OH, and PEDOT−hydrazine catalysts (250−950 nm) arrive at the photoactive PEDOT film interface (active site) (Figure 2E) facilitating simultaneous reduction of CO₂ (electrons) and oxidation of H₂O (holes). The PEDOT nanofibrillar morphology boosts light-harvesting because internal reflection of light is multiplied within the fibrillar nanostructure. Previous studies demonstrate that introducing Lewis basic sites on the surface of a catalyst enhances CO₂ adsorption and activation.29 The enhanced CO₂ uptake capacity exhibited by PEDOT−hydrazine catalysts at 80 °C. (B) UV−vis−NIR spectra prove that both a reducing agent and base enhance light absorption in the visible region partially converting polaronic PEDOT to its neutral state. (C) Ultraviolet photoelectron valence band spectra for PEDOT−HCl, PEDOT−NH₄OH, and PEDOT−hydrazine samples predict the secondary electron edge and onset of the density of states. (D) Proposed mechanism of charge separation/transfer shows LUMO/HOMO band gaps for PEDOT−HCl, PEDOT−NH₄OH, and PEDOT−hydrazine photocatalysts under a range of wavelengths (250−950 nm).

To study reduction rates among our catalysts, we investigate CO₂ adsorption, light harvesting properties and band gap structure through CO₂ uptake data, UV−vis−NIR and ultraviolet photoelectron spectroscopy (UPS). Adsorption of CO₂ on the catalytic surface is an important parameter for catalyst evaluation because it is the first step of the photocatalytic reaction. The CO₂ uptake data evaluated from TGA measurements (Figures 3A and S10) show the following trend: PEDOT−hydrazine (0.15) > PEDOT−NH₄OH (0.113) ≈ PEDOT−HCl (0.12), confirming that low surface energy enhances CO₂ adsorption. The π-conjugated structure of PEDOT attracts delocalized p-conjugated electrons from CO₂ molecules prompting CO₂ uptake on catalytic surface sites.29−30 The enhanced CO₂ uptake capacity exhibited by hydrazine-treated PEDOT may arise from increased interactions between quadrupolar CO₂ (Lewis acid) and depolarized oxygen of the ethylenedioxy ring (Lewis base) leading to greater CO₂ solubility and permeability in PEDOT-coated membranes.30

Another key factor for improving CO₂ photoreduction efficiency is light absorption because photons provide energy
for catalysis, UV–vis–NIR spectroscopy is utilized to assess charge carrier density as a function of chemical structure and characterize changes in molecular structure caused by doping. The UV–vis–NIR spectra of PEDOT (Figure 3B) are divided into three regions, that is, neutral, polaronic and bipolaronic. The wavelengths chosen for CO2 reduction (250–950 nm) lead to higher absorbance for dedoped PEDOT–hydrazine and PEDOT–NH4OH samples thereby promoting photogeneration of charge carriers as well as enhanced photocatalytic activity. The photothermal activity of PEDOT increases in temperature during photoreduction that enhances the reaction kinetics, diffusion of CO2, and desorption of CO2. Larger absorbance in the polaronic and bipolaronic regions for the PEDOT–HCl sample is observed because doping ions (Cl−) help delocalization of the electron cloud thereby increasing electrical conductivity.

