ACS APPLIED ENERGY MATERIALS

Optimal Dye Sensitized Solar Cell and Photocapacitor Performance with Efficient Electrocatalytic SWCNH Assisted Carbon Electrode

Gurulakshmi Maddala, Meenakshamma Ambapuram, VijayaLaxmi Tankasala, and Raghavender Mitty*

Cite This: ACS Appl. Energy Mater. 2021, 4, 11225–11233

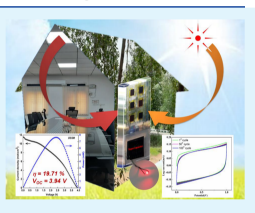


Article Recommendations

Λ	\mathbf{n}	\mathbf{n}	ГС	0
Α	U	L	ES	0
	<u> </u>	<u> </u>		_

III Metrics & More

ABSTRACT: The present work demonstrates the high photovoltaic power conversion efficiency (PCE) of 11.12% using single wall carbon nanohorn assisted carbon counter electrode based dye sensitized solar cells (DSSCs), which demonstrates a superior PCE compared to that of platinum (9.41%). This superior performance was a motivation to fabricate a dye sensitized solar module (DSSM) consisting of six series connected DSSCs arranged in a bifacial manner toward the application of building integrated photovoltaics. The DSSM demonstrated a remarkable, champion PCE of 19.71%. This high PCE, driven to fabricate an integrated device (photocapacitor), consists of a DSSM and a supercapacitor (SC). Upon two-sided illumination, the DSSM generated electrical power, and the same power is used for charging the supercapacitor. A working light emitting diode is demonstrated with discharge of the SC. The detailed fabrication strategies and results are discussed.



S Supporting Information

KEYWORDS: cost-effective carbon electrode, screen print, platinum-free, single wall carbon nanohorns, spray coat, dye sensitized solar cell, bifacial, DSSM, supercapacitor, photocapacitor

1. INTRODUCTION

The global energy requirement is crucial, and it can potentially be addressed through a clean energy conversion photovoltaic technology. Cost-effective, third-generation solar photovoltaics have become significant, so attention has been paid due to its easier fabrication process, environmental friendliness, and good energy conversion efficiency. The dye sensitized solar cell (DSSC) is identified as an affordable photovoltaic technology and has attracted more attention for its capability to deliver low-cost power generation.¹⁻³ DSSC consists of a photoelectrode (PE) of dye anchored TiO₂ nanoparticles, an electrolyte comprising redox couples (generally, I^-/I_3^-), and a counter electrode (CE). Counter electrodes play a critical role in achieving high power conversion efficiency (PCE). Developing a potential CE is essential, as CE catalysts have high conductivity, have good catalytic activity, are more stable, etc. Generally, a very thin platinum (Pt) layer will be deposited onto a conductive fluorine doped tin oxide (FTO) coated glass substrate which is used as the counter electrode, due to its extreme catalytic performance for tri-iodide reduction and good electrical conductivity.^{4,5} Due to its higher cost, rare availability leads to minimal large-scale production for power production. To overcome this, efforts have been made by researchers for the development of cost-effective catalysts, include carbonaceous materials, nitride, sulfides, etc., as promising catalysts materials.^{6–13} The present research focuses on a Pt-free, cost-effective CE as an alternative to the "champion" CE. The present study aims to develop an effective, potential CE through an easy preparation processes.

Numerous carbon-oriented materials like active carbon,¹⁴ carbon powder,^{15,16} carbon spheres,^{17,18} and fullerenes¹⁹ have been used as CEs which have shown remarkable progress. The performance of DSSCs fabricated with carbon-oriented CEs are lower compared to Pt based CEs. Improved electrocatalytic active sites through deposition of another material over CE catalysts could result in higher performance.²⁰ The single wall carbon nanohorn (SWCNH) based CE assisted DSSC exhibited a good PCE.^{21–25}

The development of bifacial DSSM provides a potential candidate for building integrated photovoltaics (BIPVs).^{26–29} The bifacial BIPV structure enables light energy from direct sunlight (coming from outside) and diffused light (indoor light) to be converted into electric power. DSSCs are recognized as the best choice for indoor light conversion (~ 200–1000 k radiance).^{30–34} The DSSC is an ideal substituent for charging in tiny electronic gadgets under indoor light conditions. The DSSC combined with a battery or super-capacitor acts as a self-powered system, demonstrating its

Received: July 15, 2021 Accepted: September 20, 2021 Published: September 30, 2021





www.acsaem.org

Article

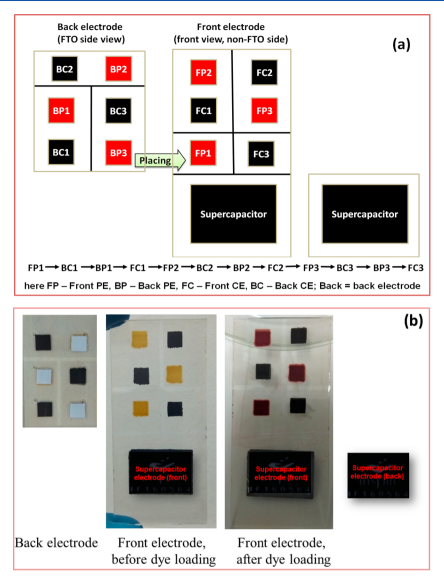


Figure 1. (a) Back and front DSSM glass after proper removal of the FTO layer (black line); and (b) resultant electrodes after material coatings.

advancement.³⁵ Many approaches are being attempted to improve the performance of self-powered systems.^{36,37}

Photocapacitors are emerging as promising, self-sufficient energy devices consisting of solar cells and supercapacitors. Over the past few years, researchers are focusing on the development of high-performance photocapacitors with higher energy density and good specific capacitance. The photocapacitor is capable of converting the electromagnetic radiation from the sun into electricity and storing the converted power. Most commonly, reported studies reveal a single solar cell charging a supercapacitor (SC). This limits the final voltage of the storage section such that it is not sufficient to drive most applications, like low-power electronics. To address this concern, here we proposed a device that consists of a bifacial DSSM (constituted series of six DSSCs) as a light harvester and an ionic liquid based supercapacitor as a storage unit. The two efficient SWCNH/carbon assisted identical electrodes are used for SC fabrication and are filled with an ionic EMIBF₄ electrolyte. Upon two-sided illumination of the bifacial DSSM, the delivered voltage is used for charging the supercapacitor.

Here, SWCNH supported carbon is used as the CE for the DSSC as well as for SC electrodes. The DSSM delivered a V_{oc}

of 3.94 V with an overall efficiency of 19.71%. The enhanced performance is reflected from its higher conductivity, excellent electrochemical properties, and the high current density of SWCNH/carbon CEs.

2. EXPERIMENTAL SECTION

Photoelectrode Preparation. Photoelectrodes were prepared by the procedure discussed here. First, FTO (fluorine doped tin oxide) (Greatcell Solar, TEC7, 7 Ω /square) conducting glass substrates were carefully washed with detergent liquid and eroded in sequence with deionized water, absolute ethanol, acetone, and finally 2-propanol for 10 min each to eliminate organic contaminants if any. A compact film of titanium dioxide solution (Solaronix, T-L/SC) was layered over a cleaned FTO glass substrate, which was then heated to 70 °C for 30 min. A 13–15 μ m TiO₂ film is realized by screen printing 18 NR-T transparent titanium dioxide paste (18-NR, Greatcell Solar) and sintering to 500 °C (30 min). The treatment of a compact layer like that described above is again repeated. Then, the substrates were cooled to 100 °C and immersed in a solution of N719 dye (0.3 mM in absolute ethanol) in dark conditions for 16 h. Finally, electrodes were washed gently with ethanol to remove any unanchored molecules of N719 dye.

