




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Pyrene-based covalent organic polymers with nano carbonaceous composites for efficient supercapacitive energy storage†

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Covalent organic polymers (COPs) show great potential as supercapacitor electrodes due to their tunable properties, stability, and high capacitance. In this work, we prepared a covalent organic polymer (Py-DSDA-COP) through a Schiff base reaction of the 1,3,6,8-tetrakis(4-formylphenyl)pyrene monomer (Py-Ph-CHO) and 4,4'-disulfanediyldianiline (DSDA), validated its chemical structure, and investigated its thermal stability, porosity, morphology, and electrochemical properties. The Py-DSDA-COP displayed high semi-crystallinity, good porosity, and a specific capacitance of 56.18 F g⁻¹. To improve its electrochemical performance, we incorporated different nano-carbonaceous materials [fullerene (C60), MWCNTs, and SWCNTs] into Py-DSDA-COP and compared their characteristics. The Py-DSDA-COP/SWCNTs showed the highest specific capacitance (171 F g⁻¹) at 1 A g⁻¹ and highest energy density (23.7 W h kg⁻¹) with a capacity retention of 93% after 2000 cycles, which is superior to those of Py-DSDA-COP/MWCNTs and Py-DSDA-COP/C60 nanocomposites. These superior electrochemical characteristics of Py-DSDA-COP/SWCNTs are attributed to the excellent π - π stacking interactions, higher EDLC behavior, higher conductivity, lower diameter, and higher surface area of SWCNTs than those of MWCNTs.

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Introduction

Covalent organic polymers (COPs) are a class of organic materials composed of repeating units linked by covalent bonds, resulting in a three-dimensional network structure.¹⁻⁶ They are typically made up of light elements such as carbon, nitrogen, and oxygen, and can have diverse chemical structures, allowing for a wide range of properties and potential applications.⁷⁻¹² In recent years, there has been a growing amount of attention towards COPs because of their unique properties, including high surface area, tunable porosity, and chemical stability.¹³⁻²⁰ COPs can also be synthesized in a variety of ways, including through the polymerization of small molecules or through self-assembly of pre-formed building blocks.²¹⁻²⁵ Some potential applications of COPs include gas storage materials for hydrogen and carbon dioxide,²⁶⁻²⁸ heterogeneous catalysts for organic

reactions,²⁹⁻³¹ and electrode materials for supercapacitors and battery devices.³²⁻³⁵ Supercapacitors (SCs), in particular, have emerged as potential electrochemical energy storage systems and have gained popularity due to their outstanding high-power density, quick charge/discharge processes, exceptional stability, and environmental friendliness.³⁶⁻⁴⁵ They have the capability to perform like batteries and traditional capacitors. Electrostatic interactions between the electrolyte ions and the carbon electrodes lead to storing energy in them. One of the key advantages of COPs for supercapacitor applications is their high surface area, which can be tuned by adjusting the size and shape of the monomers used to synthesize the polymer.⁴⁶⁻⁴⁸ The significant surface area offered by these materials provides numerous active sites for electrochemical reactions to occur which can enhance the energy storage capacity of the supercapacitor.^{49,50} Additionally, COPs can be easily synthesized using simple and cost-effective methods, which makes them attractive for large-scale production. However, one of the current limitations of COPs for supercapacitor applications is their relatively low electrical conductivity, which can limit their charge/discharge rate.⁵¹⁻⁵⁴ Incorporating conducting carbon materials (such as CNTs and graphene) into COFs, CMPs, and MOFs is an effective way to boost their conductivity, improving their electrochemical performance.⁵⁵⁻⁵⁸ The Py-OXD-CMP/CNT nanocomposites prepared by our group have exhibited a capacitance of 504 F g⁻¹.⁵⁵ In addition, our group has developed the TBN-Py-CMP/

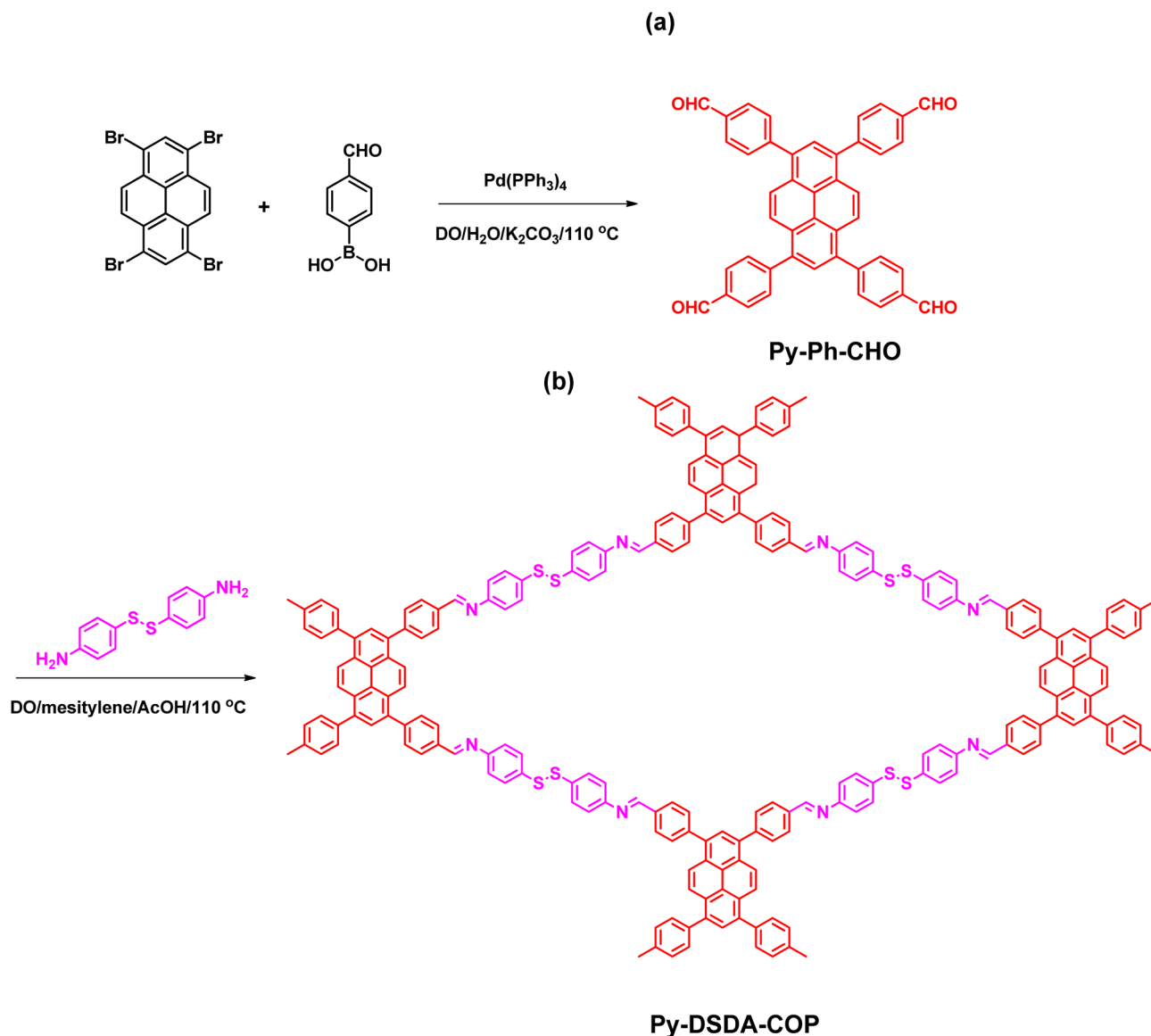
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Scheme 1 Synthesis of (a) Py-Ph-CHO and (b) Py-DSDA-COP.

CNT nanocomposite, which has exhibited a specific capacitance of 430 F g^{-1} .⁵⁶

The Duan group found a capacitance of 365 F g^{-1} for a CoPc-CMP/CNT material.⁵⁷ Lin *et al.* synthesized a CNT@Mn-MOF composite for supercapacitor application.⁵⁸ We hypothesized that combining nano carbonaceous materials with COPs would be a suitable way to develop active electrode materials for supercapacitor application. Therefore, in this study, we blended three different carbon materials (fullerene (C₆₀), MWCNTs, and SWCNTs) into Py-DSDA-COP to improve and compare the electrochemical performance of COP-based supercapacitors. The preparation of Py-DSDA-COP through Schiff base coupling of Py-Ph-CHO and 4,4'-disulfanediyldianiline (DSDA) is shown in Scheme 1. The chemical structure, thermal stability, nature, porosity, and electrochemical performance of three electrode systems and symmetric coin cells are discussed below in detail.

Experimental section

Materials

Potassium carbonate (K₂CO₃), sodium hydroxide (NaOH), pyrene (Py), 4-formylphenylboronic acid (4-FP-BO), and nitrobenzene (C₆H₅NO₂) were purchased from Sigma-Aldrich. Mesitylene, acetic acid (AcOH), 1,4-dioxane (DO), 4,4'-disulfanediyldianiline (DSDA), acetone, and tetrahydrofuran (THF) were obtained from Alfa Aesar.

