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Solid state chemical transformations through ring-opening polymerization of ferrocene-based conjugated microporous polymers in host–guest complexes with benzoxazine-linked cyclodextrin

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ABSTRACT

Herein, we describe two ferrocene-derived conjugated microporous polymers (FFC—CMPs) prepared through Sonogashira couplings of 9-ferrocenylidene-2,7-dibromo-9*H*-fluorene (FFC) with tetraethynylpyrene (Py-T) and tetrakis(4-ethynylphenyl)ethene (TPE-T), respectively, and their properties determined using various spectroscopic techniques. These two FFC CMPs formed inclusion complexes (ICs) with a benzoxazine-linked β -cyclodextrin (CD-BZ), with host-guest interactions occurring between the ferrocene and β -CD units. We used Fourier transform infrared spectroscopy, X-ray diffraction, two-dimensional nuclear Overhauser enhancement NMR spectroscopy, solid-state ¹³C CP/MAS NMR spectroscopy, and thermogravimetric analysis to characterize these FFC—CMP/CD-BZ ICs. The BZ units in the FFC—CMP/CD-BZ ICs underwent thermal ring-opening polymerization to form new Mannich bridges featuring both intermolecular (OH…O) and intramolecular (OH…N) hydrogen bonds; the resulting TPE-FFC—CMP/poly(CD-BZ) and Py-FFC—CMP/poly (CD-BZ) ICs exhibited outstanding thermal stability, with the latter having the higher CO₂ uptake ability (1.42 mmol g⁻¹) and capacitance (46 F g⁻¹ at 0.5 A g⁻¹).

Background: Host–guest complexes formed from macrocyclic hosts possessing hydrophobic cavities such as cyclodextrins (CDs) and various guest molecules have been studied widely for their fascinating characteristics. The interaction of ferrocene-containing CMPs with hydrophobic cavities of CDs to form inclusion complexes materials with excellent thermal properties and these materials can be used in different fields including energy storage and gas capture.

Methods: Two ferrocene-based fluorene CMPs—TPE-FFC CMP and Py-FFC—CMP—through Sonogashira couplings of 9-ferrocenylidene-2,7-dibromo-9*H*-fluorene (FFC), as the main block, with tetraethynylpyrene (Py-T) and tetrakis(4-ethynylphenyl)ethene (TPE-T), were successfully synthesized through Sonogashira couplings. Then, the two FFC CMPs were attached to a benzoxazine-functionalized β -cyclodextrin (CD-BZ) through host-guest complexation. The successful synthesis of two FFC CMPs and FFC—CMP/CD-BZ ICs was investigated by using Fourier transform infrared spectroscopy, X-ray diffraction, two-dimensional nuclear Overhauser enhancement NMR spectroscopy, solid-state ¹³C CP/MAS NMR spectroscopy, and thermogravimetric analysis.

Significant Findings: The present work offers a new and facile strategy for the preparation CMPs/CD inclusion complexes through supramolecular chemistry and using these materials as an electrode in the energy devices and CO_2 uptake.

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1. Introduction

E-mail address: kuosw@faculty.nsysu.edu.tw (S.-W. Kuo). ¹ These authors contributed equally to this work. Conjugated microporous polymers (CMPs) are an emerging class of porous materials having extended conjugated skeletons in their polymeric frameworks. They are attractive because of their wide applicability in several fields, including supercapacitors, metal-ion

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2

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13

batteries, hydrogen evolution, energy storage, fuel cells, sensors, photocatalysis, phototherapy, CO₂ adsorption, and optoelectronics [1-18]. CMPs can be prepared—with a range of microporous morphologies presenting various molecular structures of conjugated polvmers-using a variety of synthetic methods, including Heck reactions, Schiff-base formation, oxidative coupling, Sonogashira coupling, cyclotrimerization, Yamamoto and Suzuki coupling [19–27]. The diversity of CMP structures leads to many having attractive properties, including high conductivities, mechanical rigidities, surface areas, porosities, and thermal stabilities. Interestingly, solid-state chemical transformations of CMPs have been performed without changing their topologies significantly, typically through post-modification of the linkages of the CMPs. For example, imine-linked CMPs have been converted to secondary amides, secondary amines, oxazoles, thiazoles, and quinolones; as a result, amide- and oxazolelinked CMPs can be synthesized from their building blocks directly [28-36]. In addition, cyclic carbamate- and thiocarbamate-linked CMPs have been obtained from imine-linked CMPs through multistep post-modifications in the solid state, retaining the high surface areas $(>780 \text{ m}^2 \text{ g}^{-1})$ and long-range order of the crystalline and porous structures of their precursors [37].