Counter Electrode Preparation, Test Cell (DSSC) Assembly. The carbon based counter electrodes were prepared via screen print

by use of a commercial carbon paste (Solaronix) and sintered to 500 $^{\circ}$ C. A solution containing 5 mg of SWCNH dispersed in 2-propanol was spray coated over a carbon coated FTO glass substrate at 100 $^{\circ}$ C to get SWCNH/carbon counter electrodes. For comparison, platinum based CEs were prepared (spin coating of a Pt solution at 2000 rpm, sintered to 500 $^{\circ}$ C).

The closed type individual test cell (DSSC) was fabricated by a sandwich of photoelectrodes and counter electrodes, with thermal adhesive film (Solaronix, 25 μ m, polymer melt film) placed in between, pressed gently on a hot plate while heating at 110 °C. Altered HI-30 electrolyte was injected via the technique of back vacuum, and holes at CEs were closed through the polymer melt film and cover glass.

Dye Sensitized Solar Module (DSSM) Fabrication and Integration with Supercapacitor. Front Electrode Preparation. Glass substrates (FTO conducting) of $8.5 \times 4 \text{ cm}^2$ size were taken and etched properly using zinc dust and HCl to disconnect electrically from one to another part as shown in Figure 1a (black lines indicate the FTO removed part). Initially, glass substrates were cleaned in ethanol and acetone in an ultrasonicator for 10 min. A 5 μ m thick carbon film was deposited to act as CE (FC1, FC2, FC3, and supercapacitor part, one electrode) using a screen print technique and heated to 100 °C. Over FC1, FC2, and FC3, SWCNHs solution (in 2propanol) was spray coated and heated to 100 °C. As the other parts on the front electrode are treated, TiO₂ paste was then deposited (FP1, FP2, FP3) by using a screen print technique, along with sintering to 500 °C for 30 min to remove binders and solvents to electrically interconnect well with the respective nanoparticles. Then, the electrode was soaked for 16 h in 0.3 mM N719 dye in ethanol solution. While soaking, the carbon part (FC1, FC2, FC3 and supercapacitor part) is sealed with Kapton tape and Surlyn to avoid damage of carbon.

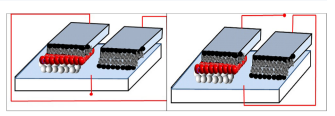
Back Electrode Preparation. FTO glass substrates of 5×3.5 cm² size were used, to have additional space for the electrode contact for measurements and etched properly to disconnect electrically from one to another part as shown in Figure 1a (black lines specify the FTO removed part). Six inlet holes of approximately 0.5 mm diameter were drilled (shown in Figure 1b) for electrolyte filling. FTO glass substrates were cleaned, and carbon films were deposited for CE parts (BC1, BC2, BC3) using screen printing and heated to 100 °C. Over BC1, BC2, and BC3, SWCNH solution was sprayed and heated to 100 °C. As the other parts on the back electrode, TiO₂ paste was then deposited (BP1, BP2, BP3) by using a screen printer and then sintered to 500 °C for 30 min. Before dye soaking, the carbon part (BC1, BC2, BC3) was sealed with Kapton tape and Surlyn to avoid damage of the carbon. Then, the electrode was soaked for 16 h in 0.3 mM N719 dye in ethanol solution, and the resulting electrode is shown in Figure 1b as after dye loading.

Assembling of DSSM. These front and back electrodes were sealed by placing a thermal adhesive film (Solaronix, polymer melt film) with a gentle hot press up to 110 °C. An acetonitrile (AcN) based redox coupled altered electrolyte (altered HI-30) was injected using a back vacuum filling technique via 6 holes followed by its perfect seal.

Integrated Device. Another FTO glass substrate of size 3.5×4 cm² was cleaned thoroughly, and the carbon paste was screen printed (deposition area 2×3 cm²) and heated to 100 °C. Then, the SWCNH solution was sprayed and sintered at 500 °C; this acted as a second electrode for the supercapacitor. A separator consisting of a Surlyn sheet was placed between the two supercapacitor electrodes, and a liquid electrolyte of EMIBF₄ was filled and gently sealed. Figure 2 presents the resultant photocapacitor (integrated the dye sensitized solar cell with the supercapacitor).

3. CHARACTERIZATION AND MEASUREMENTS

Microstructural studies of carbon, SWCNH, and SWCNH decorated carbon bilayer films and cross sections are examined by SEM (scanning electron microscopy, JEOL, JSM-IT500) and also by EDAX through APEX. X-ray diffraction studies



www.acsaem.org

Figure 2. Photo supercapacitor connections under illumination and in the dark.

were recorded by a Rigaku Miniflex diffractrometer in between 0° and 80° of 2θ value. Electrocatalytic activity was observed through cyclic voltammetry (CV) with an electrochemical workstation CompactSTAT.h IVIUM technology with threeelectrode configurations. Ag/Ag⁺ was used as the reference electrode, by use of acetonitrile solution containing 0.5 mM I_2 , 0.1 M LiClO₄, and 5 mM LiI, at a scan rate of 50 mV s⁻¹. Photocurrent density (I)-voltage (V) characterization of the developed test devices was examined with a solar simulator (PEC-L01, Peccell Inc.) containing a spectral filter (AM 1.5) and source meter (2401N Keithley). The light intensity of the solar simulator is adjusted to 100 mW/cm². The Tafel polarization plots and EIS (electrochemical impedance spectroscopy) were carried out for symmetrical dummy test cells (CompactSTAT.h workstation), with 10 mV of amplitude between frequencies of 1 Hz and 1 MHz.

4. RESULTS AND DISCUSSION

The microstructural studies of carbon, SWCNH, and SWCNH decorated carbon based electrodes are examined by SEM studies, and the resulting images are presented as Figure 3. The

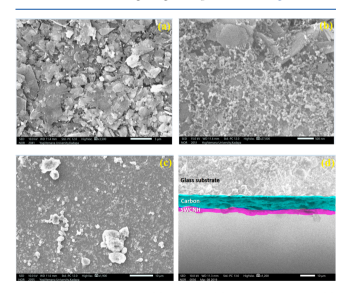


Figure 3. SEM images of (a) carbon, (b) SWCNH/carbon, (c) SWCNH, and (d) cross-sectional view of SWCNH/carbon.

pristine carbon had flakes of smooth and thin layer structures shown in Figure 3a. Figure 3b represents a SWCNH/carbon based electrode having an interconnected nature. Carbon flakes covering the SWCNH layer are observed for the SWCNH coated carbon electrode, shown in Figure 3b. Figure 3c presents the SWCNH coated electrode, consisting of agglomerated SWCNHs. Figure 3d represents the crosssectional view of SWCNH over carbon coated electrodes. The carbon film thickness of 12 μ m is present, over which a

Article

thin SWCNH layer is observed. The EDAX spectra are represented in Supporting Information Figure S1. The X-ray diffraction (XRD) studies are performed and presented as Figure S2 of Supporting Information.

The performance of the electrocatalytic activity of the developed counter electrodes is monitored in an I^-/I_3^- redox based solution by use of cyclic voltammetry (CV) studies having a three-electrode configuration. The platinum (Pt) wire acts as the counter electrode, and the Ag/Ag⁺ acts as the reference electrode in acetonitrile based solution consisting of 10 mM LiI, 0.1 M LiClO₄, and 1 mM I₂.