Preparation of 1,3,6,8-tetrabromopyrene (Py-Br₄)

Py-Br₄ was obtained as a green solid (17 g) by refluxing 80 mL of C₆H₅NO₂, 8 g of Py (40 mmol), and Br₂ (9.2 mL, 176 mmol) at 120 °C for 24 hours, according to Scheme S1.† The FTIR spectrum (Fig. S1†): 3054 cm^{-1} (aromatic CH) and 681 cm^{-1} (C-Br).

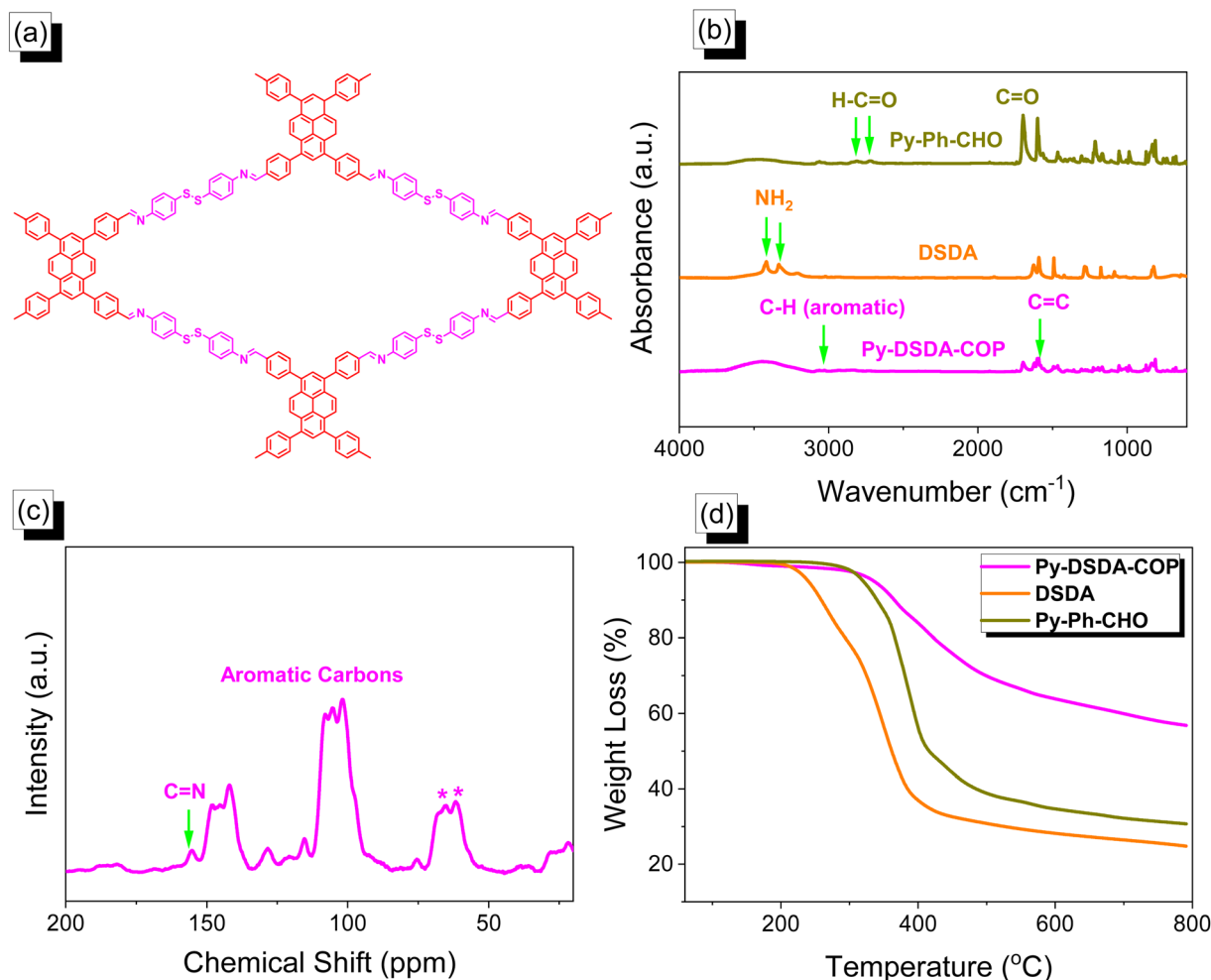


Fig. 1 (a) Chemical structure of Py-DSDA-COP. (b) FTIR spectra of Py-Ph-CHO, DSDA and Py-DSDA-COP. (c) Solid state ¹³C NMR of Py-DSDA-COP. (d) TGA of Py-Ph-CHO, DSDA and Py-DSDA-COP.

Preparation of 1,3,6,8-tetrakis(4-formylphenyl)pyrene (Py-Ph-CHO)

4-FP-BO (4.40 g, 29.00 mmol), K₂CO₃ (5.30 g, 37.86 mmol), Py-Br₄ (2.50 g, 4.83 mmol), and Pd(PPh₃)₄ (0.23 g, 0.25 mmol) were refluxed in 80 mL of DO and 40 mL of H₂O at 110 °C under N₂ for 72 h. After working up the reaction mixture with 100 mL of HCl (2 N), a yellow solid was obtained which was washed with water and MeOH to yield pure Py-Ph-CHO [Scheme 1(a), 2.13 g]. FTIR (cm⁻¹, Fig. 1(a)): 3061, 2813–2732 (aldehydic CH), 1706 (C=O), and 1598 (C=C). ssNMR (Fig. S2[†]): 182.38 (CHO), 143.85–121.36 (aromatic carbons).

Preparation of Py-DSDA-COP

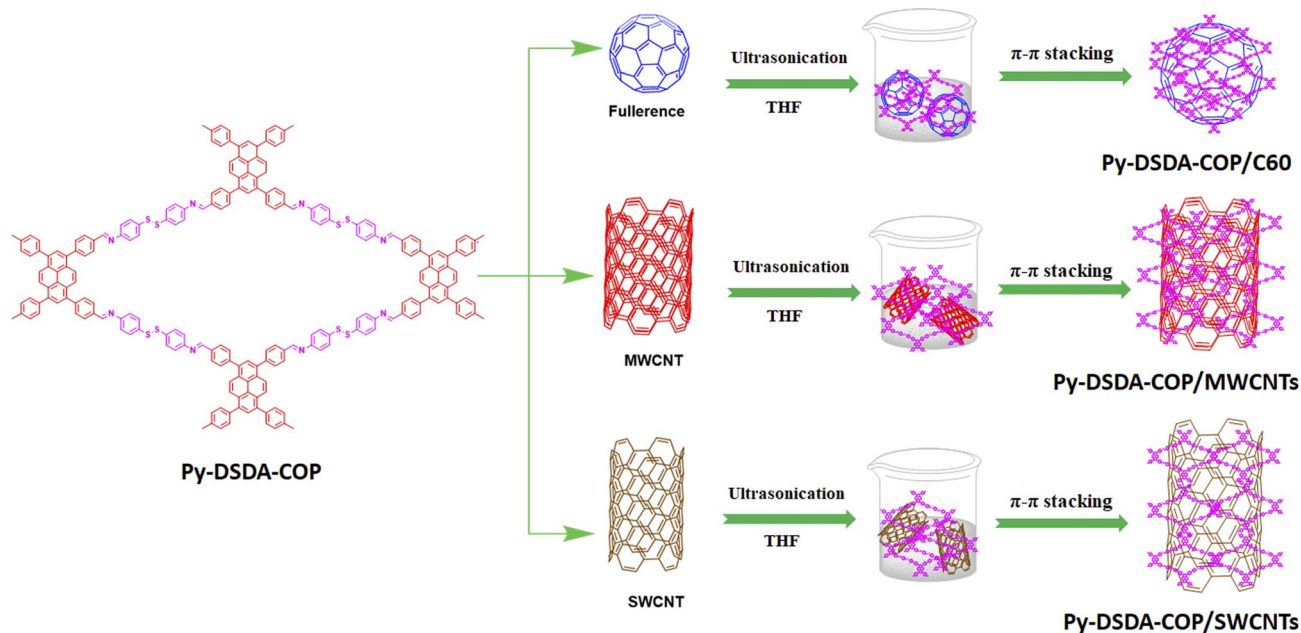
The solution of Py-Ph-CHO (0.15 g, 0.24 mmol), DSDA (0.12 g, 0.48 mmol), mesitylene (5 mL), acetic acid (0.65 mL), H₂O (0.35 mL), and DO (5 mL) was degassed two times under vacuum and then stirred at 75 °C for 72 h, filtered and washed with THF, H₂O, MeOH, and acetone to get a yellow solid, which was dried in a vacuum oven overnight.

Preparation of Py-DSDA-COP/nanocomposites

Py-DSDA-COP (2 wt%), and C₆₀, MWCNTs, and SWCNTs (2 wt%) were individually dissolved in THF and sonicated overnight, and then the solvent was removed by rotary evaporation to afford Py-DSDA-COP/C₆₀, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites, respectively as shown in Scheme 2.

Results and discussion

Pyrene is a polycyclic aromatic hydrocarbon that has been explored for various applications, including its potential use in energy storage due to its high electrical conductivity, surface area, porous structure, and redox activity. These characteristics make pyrene a good candidate to be used as an active electrode material for supercapacitors. The synthesis of a porous COP material based on Py-Ph-CHO and DSDA to afford Py-DSDA-COP is shown in Scheme 1. The Py-Ph-CHO monomer was synthesized *via* the Suzuki coupling reaction of Py-Br₄ with 4-FP-BO in the presence of DO/K₂CO₃/H₂O at 110 °C for 72 h [Scheme 1(a)].