To understand the effect of doping on charge carrier density and mobility for enhanced CO2 photoreduction, UPS is utilized for determining the valence band (VB), conduction band (CB), and Fermi level of a PEDOT catalyst (Figure 3C). Long tails in the VB spectrum of each PEDOT sample indicate different convoluted peaks and PEDOT–hydrazine in particular exhibits clear steps suggesting the presence of multiple hybridized states between VB and CB. The work function (Fermi level energy) of PEDOT is determined by calculating the difference between the secondary electron cutoff and the energy of an incident photon. The gap between the Fermi level and the VB maximum is the onset of photoemission for valence electrons and the CB is calculated from the VB maximum, Fermi level and band gap (Figures S11–S13). An energy level diagram shows that the CB level for all PEDOT catalysts meets CO2 reduction requirements by lying at potentials lower than that of CO2 redox couples (Figure 3D). Hydrazine reduction dedopes PEDOT, increases its Lewis basicity, makes the photoreduction process thermodynamically more favorable (Figure 3D) and results in higher CO2 adsorption thereby enhancing photocatalytic activity. PEDOT catalysts are excited by absorbing energetic photons and produce electron–hole pairs as opposed to conventional inorganic catalysts that only allow direct excitation of electrons from the VB to the CB. Further study is required to determine whether doping or dedoping changes the charge carrier density or charge carrier mobility. On the basis on our results, PEDOT–hydrazine possesses a narrower band gap requiring less excitation energy for promoting facile excitation, and the higher Fermi level also prevents the recombination process of photoexcited electrons and holes thereby facilitating photocatalytic activity.

**CONCLUSION**

In summary, we present a vapor phase-deposited nanofibrillar PEDOT coating that serves as a novel single catalyst for CO2 photoreduction. Our coating readily conforms to a substrate’s shape and when deposited on a superhydrophobic PTFE membrane, it’s characterized by high surface area and high electronic conductivity. The hydrazine-treated PEDOT catalyst demonstrates the highest CO production rate under 250–950 nm wavelengths. We achieve a state-of-the-art CO yield rate (3000 μmol gcat−1 h−1) with 100% CO yield under a stable regime (>10 h) surpassing the top performing single catalyst systems by 2 orders of magnitude. This work overcomes challenges associated with cocatalyst systems and opens up a new avenue for CO2 photoreduction utilizing nanofibrillar conducting polymers. Our single catalyst system represents a novel material serving as the next generation building block for developing future organic–inorganic cocatalyst systems with the potential to catapult conversion efficiencies to new levels.

**METHODS**

**Materials.** Fluoropore membrane was purchased from MilliporeSigma. 3,4-ethylenedioxythiophene (EDOT, 97%), chlorobenzene (99%), hydrochloric acid (37%), ammonia hydroxide (25%), and hydrazine hydrate (50%) were purchased from Sigma-Aldrich and used as received.

**Deposition of Nanofibrillar PEDOT Film.** A solid-oxidant precursor, Fe3O4, was sputter deposited over polytetrafluoroethylene (PTFE) membrane via physical vapor deposition (Kurt J. Lesker PVD 75 RF and DC). All syntheses were performed at 140 °C for 1.5 h in glass reactors, each containing a substrate, a reservoir with 20 μL concentrated hydrochloric acid (HCl) and a reservoir with 200 μL of a 1.56 M EDOT/chlorobenzene solution. After 1.5 h, the electrodes were immediately removed from the reactors and cooled at room temperature and purified with 6 M HCl.

**PEDOT Characterization.** Scanning electron micrographs and energy-dispersive X-ray spectra were collected using a JEOL 7001LVF FE-SEM. Transmission electron micrographs were obtained using a JEOL 2100 TEM by drop-casting a dispersion of PEDOT nanofibers on a TEM grid. Raman spectra were obtained using a Renishaw inVia confocal Raman spectrometer mounted on a Leica microscope with a 20x objective and 785 nm wavelength diode laser serving as an illumination source. A low power was necessary to mitigate heating of conducting polymer samples. A Bruker d8 Advanced X-ray diffractometer was utilized to collect powder X-ray diffractograms of pulverized samples at room temperature using a Cu Kα radiation source (λ = 1.5406 Å) and LynxEye XE detector (operating at 40 kV and 40 mA); the sample holder was rotated at 30 rpm with a scan step of 0.02°. Current–voltage (I–V) curves were obtained with a built-in-house 3D printed probe station using two gold needles 1.24 mm apart. Four-point probe sheet resistance measurements were carried out using a Keithley 2450 SourceMeter with a Signatone SP4 four-point probe head. Ultraviolet–visible–near-infrared (UV–vis–NIR) spectra were collected on a Cary 5000 UV–vis–NIR spectrophotometer. Solid PEDOT films were dispersed in HCl (37%), NH4OH, and hydrazine. X-ray photoelectron spectroscopy (XPS) was conducted on solid samples using a PHI 5000 VersaProbe II with an Al 1486.6 eV Mono-X-ray source at 51.3 W, a beam diameter of 100–200 μm and a 1 V neutralizer at 15 μA. Ultraviolet photoemission spectroscopy (UPS) measurements required a Physical Electronics 5000 VersaProbe II Scanning ESCA Microprobe and were performed to determine the valence band, conduction band and Fermi level position. He I (21.2 eV) ultraviolet source was used and a 5 V bias was applied to the sample to observe a clear secondary electron edge.