The prepared working electrodes (carbon, SWCNH, SWCNH/carbon, and Pt solution coated FTO glass substrates) are inserted into the solution, and ultrapure N_2 (nitrogen) gas is purged well before the start of the scans for better performance. The witnessed results are presented in Figure 4, in which the two pairs of prominent of peaks are

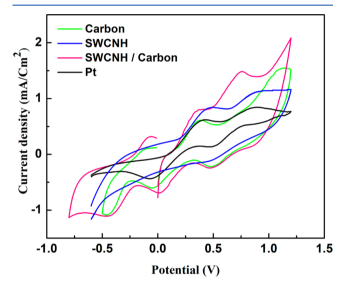


Figure 4. *C*-*V* curves of carbon, SWCNH, SWCNH/carbon, and Pt electrodes.

clearly observed. The two pairs of redox peaks, anodic peaks and cathodic peaks, are observed for carbon, SWCNH/carbon, Pt, and SWCNH. The cathodic peak reflects the reduction of tri-iodide, and anodic peaks reflect the oxidation of iodide and tri-iodide. SWCNH/carbon presented oxidation and reduction peaks similar to those for Pt. This specifies that the SWCNH/ carbon electrode displays improved catalytic activity compared with that of Pt. The redox reaction that occurs at the CE of DSSC in the cyclic voltammogram is owed to the reduction of iodine molecules. The SWCNH/carbon electrode presented both higher oxidation and reduction current density in a comparison with those of Pt, predicting a wild rate of triiodide reduction.³⁸ For the carbon working electrode, there is another pair of peaks found; this is associated with oxidation and reduction of carbon itself.³⁹

The Tafel slope is an intrinsic characteristic of electrocatalyst materials that is determined by the rate-limiting step.⁴⁰ Figure 5 represents Tafel polarization curves with logarithmic current density (log mA) as a voltage function for oxidation and reduction reactions of carbon, SWCNH/carbon, SWCNH, and Pt electrodes. Tafel polarization was accomplished with symmetric dummy test cells (two similar counter electrodes sandwiched, with filled electrolyte into the cell to become

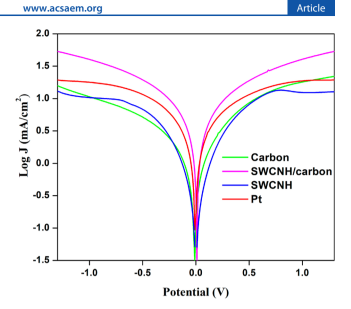


Figure 5. Tafel polarization curves of carbon, SWCNH, SWCNH/ carbon, and Pt CEs.

symmetric dummy test cells). From Figure 5, the SWCNH/ carbon electrode performed a larger exchange current density $(J_{\rm O})$ than the carbon, SWCNH, which implies a higher electrocatalytic activity and reduced charge-transfer resistance at the electrode–electrolyte interface. Tafel curve branches of anodic and cathodic nature for the SWCNH/carbon electrode presented a greater slope representing a higher exchange current density.

Figure 6 depicts photo current density-voltage (J-V) curves; its resultant photovoltaic parameter values with

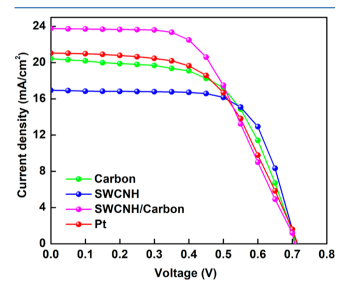


Figure 6. I-V curves of carbon, SWCNH/carbon, SWCNH, and Pt CE based DSSCs.

acetonitrile based electrolyte are given in Table 1. The photovoltaic performance with N719 dye based DSSC accumulated with prepared electrodes and derived performance of photovoltaic parameters, like short-circuit current density, efficiency, fill factor, open-circuit voltage, is tabulated in Table 1. DSSC with carbon CE shows $J_{sc} = 20.44 \text{ mA/cm}^2$; V_{oc} is 0.71 V, with a fill factor of 0.63, yielding a PCE of 9.14%. In contrast, the cell that ended with SWCNH/carbon

Table 1. Fabricated DSSC Photovoltaic Parameters

parameter	carbon	SWCNH	SWCNH/carbon	Pt	
$V_{\rm oc} (V)^a$	0.71	0.72	0.72	0.71	
$J_{\rm sc} ({\rm mA/cm^2})^a$	20.44	16.99	23.76	21.04	
fill factor (FF) ^a	0.63	0.63	0.65	0.63	
efficiency (η) $(\%)^a$	9.14	7.74	11.12	9.41	
${}^{a}V_{oc}$: ±30 mV. J_{sc} : ±0.20 mA/cm ² . FF: ±0.03. η : ±0.10.					

harvested $\eta = 11.12\%$ with $V_{oc} = 0.72$ V, $J_{sc} = 23.76$ mA/cm², and FF = 0.65. For comparison, SWCNH and Pt based test cells are fabricated, and their photovoltaic parameters are tabulated in Table 1.

The overall performance of the SWCNH/carbon based DSSC is high due to the conductivity of carbon and the high electrochemical activity of SWCNH; the composite DSSC exhibited a higher PCE. The stability of the materials and device performance are provided in the Supporting Information, Figure S3a,b.

EIS (electrochemical impedance spectroscopy) measurements were performed to examine the interfacial chargetransfer process that occurs at the electrode–electrolyte– electrode interfaces of symmetrical dummy test cells. The Nyquist plots consist of two semicircles as shown in Figure 7.

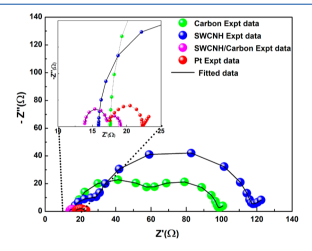


Figure 7. EIS of dummy test cells made with the same CEs.

At the higher-frequency region, the first semicircle intercept on the real axis is assigned to series resistance (R_S) of the FTO– SWCNH/carbon interface, whereas, at the lower-frequency range, the intercept on the real axis signifies $R_{\rm CT}$ (chargetransfer resistance). The R_S and $R_{\rm CT}$ values are evaluated through the fitted equivalent circuit (inset of Figure 7), tabulated in Table 2.

From this, we can conclude that SWCNH/carbon offers a better performance. The R_{CT} value depends on the counter electrode's catalytic materials; the fitted R_{CT} for the SWCNH/ carbon CE is effective compared to that of the Pt CE; the

Table 2. EIS Parameters and J_0 of Dummy Test Cells Fabricated with the Same CEs

parameter	SWCNH	carbon	SWCNH/carbon	Pt
Rs	15.92	17.60	13.81	17.41
$R_{\rm CT}$	21.01	37.90	19.40	19.68
$J_{\rm O}~({\rm mA/cm^2})$	1.50	3.77	10.43	8.75

higher value R_{CT} of SWCNH/carbon indicates a higher ability for the reduction of I_3^- to I^- in an electrolyte solution compared with that of the Pt CE.

www.acsaem.org

The Bode phase plots are presented as Figure 8; from the peak values, the electron lifetime (τ_e) is evaluated for carbon,

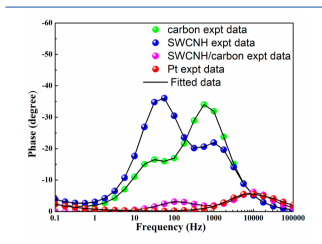


Figure 8. Symmetric dummy cell Bode phase plots.

SWCNH, SWCNH/carbon, and Pt based devices. Here, the frequency peaks confirm the charge transfer at different interfaces for these devices. The SWCNH/carbon test device's characteristic peak frequency is present at the lower-frequency side, demonstrating a longer electron lifetime of 1.3 ms, which is evaluated by using the equation $\tau_e = 1/2\pi f_{max}$.