Benzoxazine (BZ) units undergo ring-opening polymerization (ROP) to form polybenzoxazines (PBZs) featuring strong intramolecular (OH…N) and intermolecular (OH…O) hydrogen bonds, thereby enhancing the thermal stabilities and dielectric constants [38–46]. In previous studies, we synthesized porous organic polymers featuring BZ-linkages through direct one-pot Mannich reactions and multistep syntheses using tetraphenylethylene (TPE) as the building block [47,48]. We performed post-modifications of the linkages of these CMPs through the formation of covalent bonds; to the best of our knowledge, noncovalent bonding has not been reported previously to introduce BZ-linkages into CMPs.

Supramolecular interactions are exploited in many different fields, including catalysis, biochemistry, organic synthesis, and electronics. They involve various types of intermolecular noncovalent bonding, typically van der Waals interactions, metal coordination, hydrophobic interactions and π -stacking [49–52]. Host–guest complexes formed from macrocyclic hosts possessing hydrophobic cavities [e.g., cyclodextrins (CDs), calixarenes, and cyclophanes] and various guest molecules have been studied widely for their fascinating characteristics [53, 54]. By allowing noncovalent interactions to exist between their components, various supramolecular host-guest inclusion complexes (ICs) can be formed in a facile manner for practical use. CDs are among the most popular macrocyclic receptors used to introduce supramolecular interactions; they are cyclic oligomers of α -1,4-D-glucopyranose, with α -, β -, and γ -CD being most common, differentiated by their ring sizes. The hydrophobic cavities of CDs interact with various guest molecules, including azobenzene, polyethylene glycol, adamantane, and ferrocene units, forming ICs [55,56]. β -CD has been the most widely used of the CDs because of its commercial availability, unique structure, outstanding biocompatibility, and facile chemical modification [55]. Ferrocene units can be incorporated into the hydrophobic cavity of β -CD through supramolecular hydrophobic interactions, forming ICs of equal mole ratio. For example, Yuan et al. prepared supramolecular graft copolymers featuring host–guest interactions between β -CD in the main chain and ferrocene units and investigated their responsiveness for electrochemical redox applications [57]. Guo et al. obtained amphiphilic polymers from β -CD dextran and a ferrocene-terminated polycaprolactam, based on a host-guest strategy; they used cyclic voltammetry (CV) and two-dimensional (2D) NMR spectroscopy to confirm the presence of the ICs formed between the two units [58]. Ni and coworkers fabricated a graft copolymer (Dex- β -CD/Fc-PLA) through host-guest complexation of building blocks featuring ferrocene and β -CD moieties [59].

The introduction of organometallic compounds (in particular, ferrocene) in the skeletons of CMPs can lead to excellent electrical, optical, sensing, catalytic, magnetic, thermal, and redox properties. Indeed, ferrocene-containing CMPs have been used in various applications, including gas storage, lithium batteries, dye removal, catalysis, magnetic switches, and memory devices [60–67]. For example, Liu et al. used Yamamoto coupling to prepare two ferrocene-based CMPs displaying good thermal stability and gas storage capacity [68]. Wong et al. synthesized ferrocene-based hyperbranched polymers through Sonogashira coupling of 9-ferrocenylidene-2,7-diiodo-9*H*fluorene and tri(4-ethynylphenyl)amine; by controlling the morphology of these synthesized polymers, they obtained nanostructures suitable for use in water treatment and lithium-ion batteries [69].

In this study, we synthesized two ferrocene-based fluorene CMPs—TPE-FFC CMP and Py-FFC—CMP—through Sonogashira couplings of 9-ferrocenylidene-2,7-dibromo-9*H*-fluorene (FFC), as the main block, with tetraethynylpyrene (Py-T) and tetrakis(4-ethynylphenyl)ethene (TPE-T), respectively [Schemes 1(a) and 1(b)]. We then attached the two FFC CMPs to a benzoxazine-functionalized β -cyclodextrin (CD-BZ) [70] through host—guest complexation of the β -CD and ferrocene moieties to form FFC/CD-BZ CMP ICs [Schemes 1 (c) and 1(d)]. The formation of the ICs between the FFC—CMPs and β -CD was confirmed using 2D nuclear Overhauser enhancement (2D NOESY) NMR spectroscopy. To the best of our knowledge, these FFC—CMP/CD-BZ ICs are the first examples of CMPs possessing BZ units introduced through host—guest interaction of FFC and β -CD units; furthermore, these BZ units underwent ROP upon thermal treatment, without the need for a catalyst or curing agent [52].