**Evaluation of PEDOT films for CO2 photoreduction.** Photocatalytic activity of the synthesized PEDOT films was evaluated using a photoreduction analysis system (Figure 3D) which was discussed in our previous publications. The photoreduction analysis system contains a lab-built continuous flow reactor, Xe arc lamp (Oriel 66021, Newport Co.) and gas chromatograph (GC, 6895N, Agilent Technologies, Inc.). For testing, the PEDOT film was deposited on glass and placed in a lab-built continuous flow reactor. High purity compressed
CO₂ was used as the source gas and its flow rate was fixed at 20 mL min⁻¹ by a mass flow controller (OMEGA engineering Inc.). The gas stream continuously passed through a water bubbler to generate a mixed flow of CO₂ and water vapor. Then, the mixed flow was introduced into the continuous flow reactor and a quartz window enabled positioning a sample vertically to face the light (inset photo of Figure 3E). The PEDOT film was illuminated by the Xe arc lamp, which was operated at 400 W within a UV–vis–NIR wavelength region (250–950 nm). The composition in the gas streams was continuously measured by a GC equipped with a porous layer open tubular (PLOT) capillary column (Supelco Carbobox-1010) and a thermal conductivity detector (TCD); helium was used as a carrier gas for the GC. In all experiments, the Xe arc lamp was initiated after the system was stabilized and approximately 3 mL min⁻¹ of gas were introduced to the GC. All experiments were performed at least in duplicates and no unexpected or unusually high safety hazards were encountered.

## ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscentsci.1c00712.

Photograph of a pristine PTFE film shows macroscale morphology, scanning electron micrograph of a PEDOT nanofibrillar film, I–V curves of unpurified and purified PEDOT films (purification is carried out via repeated washes in concentrated HCl), conductivity measurement carried out using a four-point probe station, TGA of pristine PTFE and of PEDOT-coated PTFE, scanning electron micrographs of PEDOT catalysts after treatment with HCl, NH₄OH, and hydrazine, energy-dispersive X-ray spectra of PEDOT catalysts after treatment with HCl, NH₄OH, and hydrazine, UV–vis–NIR measurement data after CO₂ photoreduction, time-dependent production rates of CO versus PEDOT and PEDOT–FeCl₃ catalysts under illumination (250–950 nm), TGA isotherm for CO₂ adsorption, ultraviolet photoelectron valence band spectra for PEDOT–HCl sample predict the secondary electron edge and onset of the density of states, ultraviolet photoelectron valence band spectra for PEDOT–NH₄OH sample predict the secondary electron edge and onset of the density of states, and ultraviolet photoelectron valence band spectra for PEDOT–hydrazine sample predict the secondary electron edge and onset of the density of states (PDF).

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**Author Contributions**

Y.D., S.J., M.K., P.B., and J.M.D designed the experiments. Y.D., S.J., and M.K. carried out syntheses, device fabrication, characterization, and data analysis. Y.D., S.J., and M.K. analyzed data and wrote the paper. Y.D., S.J., M.K., R.W., H.Y., P.B., and J.M.D contributed to the discussion and editing of the Research Article.

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**Notes**

The authors declare no competing financial interest.

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