Bifacial DSSM Performance. Dye sensitized solar cells are potentially useful as photovoltaic technology for indoor as well as outdoor applications. In particular, a bifacial DSSC is a promising candidate for oriented building integrated photovoltaics (BIPVs), which enables harvesting both direct sunlight and diffused light (indoor light) for power conversion. The present work demonstrated a W-type module which consists of six individual DSSC test cells in a connected series, named DSSM; the potential for power conversion of sunlight and indoor light is illustrated in Figure 9.

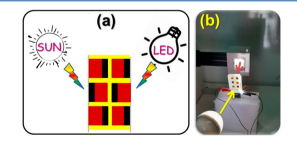


Figure 9. Developed W-type DSSM module taking advantage of both (a) sunlight and (b) indoor light conversion.

The developed bifacial DSSM was exposed to light power of 100 mW/cm² in addition to indoor light of 1000 lx; the performance is measured, and the obtained results of J-V characteristic performance are depicted in Figure 10. Photovoltaic parameters are summarized in Table 3. Here, the DSSM part exists in the W-type design in which photoelectrodes interchange between the front side and the back side of the submodule as represented in Figure 1. The DSSM revealed a

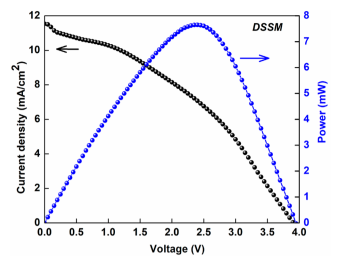


Figure 10. J-P-V plot of fabricated dye sensitized solar module.

Table 3. Photovoltaic Parameters of DSSM

DSSM
3.94
11.53
0.53
19.71

 $V_{\rm OC}$ of 3.94 V, $J_{\rm SC}$ of 11.53 mA/cm², and FF of 0.53, which leads to a PCE (power conversion efficiency) of 19.71%.

Figure 10 also presents the power-voltage characteristic of bifacial DSSM. Here, particular attention has to be paid to the shape of the curve, since that can provide precious information about the photocharging process. The power has a maximum value at 2.4 V. As reported, the overall efficiency increases quickly in the first part of the photocharge, since the storage of energy increases almost linearly in this region.

Supercapacitor Performance. The performance of the fabricated supercapacitor using SWCNH/carbon is made through cyclic voltammetry, EIS, and galvanostatic chargedischarge (GCD) measurements. The supercapacitor is loaded with efficient electrolyte. Cyclic voltammetry plots are recorded from 10 to 200 mV s⁻¹ scan rates over a voltage window from 0 to 1.5 V, and the results are shown in Figure 11a. The obtained curves are in a quasirectangular shape. From CV, the SWCNH/carbon based SC encircles a larger area, which indicates superior capacitance performance. The GCD plots are measured in the range of current densities from 0.5 to 1.5 mA/cm^2 over the voltage window from 0 to 1 V (Figure 11b), which is shown in a nearly triangular shape. Areal capacitance (ASC), power density (P), and energy density (E)in the two-electrode mode are evaluated by using the equations mentioned below, and the evaluated parameter values are tabulated in Table 4.

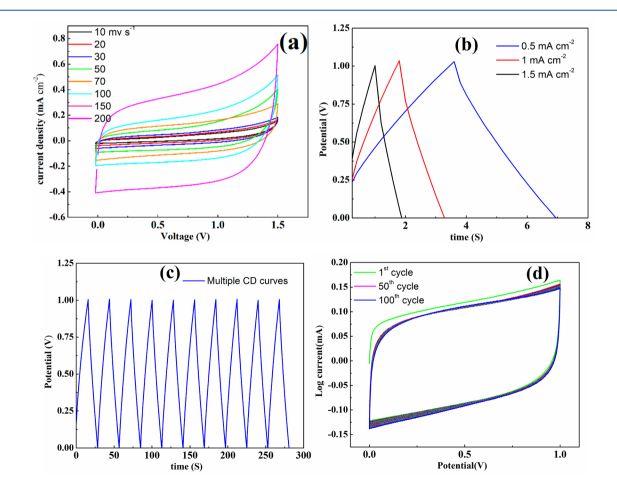


Figure 11. (a) CD curves of SWCNH/carbon SC at different charge–discharge current densities. (b) CV curves of the SC from 10 mV s⁻¹ to 200 mV s⁻¹ scan rates. (c) Galvanostatic charging and discharging curves versus time plot. (d) 100 cycles of CV curves at a scan rate of 100 mV s⁻¹.

$$P(W kg^{-1}) = E/\Delta t$$

 Table 4. Electrochemical Properties of Symmetric

 Supercapacitors

current density, I (mA/cm ²)	ASC (F g^{-1})	E (Wh kg ⁻¹)	$P (W kg^{-1})$
0.5	74.27	38.16	6.51
1.0	72.21	37.43	13.72
1.5	64.19	35.48	19.71

In the above equations, the current density is I in mA/cm^2 , the discharge time is Δt in s, and the voltage window ΔV is in V. From a decrease in current density from 1.5 to 0.5 mA/cm^2 , the discharge time is increased, which confirms that the charge storing capacity of the developed material is higher owing to the accessibility of a greater number of active sites which are electrochemically available and have a lower charge-transfer resistance. The SWCNH/carbon electrode galvanostatic charge-discharge performance up to 10 cycles for voltage starting from 0 to 1 V is demonstrated in Figure 11c, and the obtained curves have a triangular-like shape. The areal specific capacitance (ASC), energy density (E), and power density (P)are evaluated, with the values of 74.27 F g^{-1} , 38.16 Wh kg^{-1} , and 6.51 W kg⁻¹, respectively, by applying a current density of 0.5 mA/cm². The SWCNH/carbon electrode is higher in electrical conductivity, with greater electrocatalytic active surface sites which improve the overall storage of charge (Figure 11b). Figure 11d shows that the CV is constant during 100 cycles of measurement, evidencing stable performance.

An electrochemical impedance spectroscopy measurement for SC is also performed, and the results are depicted in Figure 12. At the higher-frequency side, the series resistance value found as R_S is 33.98 Ω for the SWCNH/carbon based supercapacitor in EMIBF₄ based acetonitrile electrolyte.

DSSM Working Demonstration through an Electronic LED. The DSSM delivered a higher power conversion

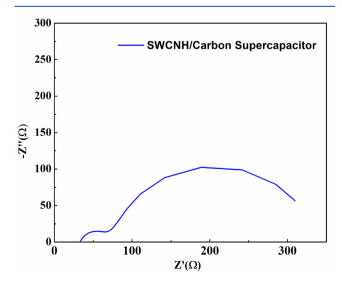


Figure 12. Electrochemical impedance spectroscopy of supercapacitor.

11231

www.acsaem.org

efficiency (η) of 19.71% and the energy density of the most of reported studies.^{20,41–43} By considering the advantage of this as well as bifacial illumination of the 100 mW/cm² light condition at one side, and the indoor light of 1000 lx on the other side, DSSM which demonstrated an open-circuit voltage of 3.94 V (experimental setup shown in Figure 13) is adequate

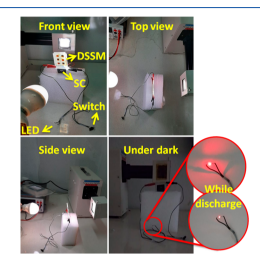


Figure 13. DSSM illumination from both sides, under dark condition discharge of SC lighting an LED.

to charge the supercapacitor. With the discharge of SC, it lit up a light emitting diode (LED) which is shown as an inset for Figure 13, demonstrating its potential as a power source. This is suggesting the better electrochemical features for the SWCNH/carbon device. Figure 13 evidenced lighting of the LED by the supercapacitor under dark conditions.