2. Experimental

2.1. Materials

Potassium hydroxide (KOH), triphenylphosphine (PPh₃), and copper iodide (CuI) were purchased from Alfa Aesar. Tetrakis(triphenylphosphine)palladium(0) [Pd(PPh₃)₄] was obtained from Sigma–Aldrich. Ethanol (EtOH), triethylamine (Et₃N), and *N*,*N*-dimethylformamide (DMF), methanol (MeOH), tetrahydrofuran (THF), and ferrocene carboxaldehyde were obtained from Acros. Tetrakis(4-bromophenyl)ethylene (TPE-Br₄), 1,3,6,8-tetrabromopyrene (Py-Br₄), 2,7-dibromo-9*H*-fluorene (F-Br₂), 1,3,6,8-tetraethynylpyrene (Py-T) 1,1,2,2-tetrakis(4-ethynylphenyl)ethene (TPE-T), FPy-CMP, and CD-BZ (Scheme S2) were prepared using previously reported procedures [1,16,48,70-74].

2.2. FTPE-CMP

A solution of PPh₃ (6.0 mg, 0.023 mmol), Pd(PPh₃)₄ (27 mg, 0.023 mmol), F-Br₂ (0.15 g, 0.46 mmol), TPE-T (0.10 g, 0.23 mmol), and CuI (4.0 mg, 0.023 mmol) in dry DMF and Et₃N (14 mL) was degassed (using vacuum pump and N₂) and then heated under reflux at 110 °C (oil bath) for 3 days. The orange precipitate was filtered off, washed thoroughly with DMF, MeOH, and THF, and then dried in an oven for 2 days [Scheme S1(a)]. FTIR (KBr, cm⁻¹): 3059 (aromatic C–H stretching), 2191 (C=C stretching), 1601 (C=C stretching) [Figure S1(a)].

2.3. FPy-CMP

A solution of Pd(PPh₃)₄ (27 mg, 0.023 mmol), F-Br₂ (0.15 g, 0.46 mmol), PPh₃ (6.0 mg, 0.023 mmol), Py-T (0.1 g, 0.23 mmol), and CuI (4.0 mg, 0.023 mmol) in dry DMF and Et₃N (14 mL) was degassed (using a vacuum pump and N₂) and then heated under reflux at 110 ° C (oil bath) for 3 days. The orange precipitate was filtered off, washed thoroughly with DMF, MeOH, and THF, and then dried in an oven for 2 days [Scheme S1(b)]. FTIR (KBr, cm⁻¹): 3059 (aromatic C–H stretching), 2191 (C=C stretching), 1601 (C=C stretching) [Figure S1(b)].

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13

2.4. 9-Ferrocenylidene-2,7-dibromo-9H-fluorene (FFC)

A mixture of F-Br₂ (0.80 g), KOH (0.27 g), and ferrocene carboxaldehyde (0.58 g) in EtOH (50 mL) was heated under reflux at 70 °C for 24 h. The red precipitate was washed with EtOH and dried in an oven at 80 °C for 24 h (Scheme 1). FTIR (KBr, cm⁻¹): 3095 (aromatic C–H stretching). ¹H NMR (500 MHz, CDCl₃, δ , ppm, Figure S2): 8.40, 7.88, 7.56, 7.56, 7.46, 4.72, 4.56, 4.26. ¹³C NMR (125 MHz, CDCl₃, δ , ppm, Figure S3): 141.6, 138.4, 138.3, 135.9, 131, 130.5, 130.1, 129.7, 127, 122.8, 121, 120.9, 120.6, 80, 70.9, 70.6, 69.7. (+)ESI-MS: *m/z* 521 (C₂₃H₁₆FeBr₂) (Figure S4).

2.5. TPE-FFC-CMP

A solution of CuI (8.0 mg, 0.040 mmol), FFC (0.24 g, 0.46 mmol), Pd (PPh₃)₄ (50 mg, 0.045 mmol), TPE-T (0.10 g, 0.23 mmol), and PPh₃ (12 mg, 0.050 mmol) in dry DMF and Et₃N (14 mL) was degassed (using a vacuum pump and N₂) and then heated under reflux at 110 °C (oil bath) for 3 days. The brown solid was filtered off and washed thoroughly

with DMF, MeOH, and THF. FTIR (KBr, cm^{-1}): 3095 (aromatic C–H stretching), 2197 (C=C stretching), 1628 (C=C stretching) [Fig. 1(b)].