Scheme 2 Preparation of Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs and Py-DSDA-COP/SWCNTs nanocomposites.

Finally, Py-DSDA-COP was obtained by a Schiff base reaction of Py-Ph-CHO and DSDA in the presence of dioxane/mesitylene at 110 °C [Scheme 1(b)]. The resultant porous Py-DSDA-COP

material is insoluble in acetone, methanol, and THF. The chemical structure of Py-DSDA-COP [Fig. 1(a)] was validated through FTIR and ^{13}C solid-state NMR as shown in Fig. 1(b) and

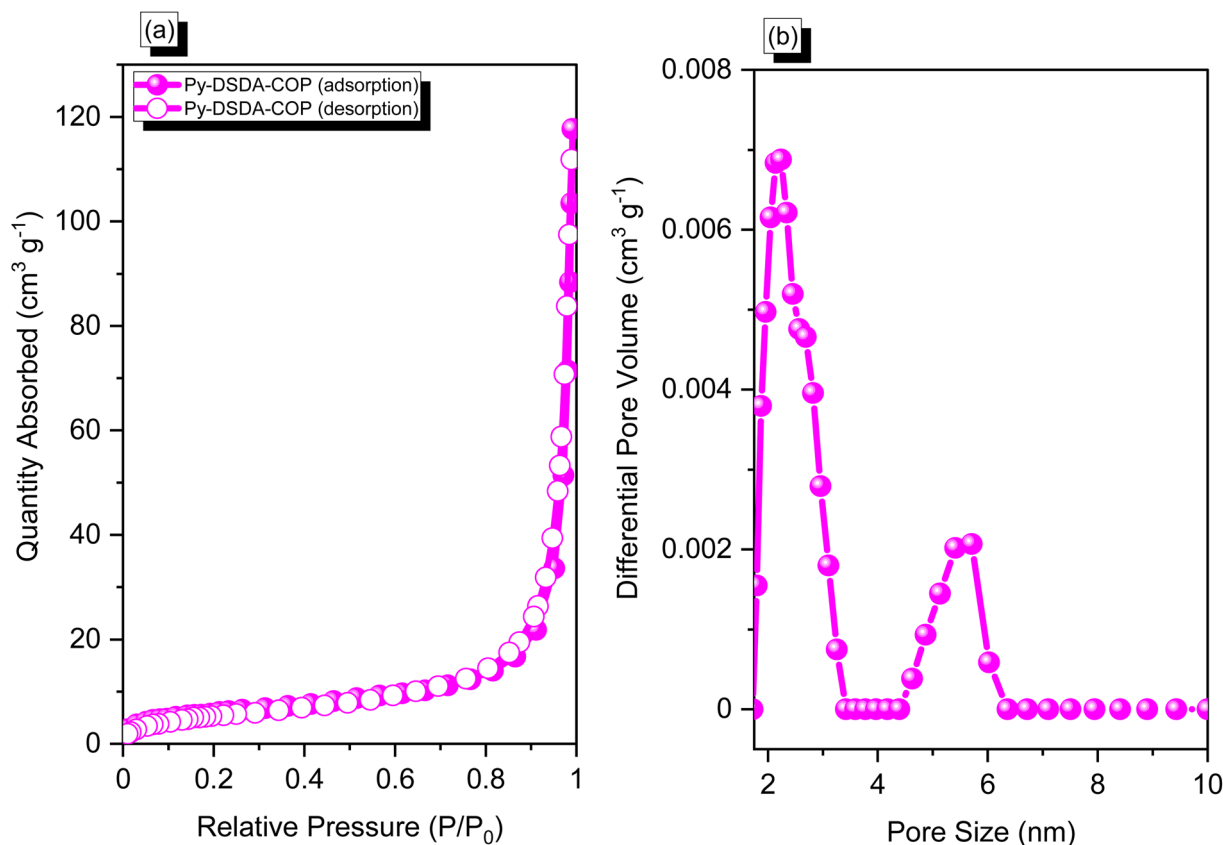


Fig. 2 (a) N_2 adsorption/desorption curve and (b) pore size of Py-DSDA-COP.

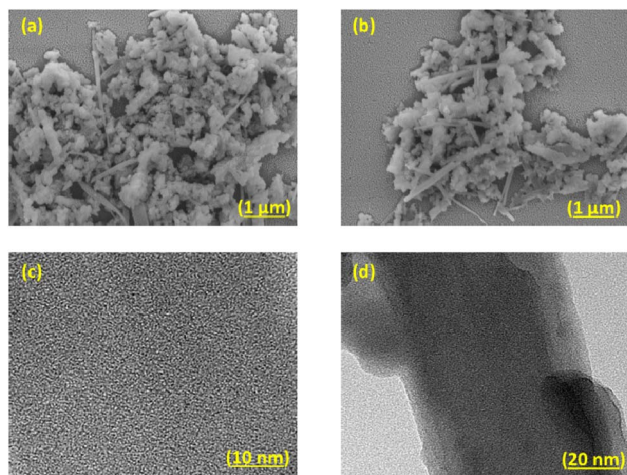


Fig. 3 SEM (a and b) and TEM images (c and d) of Py-DSDA-COP.

(c). The FTIR spectrum of Py-Ph-CHO showed peaks at 2813–2732 and 1706 cm^{-1} , representing the aldehydic C–H and C=O groups, respectively (Fig. 1(b)). The absorption bands of the NH_2 group in the DSDA monomer appeared at 3411 and 3328 cm^{-1} . The FTIR spectrum of Py-DSDA-COP showed a peak at 1605 cm^{-1} corresponding to C=N, with the complete absence of N–H and C=O peaks from the starting monomers Py-Ph-CHO and DSDA, respectively. Furthermore, Py-DSDA-COP exhibited a peak at 1592.11 cm^{-1} representing an aromatic C=C unit. The solid-state NMR of Py-DSDA-COP showed signals at 156.5 ppm for C=N and 148.5–128.0 ppm for

aromatic carbons (Fig. 1(c)). The FTIR and NMR analyses confirmed the successful synthesis of Py-DSDA-COP. The thermal stabilities of Py-DSDA-COP and its ingredients (Py-Ph-CHO and DSDA) were measured through thermogravimetric analysis, as shown in (Fig. 1(d)).

The thermal decomposition temperature (T_{d5} , T_{d10}) and char yield were observed as 240 °C, 258 °C, and 25 wt% for DSDA, while 320 °C, 340 °C, and 31 wt% for Py-Ph-CHO, respectively. We observed that after the Schiff base reaction of Py-Ph-CHO and DSDA, the thermal stability of the resultant porous material (Py-DSDA-COP) improved. The values of T_{d5} , T_{d10} , and char yield were 335 °C, 360 °C, and 57 wt%, respectively. Moreover, FTIR analysis [Fig. S3†] of Py-DSDA-COP shows distinct peaks at 3058 and 1592.11 cm^{-1} , corresponding to the aromatic CH and C=C groups. These peaks serve as evidence of the robustness and integrity of the chemical structure of the Py-DSDA-COP framework, even after exposure to various temperatures ranging from 25 to 180 °C. Notably, the absence of an absorption peak at 2515 cm^{-1} in the Py-DSDA-COP FTIR spectrum, typically associated with –SH groups, further supports the stability of the S–S bond within the material.

The nature of the Py-DSDA-COP material was validated by the powder X-ray diffraction (XRD) pattern, as shown in Fig. S4.† The result revealed the semi-crystallinity behavior of Py-DSDA-COP from the existence of some sharp peaks in its XRD profile. The porosity of Py-DSDA-COP was examined by N_2 absorption at 77 K, as shown in Fig. 2. The Py-DSDA-COP showed a type II isotherm curve according to IUPAC. The Py-DSDA-COP exhibited rapid intake at high pressure demonstrating a mesoporous structure as shown in Fig. 2(a). Furthermore, we observe a S_{BET} , pore size, and pore volume of