The overall photoconversion and storage efficiency (η_{overall}) of the photocapacitor (PC) device are calculated by use of the following equations.

$$\eta_{\text{overall}} = (E_{\text{PC}} \times A_{\text{SC}}) / (E_{\text{light}} \times t_{\text{ch}} \times A_{\text{DSSC}})$$

In the above equation, $E_{\rm PC}$ is the energy density of PC, $E_{\rm light}$ is the incident light power density (100 mW/cm²), $t_{\rm ch}$ is the photocharging time, $A_{\rm SC}$ is the effective active surface area of supercapacitor part, and $A_{\rm DSSC}$ is the effective active surface area of the solar cell part in PC.

The energy storage efficiency of the PC is given by the following equation.

$$\eta_{\rm storage} = \eta_{\rm overall} / \eta_{\rm conversion}$$

In the above equation, η_{storage} is the energy storage efficiency of the supercapacitor part in the PC, and $\eta_{\text{conversion}}$ is the PCE of solar cell in the PC (19.71%). Thus, η_{overall} and η_{storage} of the PC device are determined to be 10.46% and 53.3%, respectively.

5. CONCLUSIONS

The photovoltaic performance of a single wall carbon nanohorn (SWCNH) assisted carbon counter electrode (CE) based DSSC demonstrated a power conversion efficiency (PCE) of 11.12%, which is a motivation to develop a submodule (DSSM) which consists of six DSSCs connected in series. This setup acted as a bifacial module. This notable performance drove the fabrication of a SWCNH/carbon based photocapacitor, which exhibited excellent performance and

demonstrated an electronic light emitting device (LED) function upon discharge of the SC. The DSSM is capable of converting light illuminated from both sides into electric power, which showed a remarkable power conversion efficiency of 19.71% and an open-circuit voltage of 3.94 V; this reflects the potential for building integrated photovoltaics (BIPVs).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsaem.1c02087.

EDAX, XRD, and details about the stability of the materials and test devices (PDF)

AUTHOR INFORMATION

Corresponding Author

Raghavender Mitty – Department of Physics, Yogi Vemana University, Kadapa 516005, India; o orcid.org/0000-0002-0958-4378; Email: toraghavender@rediffmail.com

Authors

- **Gurulakshmi Maddala** Department of Physics, Yogi Vemana University, Kadapa 516005, India
- Meenakshamma Ambapuram Department of Physics, Yogi Vemana University, Kadapa 516005, India
- VijayaLaxmi Tankasala Department of Mathematics, N.T.R. Government Degree College for Women, Mahabubnagar 509001, India

Complete contact information is available at: https://pubs.acs.org/10.1021/acsaem.1c02087

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

G.M. is thankful to the Department of Science and Technology (DST)-INSPIRE for financial support through IF 160564. R.M. thanks DST-SERB (EMR/CRG) EMR/2016/007049 for the utilized equipment purchased through this financial support. A.M. is thankul to DST for WOS-B through DST/WOS-B/ER-7/2021.

REFERENCES

(1) Hagfeldt, A.; Boschloo, G.; Sun, L.; Kloo, L.; Pettersson, H. Dye Sensitized Solar Cells. *Chem. Rev.* **2010**, *110*, 6595–6663.

(2) Li, L. L.; Diau, E. W.G. Porphyrin-Sensitized Solar Cells. Chem. Soc. Rev. 2013, 42, 291–304.

(3) Yella, A.; Lee, H. W.; Tsao, H. N.; Yi, C. Y.; Chandiran, A. K.; Nazeeruddin, M. K.; Diau, E. W. G.; Yeh, C. Y.; Zakeeruddin, S. M.; Gratzel, M. Porphyrin-Sensitized Solar Cells with Cobalt (II/III) Based Redox Electrolyte Exceed 12% Efficiency. *Science* **2011**, *334*, 629–634.

(4) Ito, S.; Murakami, T. N.; Comte, P.; Liska, P.; Grätzel, C.; Nazeeruddin, M. K.; Grätzel, M. Fabrication of thin film dye sensitized solar cells with solar to electric power conversion efficiency over 10%. *Thin Solid Films* **2008**, *516*, 4613–4619.

(5) Gao, F.; Wang, Y.; Shi, D.; Zhang, J.; Wang, M.; Jing, X.; Humphry-Baker, R.; Wang, P.; Zakeeruddin, S. M.; Grätzel, M. Enhance the Optical Absorptivity of Nanocrystalline TiO2 Film with High Molar Extinction Coefficient Ruthenium Sensitizers for High Performance Dye-Sensitized Solar Cells. J. Am. Chem. Soc. **2008**, 130, 10720–10728. (6) Du, Y. F.; Fan, J. Q.; Zhou, W. H.; Zhou, Z. J.; Jiao, J.; Wu, S. X. One-Step Synthesis of Stoichiometric $Cu_2ZnSnSe_4$ as Counter Electrode for Dye-Sensitized Solar Cells. *ACS Appl. Mater. Interfaces* **2012**, *4*, 1796–1802.

(7) Gong, F.; Wang, H.; Xu, X.; Zhou, G.; Wang, Z. S. In Situ Growth of $Co_{0.85}$ Se and $Ni_{0.85}$ Se on Conductive Substrates as high-performance Counter Electrodes for Dye-Sensitized Solar Cells. *J. Am. Chem. Soc.* **2012**, *134*, 10953–10958.

(8) Wang, M.; Anghel, A. M.; Marsan, B. t.; Cevey Ha, N. L.; Pootrakulchote, N.; Zakeeruddin, S. M.; Grätzel, M. CoS Supersedes Pt as Efficient Electrocatalyst for Triiodide Reduction in Dye Sensitized Solar Cells. J. Am. Chem. Soc. **2009**, 131, 15976–15977.

(9) Wu, M.; Lin, X.; Wang, Y.; Wang, L.; Guo, W.; Qi, D.; Peng, X.; Hagfeldt, A.; Grätzel, M.; Ma, T. Economical Pt-Free Catalysts for Counter Electrodes of Dye-Sensitized Solar Cells. *J. Am. Chem. Soc.* **2012**, *134*, 3419–3428.

(10) Roy-Mayhew, J. D.; Bozym, D. J.; Punckt, C.; Aksay, I. A. Functionalized Graphene as a Catalytic Counter Electrode in Dye Sensitized Solar Cells. *ACS Nano* **2010**, *4*, 6203–6211.

(11) Li, G. R.; Song, J.; Pan, G. L.; Gao, X. P. Highly Pt-like Electrocatalytic Activity of Transition Metal Nitrides for Dye Sensitized Solar Cells. *Energy Environ. Sci.* **2011**, *4*, 1680–1683.

(12) Wu, M. X.; Lin, X. A.; Hagfeldt, A.; Ma, T. L. Low-Cost Molybdenum Carbide and Tungsten Carbide Counter Electrodes for Dye-Sensitized Solar Cells. *Angew. Chem., Int. Ed.* **2011**, *50*, 3520– 3524.

(13) Jiang, Q. W.; Li, G. R.; Gao, X. P. Highly ordered TiN nanotube arrays as counter electrodes for dye-sensitized solar cells. *Chem. Commun.* **2009**, *44*, 6720–6722.

(14) Imoto, K.; Takahashi, K.; Yamaguchi, T.; Komura, T.; Nakamura, J.; Murata, K. High-Performance Carbon Counter Electrode for Dye-Sensitized Solar Cells. *Sol. Energy Mater. Sol. Cells* **2003**, *79*, 459–469.