2.6. Py-FFC-CMP

This CMP was prepared as a dark-red precipitate by using the procedure described above for TPE-FFC—CMP, but with the following changes: PPh₃ (20 mg, 0.060 mmol), FFC (0.34 g, 0.66 mmol), Py-T (0.10 g, 0.33 mmol), and CuI (0.01 g, 0.05 mmol). FTIR (KBr, cm⁻¹): 3095 (aromatic C–H stretching), 2197 (C=C stretching), 1628 (C=C stretching) [Fig. 1(c)].

2.7. CD-BZ/FFC and FFC/cd-BZ CMP ICs and thermal curing polymerization

CD-BZ (0.1 g) and FFC, Py-FFC—CMP, or TPE-FFC—CMP (0.05 g) were placed in distilled water (10 mL) and then the mixture was sonicated for 2 h and stirred for 12 h to form the CD-BZ/FFC and FFC/CD-BZ CMP ICs, respectively. Each FFC/CD-BZ CMP IC (0.15 g) was then



Scheme 1. Schematic representations of (a) Py-FFC—CMP, (b) TPE-FFC—CMP, (c) Py-FFC/CD-BZ CMP, (d) TPE-FFC/CD-BZ CMP, (e) Py-FFC/CD-BZ CMP after curing, and (f) TPE-FFC/CD-BZ CMP after curing.

4

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Fig. 1. FTIR spectra of (a) CD-BZ, FFC, and the FFC/CD-BZ complex; (b) CD-BZ, TPE-FFC—CMP, and the TPE-FFC/CD-BZ CMP; and (c) CD-BZ, Py-FFC—CMP, and the Py-FFC/CD-BZ CMP.

cured at 300 $^\circ C$ for 4 h to afford the FFC/poly(CD-BZ) CMP as a dark-brown solid.

3. Results and discussion

3.1. Synthesis of FFC monomer

The FFC monomer was prepared through condensation of F-Br₂ and ferrocene carboxaldehyde in absolute EtOH in the presence of KOH (Scheme 1). The chemical structure of the FFC monomer was confirmed using conventional techniques (FTIR, ¹H and ¹³C NMR, and FTMS spectroscopy; Figures S2–S4). The FTIR spectrum [Fig. 1(a)] featured a peak at 1628 cm⁻¹ representing C=C stretching, but no characteristic peak for a carboxyl group, revealing that the condensation of F-Br₂ and ferrocene carboxaldehyde had given the target FFC monomer, isolated as a red solid. The ¹H NMR spectrum of the FFC monomer (Fig. S2) featured signals at 4.26, 4.56, and 4.72 ppm representing the Cp protons of the ferrocene moiety and at 7.46–7.88 and 8.40 ppm representing the aromatic and HC=C protons, respectively.

The ¹³C NMR spectrum of the FFC monomer (Figure S3) featured peaks at 69.7, 70.6, 70.9, and 80.0 ppm representing the carbon nuclei of the Cp units, as well as signals at 120.6–138.4 and 141.6 ppm representing the carbon nuclei of the aromatic and HC=C units, respectively. The molecular weight of the FFC monomer (m/z 521), determined using (+)ESI mass spectrometry (Figure S4), was consistent with its expected calculated value (520.7).

3.2. Synthesis of Py-FFC-CMP and TPE-FFC-CMP

The two ferrocene-based fluorene conjugated microporous polymers (FFC—CMPs) were prepared through Sonogashira couplings of the FFC monomer with Py-T and TPE-T, respectively, in DMF/Et₃N with Pd(PPh₃)₄ as the catalyst [Schemes 1(a) and 1(b)]. The FTIR

spectra of these two CMPs featured peaks at 2197 cm⁻¹ for -C=Cstretching and at 3095 cm⁻¹ for C–H aromatic vibrations [Fig. 1(b) and 1(c)]. Fig. 2(b) and 2(d) present the solid-state ¹³C CP/MAS NMR spectra of Py-FFC and TPE-FFC—CMPs with signal assignments. The peaks in the range from 69.2 to 92 ppm represented the alkyne and ferrocene units in both FFC—CMP frameworks; the signals at 120 and 129–140 ppm represent the carbon nuclei of the HC=C and aromatic units, respectively. These spectral data are consistent with the chemical structures of both FFC—CMPs.