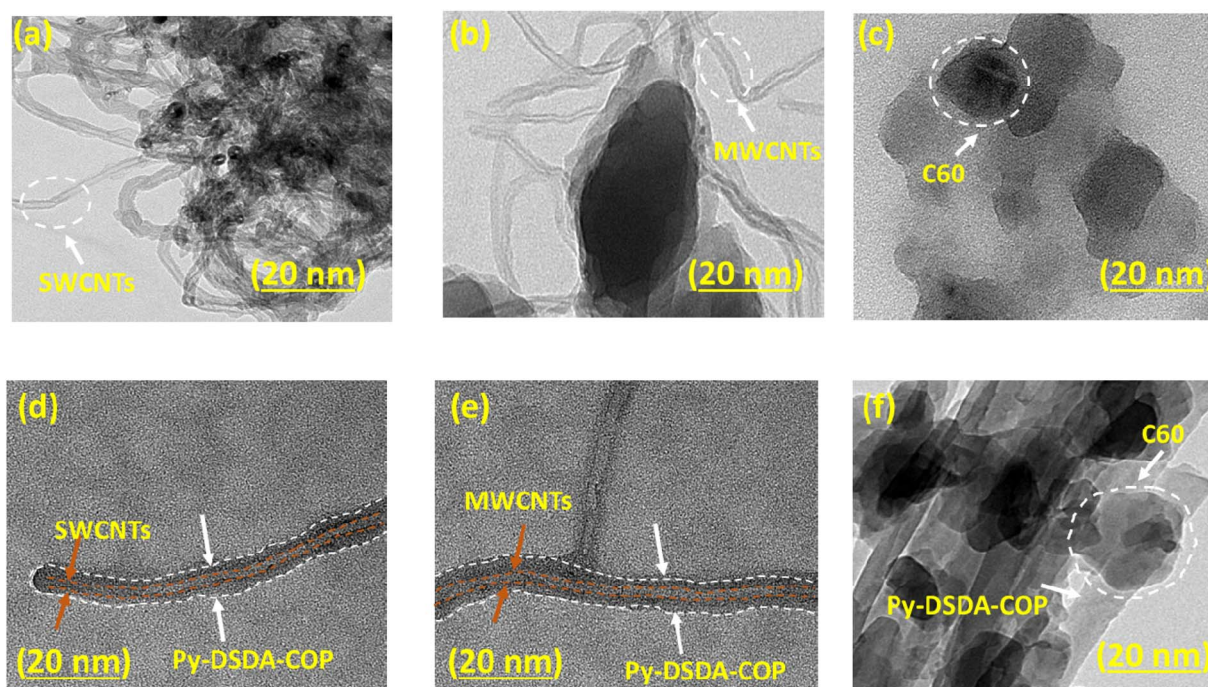


Fig. 4 TEM images of (a) SWCNTs, (b) MWCNTs, (c) C60, (d) Py-DSDA-COP/SWCNTs, (e) Py-DSDA-COP/MWCNTs and (f) Py-DSDA-COP/C60.

$45 \text{ m}^2 \text{ g}^{-1}$, 2.15 nm [Fig. 2(b)], and $0.16 \text{ cm}^3 \text{ g}^{-1}$, respectively. In summary, Py-DSDA-COP showed significant mesoporous characteristics. The SEM images of Py-DSDA-COP revealed a nanorod-like structure with aggregated particles [Fig. 3(a) and (b)], which was also validated by TEM analysis (Fig. 3(c) and (d)) with porous characteristics.

Scheme 2 outlines the process for creating Py-DSDA-COP/nanocomposite materials using π - π stacking interactions between Py-DSDA-COP and C60, MWCNTs, or SWCNTs. The process involves placing the Py-DSDA-COP and the chosen nanomaterial into a THF solution and sonicating the mixture overnight. This results in the formation of Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, or Py-DSDA-COP/SWCNTs nanocomposites, respectively. TEM images of SWCNTs, MWCNTs, and fullerene (C60) and Py-DSDA-COP nanocomposites (Py-DSDA-COP/SWCNTs, Py-DSDA-COP/MWCNTs, Py-DSDA-COP/C60) are shown in (Fig. 4(a-f)). The SWCNTs and MWCNTs were observed as nanotube-like structures, while C60 has a circular shape. The SWCNTs and MWCNTs dispersed very uniformly in Py-DSDA-COP due to excellent π - π stacking interactions between them [Fig. 4(d) and (e)]. But in the case of

C60, poor dispersion was observed with Py-DSDA-COP suggesting no interaction between them (Fig. 4(f)).

Electrochemical performance of Py-DSDA-COP and its nanocomposites with C60, MWCNTs, and SWCNTs

The electrochemical performance of C60, MWCNTs, SWCNTs, Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites was measured by performing cyclic voltammetry (CV) and galvanostatic charge-discharge (GCD) analysis through a three-electrode system (1 M KOH solution). The CV curves of all Py-DSDA-COPs samples were measured at various sweep rates (5 to 200 mV s^{-1}) corresponding to the potential window between 0 and -1 V , as shown in Fig. 5(a-d) and S5(a-c).† The CV curves for MWCNTs and SWCNTs exhibited quasi-rectangular shapes with humps, indicating the presence of electric double-layer capacitance (EDLC) and pseudocapacitive responses.

In contrast, C60 displayed oxidation and reduction peaks with humps, showcasing a more pronounced pseudocapacitive nature along with EDLC.^{59,60} Notably, the CV-integrated area for SWCNTs was significantly larger than that observed for

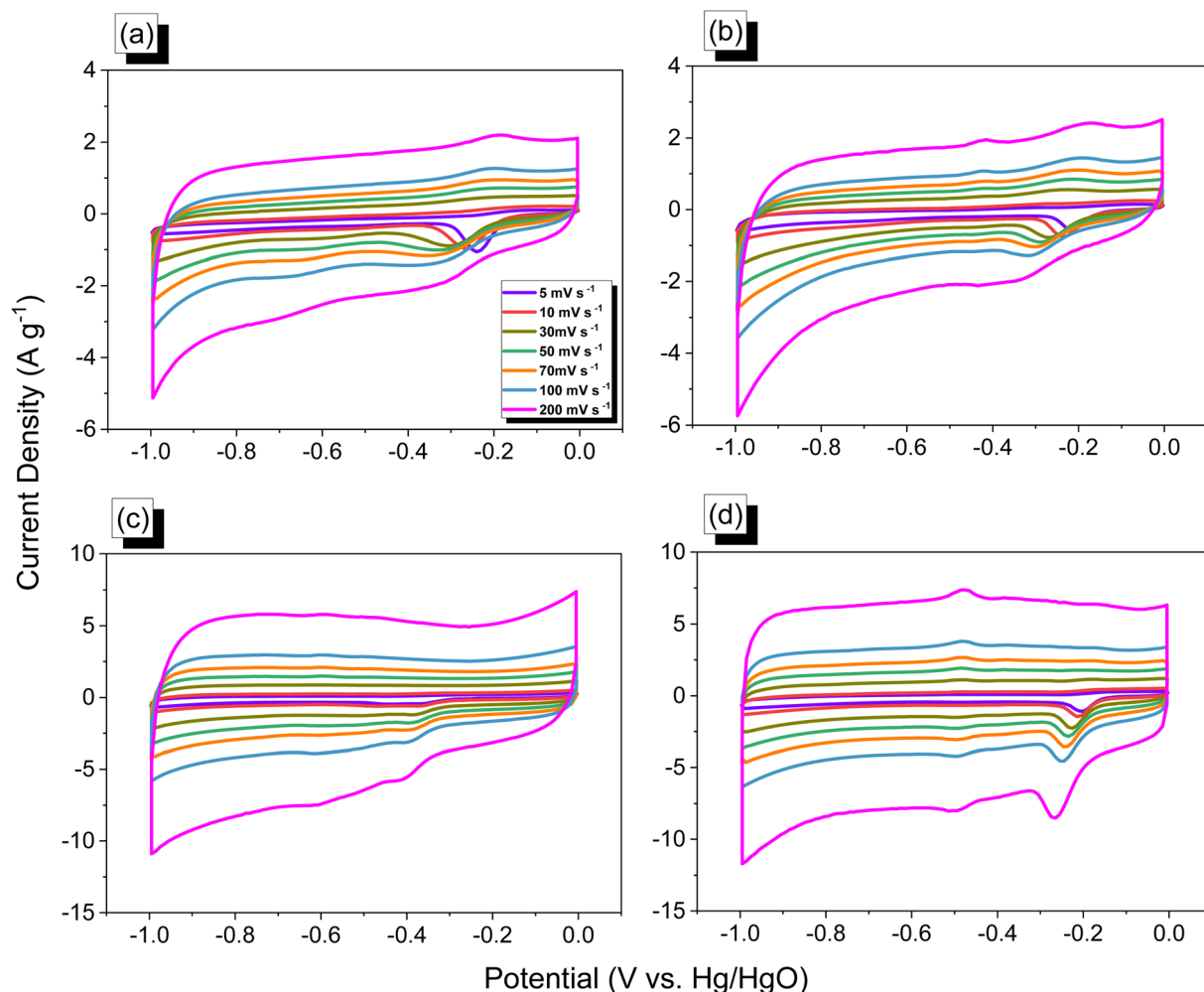


Fig. 5 CV curves of (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

MWCNTs and C60, as illustrated in Fig. S5(a–c).† The CV curves of Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites showed quasi-rectangular curves with humps, revealing that the capacitive feedback was influenced by the EDLC.^{59,60} The presence of humps suggests that faradaic redox currents induced another minor pseudocapacitance effect, due to the blending of nano carbonaceous materials and sulfur species (heteroatom influence) in these Py-DSDA-COPs electrodes.^{55,56} The CV curves of all Py-DSDA-COP electrode samples have redox peaks, corresponding to the pseudocapacitance nature, the presence of the conjugated system, and N and S atoms in the pure Py-DSDA-COP.^{55,56} Furthermore, with increasing sweep rates, current density also increased without affecting CV profiles, suggesting high-rate capability.^{55,56} The CV-integrated area of Py-DSDA-COP/MWCNTs and Py-DSDA-COP/SWCNTs nanocomposites improved as compared to pure Py-DSDA-COP, indicating their higher specific capacitance. The GCD curves of Py-DSDA-COP and its nanocomposite samples were recorded at various current densities from 1 to 20 A g⁻¹, as shown in Fig. 6(a–d) and S5(d–f).† The GCD curves of C60, MWCNTs, and SWCNTs are triangular with slight bends revealing EDLC and

pseudocapacitive behavior.^{†59,60} The specific capacitance of C60, MWCNTs, and SWCNTs at 1 A g⁻¹ was 9.8 F g⁻¹, 31.75 F g⁻¹, and 73.42 F g⁻¹ respectively. The SWCNTs showed higher electrochemical performance than C60 and MWCNTs. These GCD curves of Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites have triangular shapes with a slight bend, demonstrating both pseudocapacitive and EDLC characteristics.^{59,60} The Py-DSDA-COP nanocomposites (Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs) experienced a higher discharging time curve than those of pure Py-DSDA-COP, revealing that after blending of different nanoparticles (C60, MWCNTs, SWCNTs), the specific capacitance of Py-DSDA-COP was improved. The specific capacitance from GCD curves at 1 A g⁻¹ was observed as 56 F g⁻¹, 61 F g⁻¹, 123 F g⁻¹, and 171 F g⁻¹ for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites, respectively (Fig. 7(a)). The Py-DSDA-COP/SWCNTs showed higher specific capacitance than Py-DSDA-COP/MWCNTs, possibly due to the higher EDLC behavior, higher conductivity, and lower diameter of SWCNTs than those of MWCNTs and C60.^{61,62} In addition, based on TEM images, the successful