(15) Murakami, T. N.; Ito, S.; Wang, Q.; Nazeeruddin, M. K.; Bessho, T.; Cesar, I.; Liska, P.; Humphry-Baker, R.; Comte, P.; Pechy, P.; Grätzel, M. Highly Efficient Dye-Sensitized Solar Cells Based on Carbon Black Counter Electrodes. *J. Electrochem. Soc.* **2006**, *153*, A2255–A2261.

(16) Ramasamy, E.; Lee, W. J.; Lee, D. Y.; Song, J. S. Nanocarbon Counter electrode for Dye Sensitized Solar Cells. *Appl. Phys. Lett.* **2007**, *90*, 173103.

(17) Huang, Z.; Liu, X. H.; Li, K. X.; Li, D. M.; Luo, Y. H.; Li, H.; Song, W. B.; Chen, L. Q.; Meng, Q. B. Application of Carbon Materials as Counter Electrodes of Dye-Sensitized Solar Cells. *Electrochem. Commun.* **2007**, *9*, 596–598.

(18) Suzuki, K.; Yamaguchi, M.; Kumagai, M.; Yanagida, S. Application of Carbon Nanotubes to Counter Electrodes of Dye-Sensitized Solar Cells. *Chem. Lett.* **2003**, *32*, 28–29.

(19) Hino, T.; Ogawa, Y.; Kuramoto, N. Preparation of Functionalized and Non-functionalized Fullerene Thin Films on ITO Glasses and the Application to a Counter Electrode in a Dye-Sensitized Solar Cell. *Carbon* **2006**, *44*, 880–887.

(20) Das, A.; Ojha, M.; Subramanyam, P.; Deepa, M. A poly (3,4propylene dioxy thiophene)/carbon micro-sphere-bismuth nanoflake composite and multifunctional Co-doped graphene for a benchmark photo-supercapacitor. *Nanoscale Adv.* **2020**, *2*, 2925–2942.

(21) Ebbesen, T. W.; Lezec, H. J.; Hiura, H.; Bennett, J. W.; Ghaemi, H. F.; Thio, T. Electrical Conductivity of Individual Carbon Nanotubes. *Nature* **1996**, *382*, 54–56.

(22) Chew, S. Y.; Ng, S. H.; Wang, J. Z.; Novak, P.; Krumeich, F.; Chou, S. L.; Chen, J.; Liu, H. K. Flexible Free-Standing Carbon Nanotube Films for Model Lithium-Ion Batteries. *Carbon* **2009**, *47*, 2976–2983.

(23) Chen, Y. S.; Huang, J. H.; Chuang, C. C. Glucose Biosensor Based on Multiwalled Carbon Nanotubes Grown Directly on Si. *Carbon* 2009, 47, 3106–3112.

(24) Gurulakshmi, M.; Meenakshamma, A.; Susmitha, K.; Charanadhar, N.; Srikanth, V. V. S. S.; Narendra Babu, S.; Venkata Subbaiah, Y. P.; Venkateswarlu, Katta; Raghavender, M. A transparent

and Pt-free all-carbon nanocomposite counter electrode catalyst for efficient dye sensitized solar cells. *Sol. Energy* **2019**, *193*, 568–575.

(25) Gurulakshmi, M.; Meenakshamma, A.; Susmitha, K.; Venkata Subbaiah, Y.; Raghavender, M. Enhanced Performance of Dye-Sensitized Solar Cells (DSSCs) Based on MoS₂/Single-Walled Carbon Nanohorns Electrochemically Deposited Bilayer Counter Electrodes. *ChemPlusChem* **2020**, *85*, 2599–2605.

(26) Lodermeyer, F.; Costa, R. D.; Guldi, D. M. Review—Single-Walled Carbon Nanohorn-Based Dye-Sensitized Solar Cells. ECS J. Solid State Sci. Technol. 2017, 6, M3140–M3147.

(27) Huaulmé, Q.; Mwalukuku, V. M.; Joly, D.; Liotier, J.; Kervella, Y.; Maldivi, P.; Narbey, S.; Oswald, F.; Riquelme, A. J.; Anta, J. A.; Demadrille, R. Photochromic Dye-Sensitized Solar Cells with Light-Driven Adjustable Optical Transmission and Power Conversion Efficiency. *Nat. Energy* **2020**, *5*, 468–477.

(28) Zhang, K.; Qin, C.; Yang, X.; Islam, A.; Zhang, S.; Chen, H.; Han, L. High-Performance, Transparent, Dye-Sensitized Solar Cells for See-through Photovoltaic Windows. *Adv. Energy Mater.* **2014**, *4*, 1301966.

(29) Gokul, G.; Pradhan, S. C.; Soman, S. Dye-Sensitized Solar Cells as Potential Candidate for Indoor/Diffused Light Harvesting Applications: From BIPV to Self-Powered IoTs. *Advances in Solar Energy Research*; Springer: Singapore, 2019; pp 281–316. DOI: 10.1007/978-981-13-3302-6 9

(30) Fakharuddin, A.; Jose, R.; Brown, T. M.; Fabregat-Santiago, F.; Bisquert, J. A Perspective on the Production of Dye-Sensitized Solar Modules. *Energy Environ. Sci.* **2014**, *7*, 3952–3981.

(31) Michaels, H.; Rinderle, M.; Freitag, R.; Benesperi, I.; Edvinsson, T.; Socher, R.; Gagliardi, A.; Freitag, M. Dye-Sensitized Solar Cells under Ambient Light Powering Machine Learning: Towards Autonomous Smart Sensors for Internet of Things. *Chem. Sci.* 2020, *11*, 2895–2906.

(32) Zhang, W.; Wu, Y.; Bahng, H. W.; Cao, Y.; Yi, C.; Saygili, Y.; Luo, J.; Liu, Y.; Kavan, L.; Moser, J.-E.; Hagfeldt, A.; Tian, H.; Zakeeruddin, S. M.; Zhu, W.-H.; Grätzel, M. Comprehensive Control of Voltage Loss Enables 11.7% Efficient Solid-State Dye-Sensitized Solar Cells. *Energy Environ. Sci.* **2018**, *11*, 1779–1787.

(33) Cao, Y.; Liu, Y.; Zakeeruddin, S. M.; Hagfeldt, A.; Grätzel, M. Direct Contact of Selective Charge Extraction Layers Enables High-Efficiency Molecular Photovoltaics. *Joule* **2018**, *2*, 1108–1117.

(34) Freitag, M.; Teuscher, J.; Saygili, Y.; Zhang, X.; Giordano, F.; Liska, P.; Hua, J.; Zakeeruddin, S. M.; Moser, J. E.; Grätzel, M.; Hagfeldt, A. Dye-Sensitized Solar Cells for Efficient Power Generation under Ambient Lighting. *Nat. Photonics* **2017**, *11*, 372–378.

(35) Vega Garita, V.; Ramirez Elizondo, L.; Narayan, N.; Bauer, P. Integrating a photovoltaic storage system in one device: A critical review. *Prog. Photovoltaics* **2018**, *27*, 346 DOI: 10.1002/pip.3093.

(36) Ramadoss, A.; Saravanakumar, B.; Kim, S.-J. Fabrication and Characterization of Supercapacitors toward Self-Powered System; IntechOpen: London, 2018; pp 167–192. DOI: 10.5772/inte-chopen.73647

(37) Yuan, L.; Xiao, X.; Ding, T.; Zhong, J.; Zhang, X.; Shen, Y.; Hu, B.; Huang, Y.; Zhou, J.; Wang, Z. L. Paper-Based Supercapacitors for Self-Powered Nanosystems. *Angew. Chem., Int. Ed.* **2012**, *51*, 4934–4938.