We recorded Brunauer-Emmett-Teller (BET) isotherms at 77 K to investigate the porosities of FTPE-CMP, FPy-CMP, TPE-FFC-CMP, and Py-FFC—CMP (Fig. 3). The isotherms of FTPE-CMP [Fig. 3(a)] and FPy-CMP [Fig. 3(e)] reveal mesoporous structures for these CMP polymers (type IV according to IUPAC classification). The surface area (S_{BET}) and total pore volume (V_{total}) for FTPE-CMP were 31 m² g⁻¹ and 0.06 cm 3 g ${}^{-1}$, respectively; for FPy-CMP they were 191 m 2 g ${}^{-1}$ and 0.17 cm³ g⁻¹, respectively. Nonlocal density functional theory (NL-DFT) revealed that the pore size distributions of FTPE-CMP and FPy-CMP were 1.24-2.37 [Fig. 3(b)] and 2.14-5.85 [Fig. 3(f)] nm, respectively. The TEM images in Fig. 3(c) and 3(g) reveals that these two CMPs were mesoporous with a few microporous structures having pore sizes of 1.5–2.5 nm. We suspect that the BET surface area and pore volume of FPy-CMP were higher than those of FTPE-CMP because a pyrene unit is more planar than a TPE unit, allowing the former to feature stronger π -stacking and a more ordered structure. After attaching ferrocene units to these two CMP polymers, TPE-FFC-CMP [Fig. 3(a)] and Py-FFC-CMP [Fig. 3(e)] provided similar type IV curves for their mesoporous structures; the values of S_{BET} and V_{total} decreased, however, to 8 m² g⁻¹ and 0.04 cm³ g⁻¹, respectively, for TPE-FFC—CMP and to 50 $m^2 \ g^{-1}$ and 0.07 $cm^3 \ g^{-1}$, respectively, for Py-FFC-CMP. The pore size distributions were 0.47-2.83 nm for TPE-FFC-CMP [Fig. 3(b)] and 1.96-5.48 nm for Py-FFC-CMP [Fig. 3 (f)], as confirmed through TEM imaging [Fig. 3(d) and 3(h)] of the

5

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13



Fig. 2. ¹³C NMR spectra of (a) CD-BZ, (b) TPE-FFC-CMP, (c) TPE-FFC/CD-BZ CMP, (d) Py-FFC-CMP, and (e) Py-FFC/CD-BZ CMP.

microporous structures. Thus, we suspect that the ferrocene units attached to these two CMP polymers as side groups occupied their porous spaces, thereby decreasing the surface areas and pore volumes. The hysteresis loops for all of the CMP materials were not closed because of the flexibility of the polymeric frameworks and their swelling through gas adsorption upon elastic deformation; this behavior has been observed for several CMPs [11,57].

Although the total pore volume and surface area both decreased after attaching the ferrocene units to these two CMPs, the thermal properties of TPE-FFC—CMP and Py-FFC—CMP both increased significantly, based on TGA measurements performed under N₂ at a heating rate of 20 °C min⁻¹ (Fig. 4). For example, FTPE-CMP provided a value of T_{d10} of 357 °C and a char yield of 59.6 wt% at 800 °C, but these values increased significantly for TPE-FFC—CMP to 527 °C and 67.3 wt%, respectively. Furthermore, FPy-CMP provided a value of T_{d10} of 321 °C and a char yield of 53.2 wt% at 800 °C, but for Py-FFC—CMP they had increased significantly to 470 °C and 74.7 wt%, respectively. These values suggest outstanding thermal stability and rigidity for TPE-FFC—CMP and Py-FFC—CMP. Table 1 summarizes the BET and TGA data of these four CMP polymers.

Fig. 5 displays the X-ray diffraction (XRD) patterns of our two FFC—CMPs; weak diffraction peaks appeared at 13.3°, indicative of their amorphous structures. We used SEM (Figure S5) to study the morphologies of these two FFC—CMPs; the images revealed the aggregation of spherical structures.

3.3. Preparation and thermal curing polymerization of FFC/CD-BZ and FFC/CD-BZ CMP ICs

Because ferrocene forms an IC with β -CD, our FFC—CMPs readily underwent attachment of their ferrocene units through host–guest interactions with the water-soluble BZ-functionalized CD derivative CD-BZ in reasonable yield. The formation of the ICs was evidenced by the solution progressively becoming turbid; the uncomplexed CD-BZ was then removed by washing the ICs with acetone and water. To confirm that ICs could form between the FFC—CMPs and CD-BZ, we used 2D NOESY NMR spectroscopy to investigate the interaction of an equimolar mixture of the low-molecular-weight model compound FFC and CD-BZ (Fig. 6). Several NOE correlation peaks (green squares) existed between the signals of the protons H₃ and H₅ of CD-BZ and the signals of the protons of the substituent and Cp units (4.1–4.5 ppm) of FFC, consistent with a host–guest arrangement of FFC and CD-BZ. XRD, FTIR spectroscopy, TGA, and solid-state ¹³C CP/ MAS NMR spectroscopy provided further evidence for the formation of the FFC—CMP/CD-BZ ICs.