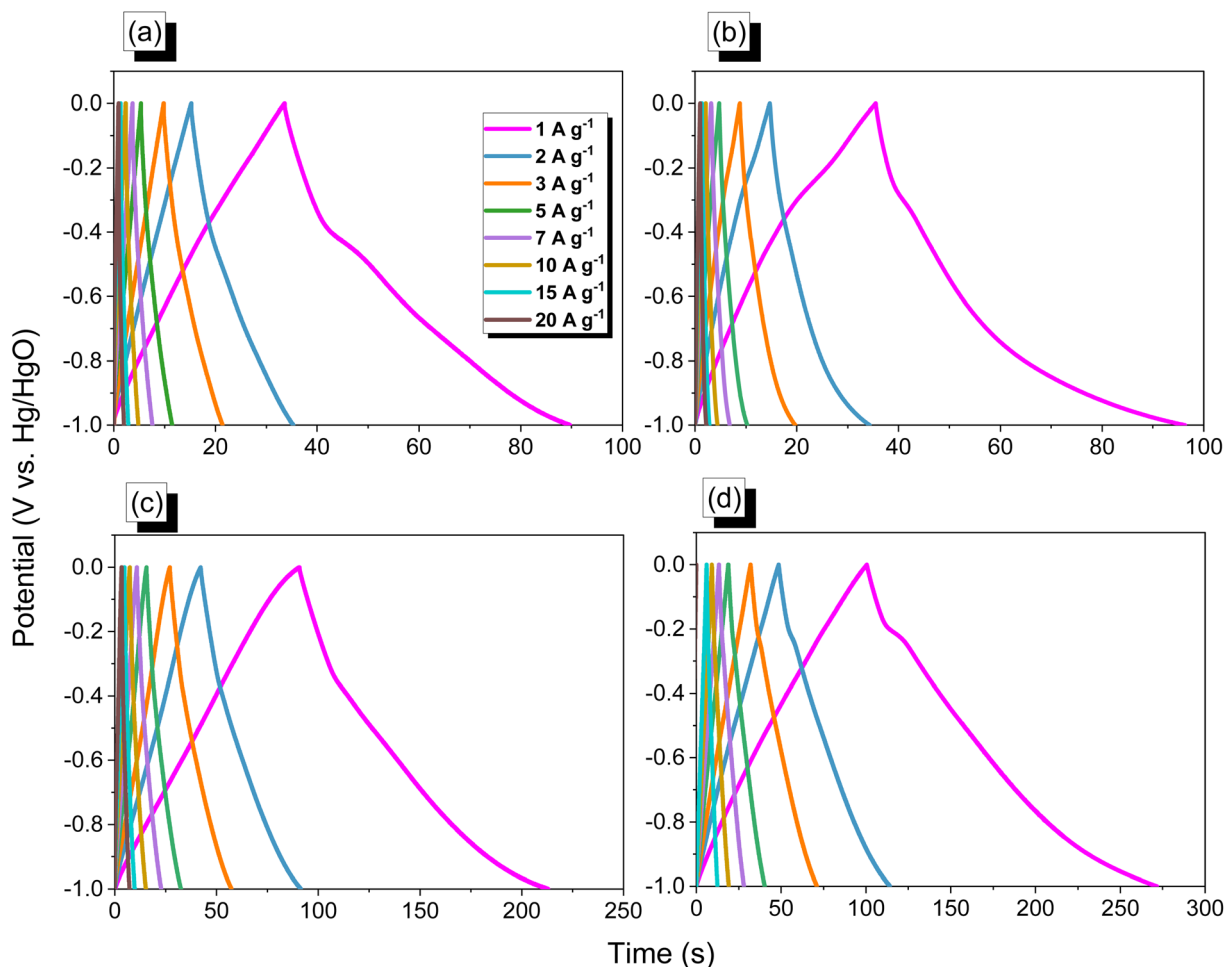


Fig. 6 GCD curves of (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

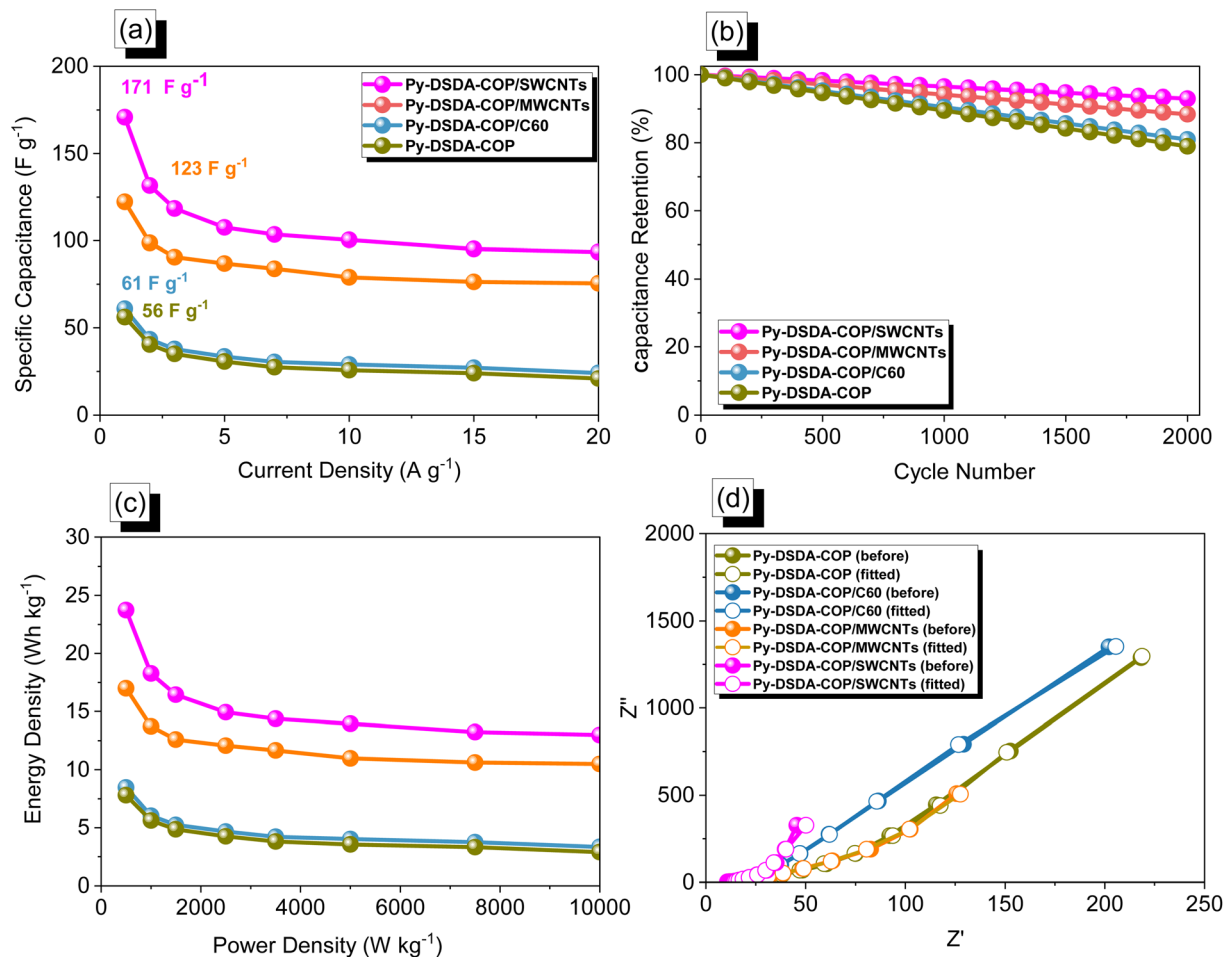


Fig. 7 (a) Specific capacity, (b) stability, (c) Ragone profiles, and (d) Nyquist plots for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs.