(38) Hong, C. K.; Ko, H. S.; Han, E. M.; Park, K. H. Electrochemical Properties of Electrodeposited PEDOT Counter Electrode for Dyesensitized Solar Cells. *Int. J. Electrochem. Sci.* **2015**, *10*, 5521–5529.

(39) Costa, C.; Mesquita, I.; Andrade, L.; Mendes, A. Photoelectrochromic devices: Influence of device architecture and electrolyte composition. *Electrochim. Acta* **2016**, *219*, 99–106.

(40) He, J.; Lee, L. T. L.; Yang, S.; Li, Q.; Xiao, X.; Chen, T. Printable Highly Catalytic Pt- and TCO-Free Counter Electrode for Dye-Sensitized Solar Cells. ACS Appl. Mater. Interfaces **2014**, *6*, 2224–2229.

(41) Scalia, A.; Varzi, A.; Lamberti, A.; Tresso, E.; Jeong, S.; Jacob, T.; Passerini, S. High energy and high voltage integrated photoelectrochemical double layer capacitor. *Sustainable Energy Fuels* **2018**, *2*, 968–977. (42) Gao, K.; Ti, D.; Zhang, Z. A photocapacitor with high working voltage and energy density. *Sustainable Energy Fuels* **2019**, *3*, 1937–1942.

(43) Sasidharan, S.; Pradhan, S. C.; Jagadeesh, A.; Nair, B. N.; Mohamed, A. A. P.; K. N, N. U.; Soman, S.; Hareesh, U. N. S. Bifacial Dye-Sensitized Solar Cells with Enhanced Light Scattering and Improved Power Conversion Efficiency under Full Sun and Indoor Light Conditions. ACS Appl. Energy Mater. **2020**, *3*, 12584–12595.



، آھينہ بيگم ، ريسر چ اسکالر ، محکمہ تاريخ ، مولانا آز اد قومی اردو يونيورسٹی حيدرآباد۔ amenabegum.s777@gmail.com

سلطت، معقيد ي امير باتيكا ومرآ مان جادى رعابا يرورى

تويينيكم (ريمرى، كار شرية، دريغ) مواديا ترزيديحش اردو برشورش، كمكي بولى، حيد آباد

سلامی آصلید بینوبی بند کی آیی منظر و متحکم سلامی گزاری ہے۔ یہ سلطنت مناصرف اعدو متلان بلک عالمی تنظر و مطام رکھتی ہے۔ جسر قمر الدین خاص نے 1724 میں اسلامیہ آصلیہ کی بلیادة الی۔ جس میں 7 فرمانرواں کزرے ہیں۔ اور تک آیادا لکا پہلا دارا لکا فسلالہ کیکن بعد میں تکا مابلی خاص نے میدر آباد کوانا دارا لکا قد اطلہ۔

اللالدية المنايدين امرائية يا يكاوما مقام ومرتية تقران وقت كه يعدّ كالله ابية الب سلامة واكرت شطر الدائق خال من بايتكان المارية تأخر دوتي - الملامة كالمحلك، المارشون، المتادقات كي سركوني اورونيوان اورمصب دارون بر تشرول كريمان كالم م المسترق الجروع - البابية علاق تل خودتكار ش-

لواب سرآ مان جاد کانام محد طلوالدین خان قال آب آب سلطان الدین خان کے مجبوبے بینے تھے۔ آب کی شادی سلطن آ منیہ ک یو کچ پی تقدران المطل الدواری دفتر اور مرجوب ملی خان کی بعثیرہ ہی درش اشدار تک ماں کے مجبوبے بینے تھے۔ آب کی شادی سلطن آ منیہ ک حاصل قتار سلطن آسف میں حدد المباقی عدالت کے علیل القدر حبد ہے جافز ہوئے۔ انہوں نے باا محاصل اپنے خدمات انجام وب انہوں نے اپنے دور مدارالمباقی میں دعایا کی خان و ہندہ آتی، ان کے بلیادی طرور توں کی سلک کو این الدی تعالی میں ت حسین کارنا سے اس طرح ال

محصیت بافتلان کی باد ماموری تک نصف محفوا و جاری کرتے ہوئ ان کی بان موری کو یکی منایا۔ اس سے کلی طاعدانوں کی کفالت جو تی۔ طریب میں قد خواہ شند تیات کے لیے ملت نظ کی سمبات فراہم کیے۔ 1893 و شن ڈالد باری سے تیاد شدہ مکانت کی دوبارہ تعیر، باز آباد کاری سے لیے سرکاری فراند کے ساتھ ساتھ اپنے واتی سرمایی سے اواد طافر مانے۔ اوار یہ بیٹی و لیزا طفال کو سنست کی منت تعلیم ا ریائش کا انتظام کیے۔ تاکہ مستقبل میں بھرمندی کے ساتھ خوال وہ کی گزار ہیں۔ ای طرح کا دارے بھی دین را اللہ کی کر شوا تین مال تذہ کی گھرانی میں مان کی تعلیم وزیدے کہ ماتھ دستان وہ کی گزار ہے۔ ای طرح کا دارت بیٹیم و مندو کی کو کی کا ان ک

امیر پائیکا اس سال نے رعایا پر دری کے برگون میں تمایاں خدمات انجام دیے ہیں۔ سلطت کا کوئی گوندان کے خدمات سے علی تین ہے۔ ان کے شدمات ملک کے شہریوں کے ساتھ احمان تعلیم کے ساتھ بنی توج انسان کی دعد دی کا شوت دیا۔ ان کی خرخوانی ، دریادی، دادود بھی میں ان کا کوئی عاتی تیس۔ argania / Enrol No. 03HUPHOS



प्रमाणित किया जाता है कि

मुहम्मद असलेम फरूखी

को

मानविकी

संकाय में

शोध-पबंध पस्तुत करने एवं सभी अपेक्षाओं को सफलतापूर्वक पूर्ण करने पर सन् 2006 में डॉक्टर ऑफ़ फ़िलॉसफ़ी

की उपाधि प्रदान की गयी है



This is to certify that

MOHD ASLEM FAROOQUI

having submitted a thesis in the

School of Humanities

and having successfully completed all the requirements

has been admitted to the degree of

Doctor of Philosophy

in the year 2006

Sendenner all yet al ander see Given under the seal of the University

Vice-Chancellor

berter / Hyderabad, ftris / Dated gart/July 31, 1007.

त सचिव Registrar

which the second

Telestione (40.27682206



Dean, Faculty of Oriental Languages

OSMANIA UNIVERSITY HYDERABAD - 500 DOT TA P

Proceedings of the Dean, Faculty of Oriental Languages, Osmania University Hyderabad-

No.

Sub. Dr. Mahmood Ali: PhD supervisor in undu Oriental Ouris shifted to abroad allote Dates 25 - 09 2013 other PhD supervisors for certain Research scholars Regarding Ref. : Letter of Dr. Mahmood Ali: dated. 26-08 2013

As per request of Dr. Mahmood Ali Reader in Urdu (Retd.) who is one of the recognised PhD supervisor in Urdu Oriental is shifted to abroad, and six Research scholars are registered under his supervision. Hence these Research scholars are allotted to the other recognised PhD supervisors in Urdu oriental. Chair person BoS in Urdu Oriental OU i recommended the following PhD supervisors along with the Research topics.