The FTIR spectra of the three ICs in Fig. 1 reveal signals for both CD-BZ and the FFC derivatives, implying that both components were present. The broad signal for the OH groups of CD-BZ near 3383 cm⁻¹ was slightly shifted to 3396 cm⁻¹ in the spectra of the ICs, consistent with the presence of host-guest complexation of CD-BZ and the FFC derivatives. Fig. 2 displays solid-state ¹³C CP/MAS NMR spectra of CD-BZ, TPE-FFC—CMP, and Py-FFC—CMP individually and of their corresponding ICs. The spectra of their ICs feature signals for both CD-BZ and the FFC-CMPs, again implying that both components were present in the ICs. The signals of the glucose units in the spectrum of the pure CD-BZ were resolved [Fig. 2(a)], whereas in the spectra of the ICs they were unresolved [Fig. 2(c) and 2(e)], suggesting the formation of a symmetrical cyclic conformation for the CD in the ICs, with each glucose unit experiencing a similar chemical environment. Fig. 5 displays the XRD patterns of CD-BZ, TPE-FFC—CMP, and Py-FFC CMP individually and of their corresponding ICs. The XRD pattern of the pure CD-BZ revealed a crystalline structure, with peaks at 12.4, 18.4, and 24.4°[70]. The peaks for CD-BZ were broader than those of pure β -CD, suggesting that the crystalline structure was slightly distorted







Fig. 3. (a, e) N₂ adsorption/desorption isotherms of (a) FTPE-CMP and TPE-FFC—CMP and (e) FPy-CMP and Py-FFC—CMP. (b, f) Pore size distribution curves of (b) FTPE-CMP and TPE-FFC—CMP and (f) FPy-CMP and Py-FFC—CMP. (c, d, g, h) TEM images of (c) FTPE-CMP, (d) TPE-FFC—CMP, (g) FPy-CMP, and (h) Py-FFC—CMP.





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6

JID: JTICE

(a)

CD-BZ

TPE-FFC-CMP

TPE-FFC/CD-BZ CMP

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Fig. 5. XRD patterns of (a) CD-BZ, TPE-FFC/CD-BZ CMP, TPE-FFC/CD-BZ CMP, and TPE-FFC/Poly(CD-BZ) CMP and (b) CD-BZ, Py-FFC-CMP, Py-FFC/CD-BZ CMP, and Py-FFC/Poly(CD-BZ) CMP.





Table 1

JID: JTICE

BET and TGA data for the CMPs prepared in this study.

Sample	Surface area (m ² /g)	Pore size (nm)	$T_{d10}(^{0}C)$	Char Yield (%)
FTPE-CMP	31	1.24–2.37	357	59.6
TPE-FFC—CMP	8	0.47–2.83	527	67.3
FPy-CMP	191	2.14–5.85	321	53.2
Py-FFC—CMP	50	1.96–5.48	470	74.7

as a result of one of the OH groups being substituted by a BZ unit. In contrast to the pattern of the pure CD-BZ, those of both FFC—CMP/ CD-BZ ICs featured amorphous halos, consistent with CD-BZ having formed ICs with the FFC units of the CMP frameworks, as displayed in Schemes 1(c) and 1(d); the CD-BZ units were isolated and, therefore, both FFC—CMP/CD-BZ ICs possessed amorphous structures.

The FFC/CD-BZ and both FFC—CMP/CD-BZ ICs were also thermally stable, based on TGA analyses (Fig. 7). The FFC/CD-BZ IC provided a value of T_{d10} of 315 °C and a char yield of 27.6 wt% at 800 °C; these values were, as expected, between those of the pure CD-BZ (T_{d10} = 236 °C; char yield = 7.4 wt%) and pure FFC (T_{d10} = 347 °C; char yield = 31.3 wt%) [Fig. 7(a)]. We observed similar phenomena for the TPE-FFC—CMP/CD-BZ IC [T_{d10} = 311 °C; char yield = 47.2 wt%; Fig. 7 (b)] and the Py-FFC—CMP/CD-BZ IC [T_{d10} = 322 °C; char yield = 61.1 wt%; Fig. 7(c)], with values again being between those of the pure CD-BZ and the corresponding FFC—CMPs. Furthermore, the TGA traces of all of the ICs revealed three-step thermal degradation: water desorption in the first step (ca. 3 wt% loss) at 100–200 °C, the FFC

derivatives decomposing primarily in the second step at 200–400 °C, and CD-BZ and the remaining FFC decomposing mainly in the final step at temperatures above 400 °C. All of the characterization data obtained through these FTIR spectroscopic, solid-state NMR spectroscopic, XRD, and TGA analyses were consistent with the formation of FFC—CMP/CD-BZ ICs.