dispersion of Py-DSDA-COP within SWCNTs results in enhanced ion mobility within the nanotubes, leading to improved ion migration efficiency and an increased capacitance. Moreover, the specific capacitance of Py-DSDA-COP didn't enhance much by incorporating C60, due to their poor interaction as discussed above in the TEM analysis. The electrochemical performance evaluation of Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs was conducted using three electrode systems, and this assessment was repeated three times. The specific capacitance values were determined from the GCD curves at a current density of $1 A g^{-1}$ and were found to be $56.18 F g^{-1}$ for Py-DSDA-COP, $60.96 F g^{-1}$ for Py-DSDA-COP/C60, $122.3 F g^{-1}$ for Py-DSDA-COP/MWCNTs, and $170.8 F g^{-1}$ for Py-DSDA-COP/SWCNTs (Fig. S6[†]). Numerous materials with high capacitance values, such as MWCNT@SACMP ($594 F g^{-1}$ at $1.0 A g^{-1}$),⁶³ pNTCDA-TPAT ($217.4 F g^{-1}$ at $0.5 A g^{-1}$),⁶⁴ PTPA@MWNT-4 ($410 F g^{-1}$),⁶⁵ and CNT@TFA-COF-3 ($338 F g^{-1}$),⁶⁶ have been reported in the literature. The stabilities of Py-DSDA-COP and their nanocomposites were also evaluated through GCD at $10 A g^{-1}$ over 2000 cycles (Fig. 7(b)). The

capacitance retention of Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites was observed as 79%, 81%, 88%, and 93%, respectively, demonstrating the high stability of Py-DSDA-COP/MWCNTs. This revealed that the stability of Py-DSDA-COP improved after blending with C60, MWCNTs, and SWCNTs. The nanorod-like structure of Py-DSDA-COP remained intact, even after undergoing 2000 cycles at a rate of $10 A g^{-1}$, as illustrated in Fig. S7[†]. We also display the Ragone plot (Fig. 7(c)), by calculating energy and power density [eqn (S2)[†]]. The energy density of the Py-DSDA-COP/SWCNTs nanocomposite ($23.7 W h kg^{-1}$) is much higher than those of Py-DSDA-COP/MWCNTs ($16.9 W h kg^{-1}$), Py-DSDA-COP/C60 ($8.4 W h kg^{-1}$) and Py-DSDA-COP ($28 W h kg^{-1}$). Furthermore, the internal resistance of these electrode materials was also investigated by electrochemical impedance spectroscopy (EIS). The fitted Nyquist plots (Fig. 7(d)) were used to determine series resistance (R_s) from an equivalent circuit (Fig. S8[†]). The series resistance was observed as 10.78, 11.02, 13.6, and 6.6 Ω for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites, respectively. The

Py-DSDA-COP/SWCNTs displayed the smallest resistance and hence had superior electrochemical performance to the other three COP electrode materials. The blending of different carbonaceous materials in Py-DSDA-COP resulted in excellent electrochemical performance compared to other porous materials (Table S1†).

Electrochemical performance of symmetric coin cells based on Py-DSDA-COP and its nanocomposites with C60, MWCNTs, and SWCNTs

The electrochemical performance of Py-DSDA-COP and its nanocomposites were also investigated by preparing symmetric CR2032 coin cells at various sweep rates (5 to 200 mV s^{-1}) corresponding to the potential window between 0.4 and -0.6 , as shown in Fig. 8. All CV curves showed quasi-rectangular curves with humps demonstrating EDLC and pseudocapacitance effect

(Fig. 8(a-d)).^{55,56} The GCD curves of all electrode materials are triangular with bend-assisted EDLC and pseudocapacitance characteristics (Fig. 9(a-d)). The specific capacitance from GCD curves at 1 A g^{-1} was observed as 17 F g^{-1} , 19 F g^{-1} , 54 F g^{-1} , and 79 F g^{-1} for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites respectively (Fig. 10(a)). The incorporation of SWCNTs into Py-DSDA-COP led to higher specific capacitance due to their good dispersion, higher EDLC behavior, higher conductivity, and lower diameter than those of MWCNTs, while C60 showed poor dispersion in Py-DSDA-COP, revealing no interaction between them. The stability of Py-DSDA-COP and its nanocomposites were also evaluated through GCD at 10 A g^{-1} over 2000 cycles for symmetric coin cells (Fig. 10(b)), showing a capacitance retention of 95%, 89.4%, 80.6%, and 78.4% for Py-DSDA-COP/SWCNTs, Py-DSDA-COP/MWCNTs, Py-DSDA-COP/C60, and Py-DSDA-COP, respectively.

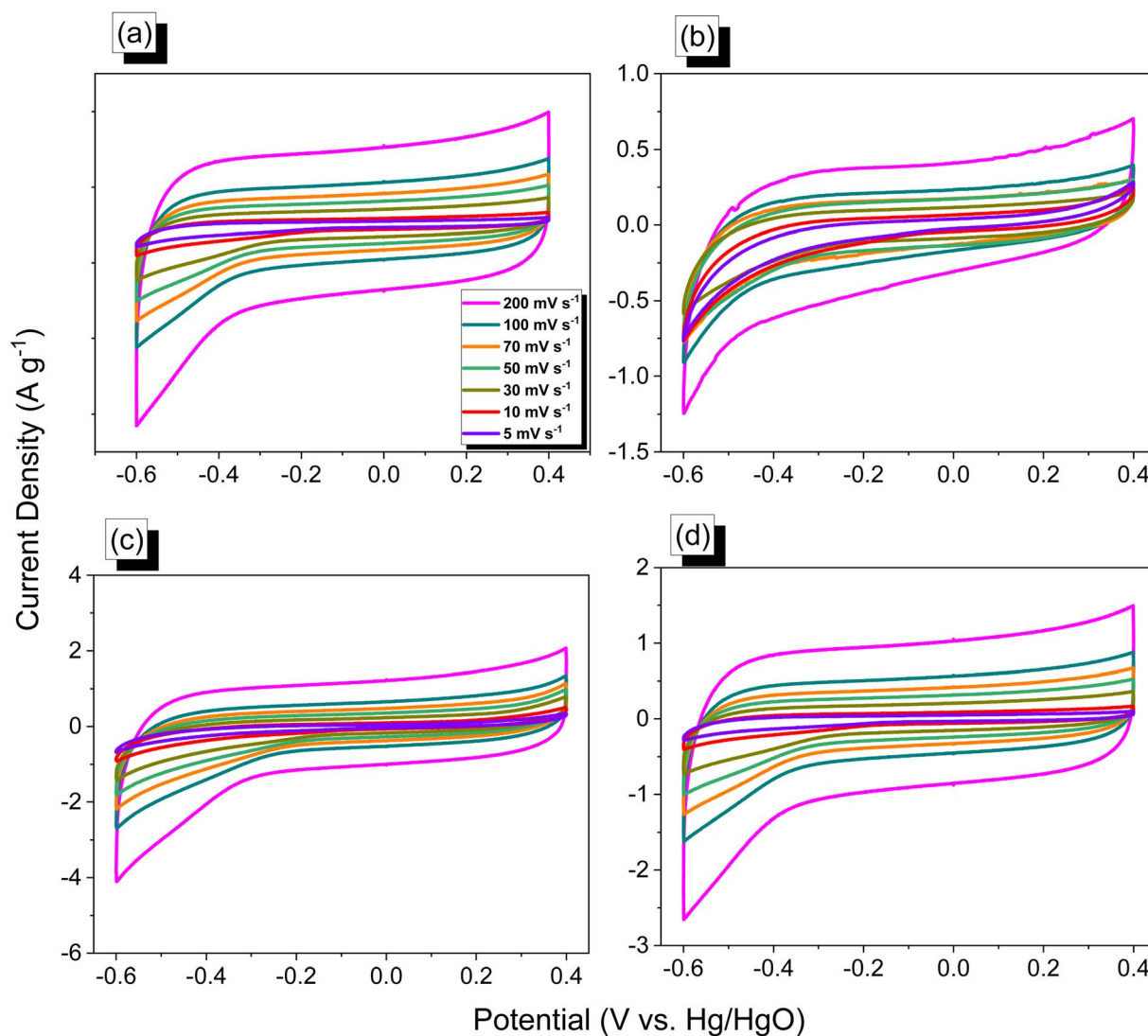


Fig. 8 CV curves of symmetric coin cells based on (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