SNo Name of the R. Scholar	Research topic	PhD supervisor
01 Syed Asrat ul Hag Ph 9346651710	Urdu zaban mein bachoun kea dab ke Aqiqi Asrath-Ek Tajola	Dr. A. M. Sved Faziuliah Chair person Bosin Urdu(O+OU Hvd PH 3849377948
02 Amena akhtar Ph. 8341883090	Warangai ke shuara-o udba ki adabi khidmath	Dr. Md Aslam Faboqui Lecturer, Gost Griraj Conege, Nicamabad PH 9247191548
03 Ali Bazahar Ph: 9247341586	Talamida e Zore aur unki adabi khidmath	Dr Mohd Shujat Ak Asset Prof. CPDUMT MANUU Hyd PH 9346258762
04 Fatima Akhtar Ph: 8341034559	Prof Md. Anwaruddin ki adabi khidmath	Dr. Md. Aslam Faonqui Lecturer: Govt Ginraj College: Nizamabad Ph 9247191548
05 Shaik Fareed Ph 9848531966	Prof Mir Turab Ali ki ilmi oʻadabi khidmath	Dr. Shaik Saleem, HoD Urdu, Anwarul Goom Degree College, Hyd Ph. 9849265753
06 Kahekashan Jabeen Ph-9052467750	Telengana ke chund namwer Afsana nigar Azadi ke baad	Dr. Md. Abrar ul Baqui HOD Urdu: Satavahana University, Karimnagar Ph. 9440717525
		Ph 9440717525



Copy to

1. Concerned PhD supervision

2. Concerned Research scholars

3. Deputy Registrar (Academic) Oc.

4 Addr - Controller of Examp (Confidential) Gill



CONFIDENTIAL SECTION EXAMINATION BRANCH NO. 647 Ph.D Exams/2022

OSMANIA UNIVERSITY HYDERABAD-500 007, LS. Dated: 01 Nov. 2022

6

PRESS NOTE

The following candidates who had presented the Thesis on the subject mentioned against each for the degree of Ph D are declared qualified for the award of Degree of Doctor of Philosophy (Ph D) of Osmania University, Hyderabad

Ph.D.

5.5	Reference No.	Name of the Candidate Father Name	Subject	Thesis Title	Supervisor: Regn. Date
1	PHI343409	Mr. D.Ravinder Naik. S.o. Bhadiya	Рћумса	Structural, Electric and Magnetic Properties Studies of Li-Mg and Li-Cr Nano Ferrites	Prof. D Ravinder Dept. of Physics, O L Hyd (02:03:2011)
2	P111343410	My, N Swathi D is N Familia	Pharmaceutica Sciences	Formulation Development and Evaluation of Mouth Dissolving Films of BCS Class II and III Drugs	Prof. D.Jaya Prakash Univ. College of Technology, O.U., Hyi- Otif, P.Amareshwar, Univ. College of Technology, O.U., Hyi- (13/04/2011)
3	PHD43411	Mr. S Bharath Goud S.o. S Pullarah Goud	Geography	Geographical Perspective of Foundation Education - An Analysis of Infrastructure and Perceptions in Mahabubriagar District	Dr. A Geetha Reddy Assoc: Professor, Centi for Indian Ocean Studies, O 11, Hvd (08.12/2006)
4	PHD43412	Ms. Sadia Saleem D.o. Abdul Malik Saleem	l'eduit menta l	ldara Aihanat Rampur Ki Elmi Wa Adabi Khisdinat	Dr. Mohammed Aslam Faroqui Asit Professor, Dept. c. Urdu, NTR Govt College (W), Mabboobnagar (14/09/2018)
475	PHD43413	Mr. Seetharamul Basani S.o. Gurusaiah	Telage	Rataka Kula Vignanam - Samajika, Samskruthika Parisheelana	Prof. Madireddy Andamma (Retd.) Dept. of Telugu, O.U., Hyd. (26:97/2010)
-fr	P111244414	Mr. G Krishna Reddy S.o. Raji Reddy	Library & Info science	Library Security Systems in the Select Libraries of Telangana: A Study	Prof. J. Nivekavardhan Librarian, Univ. Library Central University of Kerala, Kerala. (29.03/2011)
٦		Mr. Altaf Hussain Khan S.o. Ashfaq Hussain Khan		Lung (RA)	Dr. Md. Ashfaque Chand Dept. of Persian, O.U., Hyd (11/02/2013)
8	P1411441A	Mr. Poondia Radhakrishna S.o. P.Rama Subba Rao	Chemical Engineering	Neural Networks based NARMA-L2 Control of Processes with Input Maltiplicities	Prof. G Prabhaker Reddy Univ College of Technology, O.U., Hyd (17/01/2009)

(p. Vardhamiti

AddLController of Examinations

(Confidential)

Faculty of Oriental Languages - OU PhD Admissions in Urdu for the year 2014-15, 2015-16 & 2016-17

5.No	Name of the candidate	Type of	Research Topic	Name of the
1	FAREEDA BEGUM	admn.		supervisor
2	D/O WAHED KHAN	Full time	Urdu khaka nigari ke farough mein khawateen ka hissa	Dr. Rizwana Begum , Asst. Prof College of Languages Hyd.
20	HAFIZ MOHAMMED ASIF UDDIN S/O MOHD RAFEEQ AHMED	Full time	Hyderabad mein Iqbal Shanasi ka tahqeeqi wa tanqeedi jaaieza	Dr.S.A. Kareem Asst.Prof, College of Languages,Hyd
3	FAHMEEDA BEGUM D/O MOHD RAFEEQ	Full time	Dakhniyat se mutaliq kitaboun ka wazahati Isharia-1980 ke baad	Dr.Rizwana Begum , Asst.Prof College of Languages Hyd.
4	MD. MEHBOOB 5/O MD. GAFOOR	Full time	Muthahida Andhra Pradesh Urdu Academy ki Elmi wa Adabi khidmaat	Dr.Md.Aslam Faroqui, Asst.Prof, Govt. NTR College, Mahboobnagar
5/	SADIA SALEEM D/O ABDUL MALIK SALEEM	Full time	Idara e Alhasnat Rampur ki Elmi wa Adabi Khidmaath	Dr.Md.Aslam Faroqui, Asst.Prof, Govt. NTR College, Mahboobnagar
6	MOHAMMED MUJAHID ALI S/O MOHAMMED AMJAD ALI	Part time	Urdu theatre ke farough mein Hyderabad ka hissa	Dr.Md.Aslam Faroqui, Asst.Prof, Govt. NTR College, Mahboobnagar
7	MOHD SHAFIUDDIN S/O MOHD AZIZUDDIN	Part time	Jameel Shaidai ki Elmi wa Adabi khidmaath	Dr. Md. Shujat Ali MANUU , Hyderabad
8	MOHAMMED KHAJA MOHIUDDIN S/O MOHAMMED KAREEMUDDIN	Full time	Maulana Khaja Sharif ki Elmi wa Adabi khidmaath	Dr.S.A. Kareem Asst.Prof, College of Languages,Hyd
,	SYED LATEEF S/O SYED HABEEB	Full time	Hyderabad mein mizahia column nigari ka aaghaz wo irtegha	Dr. Md. Shujat Ali MANUU , Hyderabad.
0	MD. ASADULLAH KHAN S/O MOHAMMED HABEEBULLAH	Full time	Mohammed Habeeb ullah wafa : Hayat aur adabi khidmaath	Dr.S.A. Kareem Asst.Prof, College of Languages,Hyd
1	MOHD SHOUKAT FAHEEM AHMED S/O MD. JAFFAR	Full time	Hyderabad mein Ghalib shanasi ka tahqeeqi wa tanqeedi jaaleza	Dr. Md. Shujat Al MANUU , Hyderabad.
IZ	MAQBOOL HUSSAIN S/O AHMED HUSSAIN	Part time	Hyderabad ki khatoon shoura – Azadi ke baad	Dr.Rizwana Begum, Asst.Pro College of Languages Hyd.

DEAN

Faculty of Oriental Languages Dsmanie University Hyderabad