The FFC—CMP/CD-BZ ICs reported herein are the first examples of CMP materials possessing BZ units assembled through host-guest interactions of FFC and β -CD units. Most interestingly, the BZ units of these FFC—CMP/CD-BZ ICs could undergo ROP, readily converting to phenolic and tertiary amino functional groups through one-step chemical transformations in the solid state [Schemes 1(e) and 1(f)]; this process has never been reported previously for CMPs featuring host-guest interactions with an acceptable surface area and pore volume. To investigate the ROP behavior of the FFC-CMP/CD-BZ ICs in the solid state, we recorded their XRD patterns (Fig. 5); the XRD patterns of the FFC-CMP/poly(CD-BZ) ICs were both similar to those of the uncured FFC-CMP/CD-BZ ICs, indicating that no morphological change had occurred during their ROPs. Furthermore, Fig. 8 displays the FTIR spectra and TGA traces of the FFC-CMP/CD-BZ ICs before and after ROP. Fig. 8(a) and 8(b) reveals that the peaks at 937 and 1157 cm⁻¹ for the oxazine ring and C–O–C stretching, respectively, both disappeared after thermal curing polymerization of both FFC-CMP/CD-BZ ICs. All of the signals become broad after the thermal ROPs, consistent with cross-linked network structures that restricted the molecular vibrations of both FFC-CMP/poly(CD-BZ) ICs. We also investigated the thermal stabilities of the FFC-CMP/CD-



Fig. 7. TGA traces of (a) CD-BZ, FFC, and the FFC/CD-BZ complex; (b) CD-BZ, TPE-FFC—CMP, and the TPE-FFC/CD-BZ CMP; and (c) CD-BZ, Py-FFC—CMP, and the Py-FFC/CD-BZ CMP.

⁸

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13



Fig. 8. (a, b) FTIR spectra of (a) TPE-FFC/CD-BZ CMP and TPE-FFC/Poly(CD-BZ) CMP and (b) Py-FFC/CD-BZ CMP and Py-FFC/Poly(CD-BZ) CMP. (c, d) TGA traces of (c) TPE-FFC/CD-BZ CMP and TPE-FFC/Poly(CD-BZ) CMP and (d) Py-FFC/CD-BZ CMP and Py-FFC/Poly(CD-BZ) CMP.

BZ ICs before and after their thermal ROPs [Fig. 8(c) and 8(d); Table S1]. After thermal ROP of the TPE-FFC/poly(CD-BZ) CMP IC, the value of T_{d10} (364 °C) and the char yield (50.1 wt%) were both higher than those prior to the thermal ROP (T_{d10} = 311 °C; char yield = 47.2 wt%); similarly, the Py-FFC/poly(CD-BZ) CMP IC also exhibited a higher value of T_{d10} (366 °C) and a higher char yield (62.1 wt%) after its thermal ROP (prior to ROP: T_{d10} = 322 °C; char yield = 61.1 wt%). Thus, both of the FFC—CMP/poly(CD-BZ) ICs had outstanding thermal decomposition temperatures, presumably due to their greater number of intramolecular (OH…N) and intermolecular (OH…O) hydrogen bonds after their ROPs. In addition, the thermal stabilities of these two FFC—CMP/poly(CD-BZ) ICs were also higher than those of FTPE-CMP (T_{d10} = 357 °C) and FPy-CMP (T_{d10} = 321 °C), in the absence of the FFC/poly(CD-BZ) IC units; thus, the thermal curing of the CD-BZ units did, indeed, improve the thermal stability.