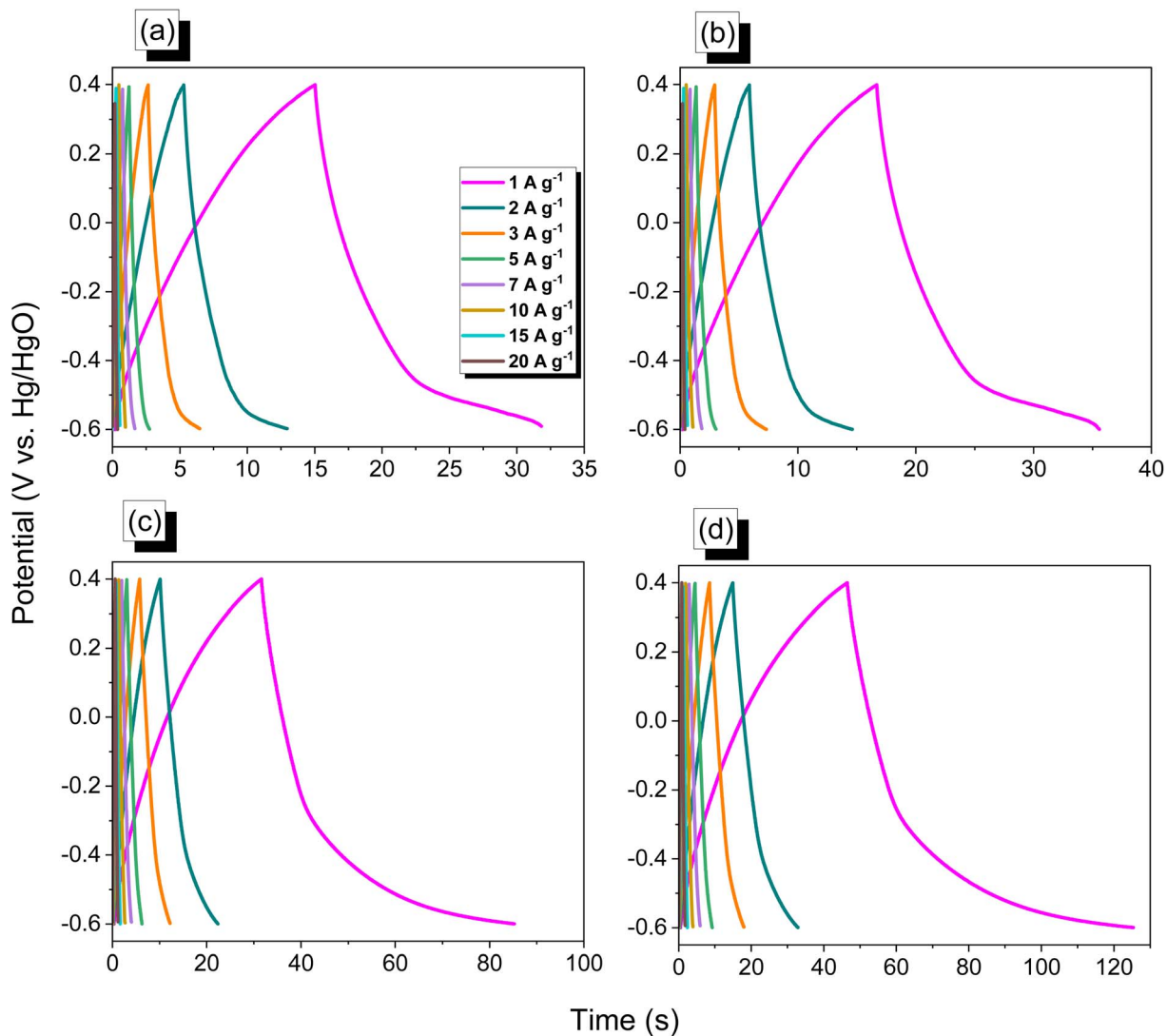


Fig. 9 GCD curves of symmetric coin cells based on (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

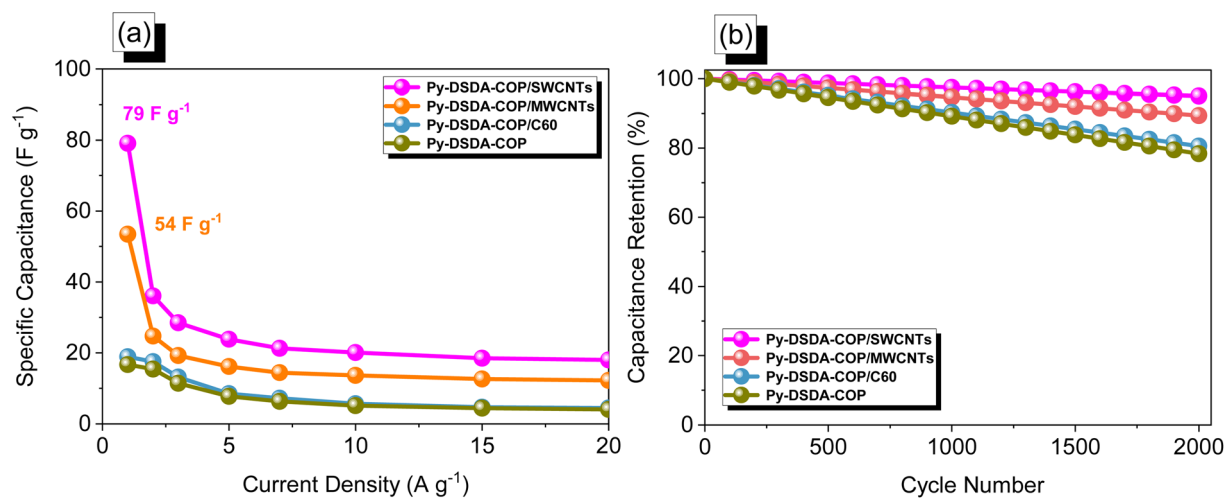


Fig. 10 (a) Specific capacity and (b) stability of symmetric coin cells based on Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs.

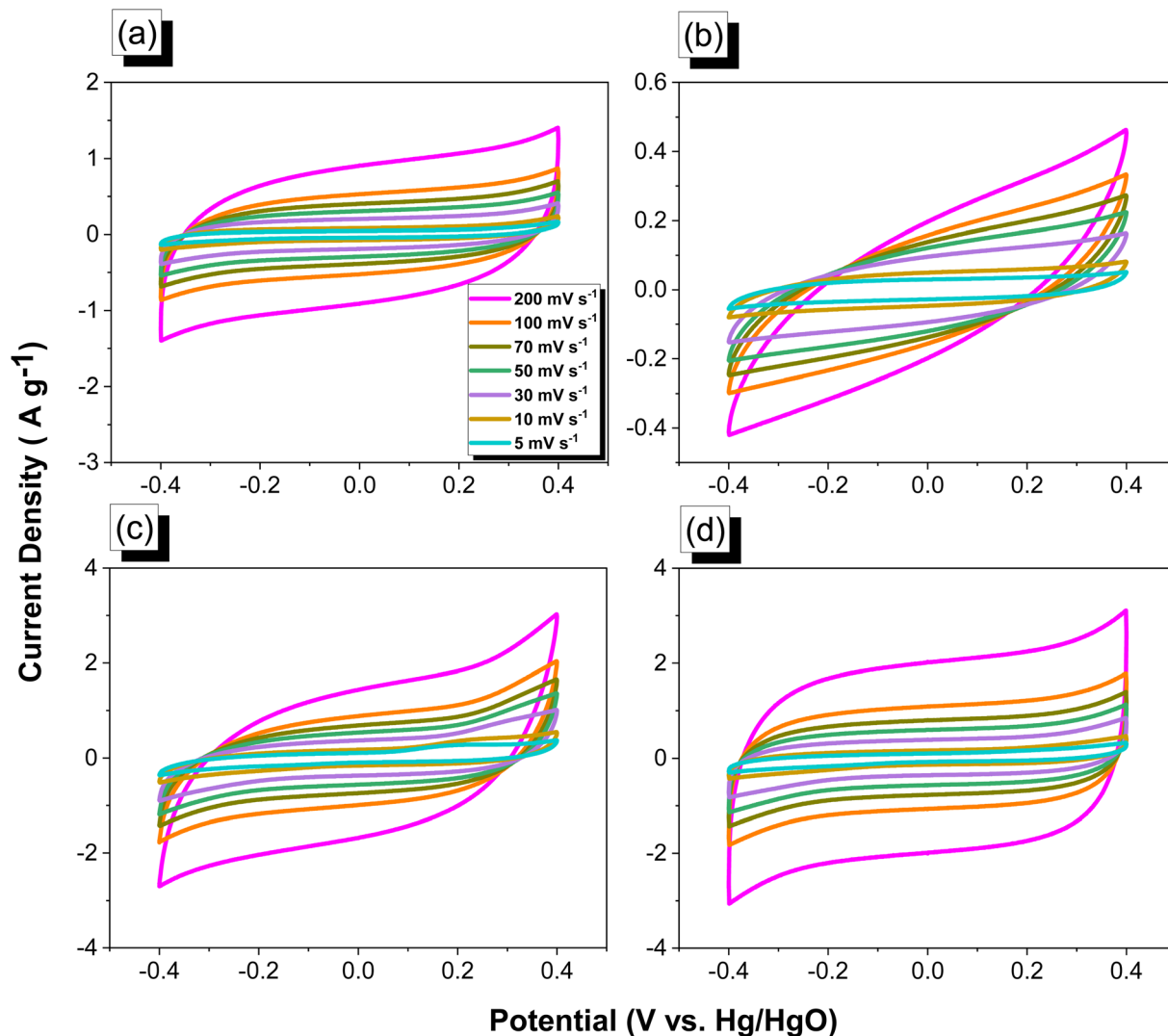


Fig. 11 CV curves of asymmetric coin cells based on (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

Electrochemical performance of asymmetric coin cells based on Py-DSDA-COP and its nanocomposites with C60, MWCNTs, and SWCNTs

We conducted experiments involving asymmetric coin cells using Py-DSDA-COP and its nanocomposites to examine their electrochemical performance. In these coin cells, activated carbon served as the negative electrode, while Py-DSDA-COP or its nanocomposites acted as the positive electrode. We conducted tests at various sweep rates, ranging from 5 to 200 mV s^{-1} , within a potential window spanning from 0.4 to -0.4 V, as illustrated in Fig. 11. The CV curves for Py-DSDA-COP, Py-DSDA-COP/MWCNTs, Py-DSDA-COP/SWCNTs, and Py-DSDA-COP/C60 exhibited nearly rectangular shapes with slight humps, as depicted in Fig. 11(a–d). However, Py-DSDA-COP/C60 displayed a non-rectangular shape. These variations in CV curve shapes were primarily due to differences in potential windows when compared to three-electrode and symmetric coin cells. GCD

analysis showed that all electrode materials exhibited triangular curves with bends, indicating a combination of EDLC and pseudocapacitive characteristics, as shown in Fig. 12(a–d). Specific capacitance measurements at 1 A g^{-1} revealed values of 30.52 F g^{-1} for Py-DSDA-COP, 36.57 F g^{-1} for Py-DSDA-COP/C60, 79.21 F g^{-1} for Py-DSDA-COP/MWCNTs, and 113.17 F g^{-1} for Py-DSDA-COP/SWCNTs nanocomposites, as demonstrated in Fig. 13(a). Notably, Py-DSDA-COP/SWCNTs exhibited the highest specific capacitance among the prepared nanocomposite electrode materials, consistent with the results obtained using three-electrode and symmetric coin cell configurations. To assess the stability of these electrode materials, we subjected them to 10 A g^{-1} cycling for 2000 cycles in asymmetric coin cells, as shown in Fig. 13(b). The capacitance retention percentages for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites were 78%, 82%, 88%, and 95%, respectively.

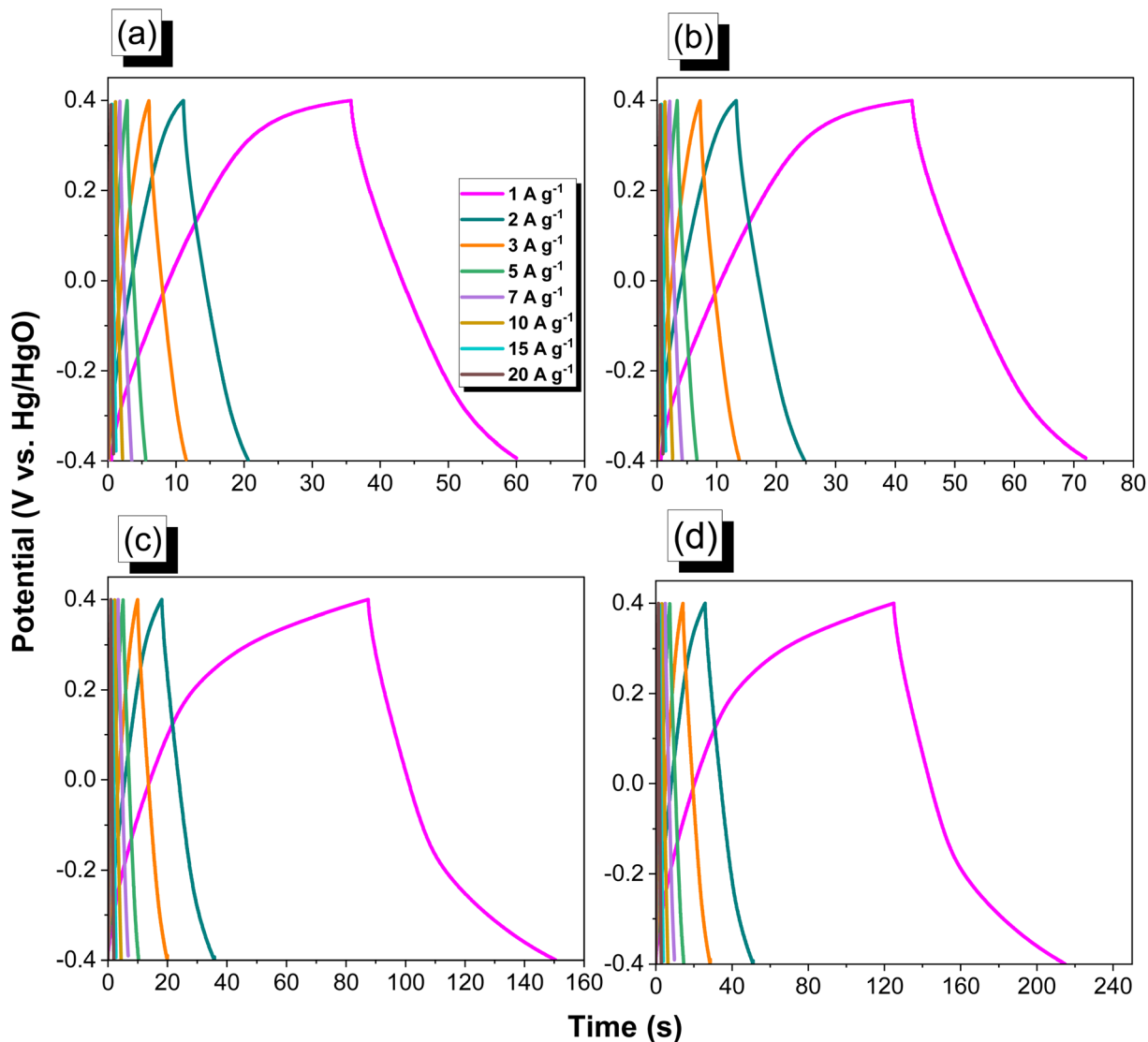


Fig. 12 GCD curves of asymmetric coin cells based on (a) Py-DSDA-COP, (b) Py-DSDA-COP/C60, (c) Py-DSDA-COP/MWCNTs and (d) Py-DSDA-COP/SWCNTs.

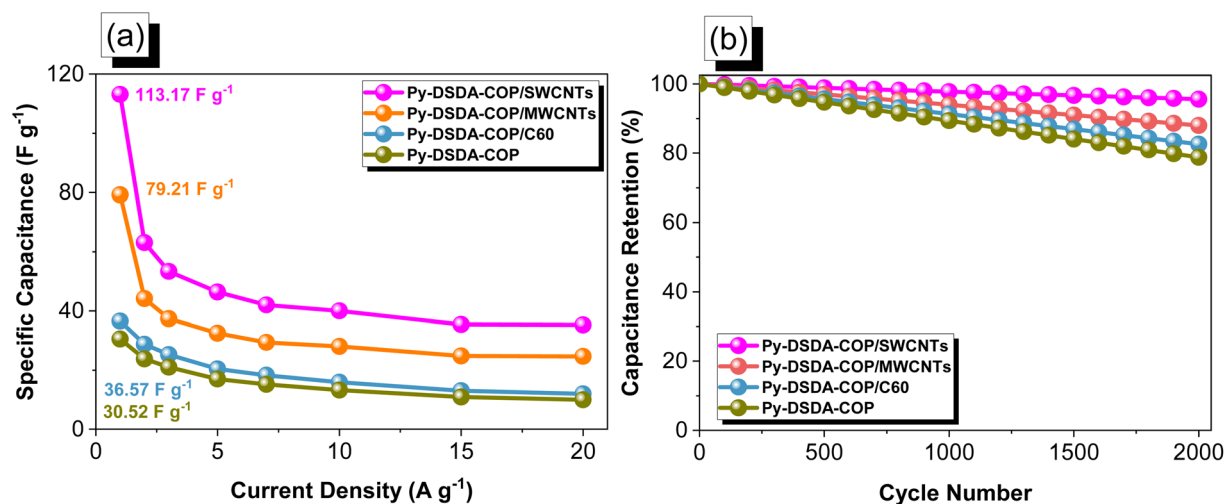


Fig. 13 (a) Specific capacity and (b) stability of asymmetric coin cells based on Py-DSDA-COP, Py-DSDA-COP, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs.

Conclusion

We used the Schiff base reaction to synthesize Py-DSDA-COP and investigate it as an electrode material for supercapacitors, demonstrating good porosity and a specific capacitance of 56.18 F g^{-1} at 1 A g^{-1} . We incorporated different carbonaceous materials into Py-DSDA-COP to afford Py-DSDA-COP/SWCNTs, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/C60 nanocomposites to improve and compare the electrochemical performance of these electrode materials. The Py-DSDA-COP/SWCNTs displayed superior specific capacitance (171 F g^{-1}) at 1 A g^{-1} and higher energy density (23.7 W h kg^{-1}) with a capacity retention of 93% after 2000 cycles than those of Py-DSDA-COP/MWCNTs (123 F g^{-1}) and Py-DSDA-COP/C60 (61 F g^{-1}) for three electrode configurations. For symmetric coin cells, the specific capacitance was observed as 79, 54, 19, and 17 F g^{-1} for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites, respectively. For asymmetric coin cells, the specific capacitance was observed as 30.52, 36.57, 79.21, and 113.17 F g^{-1} for Py-DSDA-COP, Py-DSDA-COP/C60, Py-DSDA-COP/MWCNTs, and Py-DSDA-COP/SWCNTs nanocomposites, respectively. The superior electrochemical properties of Py-DSDA-COP/SWCNTs are attributed to the good dispersion, higher EDLC behavior, higher conductivity, and lower diameter of SWCNTs compared to MWCNTs, while C60 doesn't show any compatibility with Py-DSDA-COP.

Conflicts of interest

There are no conflicts to declare.

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