3.4. CO_2 uptake

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In addition to developing a new approach for the solid-state chemical transformations of CMPs, through the application of host–guest interactions of FFC and β -CD units and subsequent ROPs of their BZ units, we suspected that the new functionality in the CMPs–the cavities and glucose units of the β -CD moieties and the phenolic OH units and Mannich bridges of the BZ moieties–might be capable of interacting with molecules of CO₂. Fig. 9 and Table S1 summarize the CO₂ capture, recorded at 298 °C and 1 bar, of FFC–CMP and the FFC–CMP/CD-BZ ICs before and after their

ROPs. Fig. 9(a) reveals that the CO₂ uptake abilities of TPE-FFC-CMP and the TPE-FFC-CMP/CD-BZ and TPE-FFC-CMP/poly (CD-BZ) ICs were 1.17, 0.81, and 1.31 mmol g^{-1} , respectively; Fig. 9 (b) reveals that for Py-FFC—CMP and the Py-FFC—CMP/CD-BZ and Py-FFC-CMP/poly(CD-BZ) ICs these values were 1.89, 1.04, and 1.42 mmol g^{-1} , respectively. The CO₂ capture ability of Py-FFC-CMP was higher than that of TPE-FFC-CMP because the former had a higher pore volume and surface area-generally favorable for CO₂ uptake in a porous material lacking polar functional groups. The CO₂ capture ability decreased for both FFC-CMP/CD-BZ ICs, presumably because the attached CD-BZ units decreased the surface area and pore volume. The CO₂ capture abilities of the FFC—CMP/poly(CD-BZ) ICs after their ROPs were, however, higher than those of their corresponding FFC-CMP/CD-BZ ICs, presumably because the high abundance of N atoms and phenolic OH groups derived from the BZ units were capable of acid/base (N···C=O), hydrogen bonding (OH···O=C), or other high-affinity interactions with CO₂ [48]. Interestingly, the CO₂ uptake of the TPE-FFC—CMP/poly(CD-BZ) IC after ROP (1.31 mmol g^{-1}) was higher than that of TPE-FFC—CMP (1.17 mmol g^{-1}), but lower than that of the Py-FFC—CMP/poly(CD-BZ) IC (1.42 mmol g^{-1}) after ROP, which itself was lower than that of Py-FFC-CMP which might be attributed to degrade some functional groups during the curing process and low N atoms contents inside Py-FFC-CMP/poly(CD-BZ) IC. As a result, the overall CO₂ capture abilities of our CMP materials were strongly correlated to their pore volumes, the surface chemical functional groups, and surface areas.

10

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Fig. 9. CO2 capture of (a) TPE-FFC/CD-BZ CMP, TPE-FFC/CD-BZ CMP, and TPE-FFC/Poly(CD-BZ) CMP and (b) Py-FFC/CD-BZ CMP, Py-FFC/Poly(CD-BZ) CMP.



Fig. 10. (a, b) CV curves of (a) Py-FFC/CD-BZ CMP and (b) Py-FFC/Poly(CD-BZ) CMP. (c, d) GCD curves of (c) Py-FFC/CD-BZ CMP and (d) Py-FFC/Poly(CD-BZ) CMP, measured at various current densities. (e) Specific capacitances of Py-FFC/CD-BZ CMP and Py-FFC/Poly(CD-BZ) CMP, measured at current densities from 0.5 to 20 A g⁻¹. (f) Cycling stabilities of Py-FFC/CD-BZ CMP and Py-FFC/Poly(CD-BZ) CMP, measured at 10 A g⁻¹ over 2000 cycles.

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13

3.5. Electrochemical performance

We measured the electrochemical performance of our CMP materials in terms of their galvanostatic charge/discharge (GCD) and CV behavior, in 1 M aqueous KOH using a three-electrode system. Figures S6(a) and S6(b) display the CV curves of TPE-FFC—CMP and Py-FFC—CMP, respectively, recorded in the potential window from +0.1 to -0.90 V (vs. Hg/HgO) at various sweep rates from 5 to 200 mV s⁻¹. The CV curves of these two FFC-CMP samples had rectangular shapes featuring humps, indicating that their capacitive responses originated from both electric double-layer capacitance (EDLC) and pseudocapacitance, arising from the presence of electron-rich phenyl rings and electroactive ferrocene backbones, in which redox processes occurred during the charging and discharging processes [71,75-79]. The ferrocene backbones within TPE-FFC—CMP and Py-FFC-CMP underwent reversible one-electron redox processes, as revealed in the voltammetric profiles [Figures S6(a) and S6(b)]. The current density provided by Py-FFC-CMP was higher than that of TPE-FFC—CMP. The peak current densities increased upon increasing the sweep rate from 5 to 200 mV s^{-1} , while the shapes of the CV curves were retained, indicative of good electron transporting properties and facile kinetics [9,80,81]. Figures S6(c) and S6(d) present the GCD curves of TPE-FFC-CMP and Py-FFC-CMP, respectively, measured at various current densities from 0.5 to 20 A g^{-1} . The GCD curves of these two FFC-CMP samples were triangular with a slight bend, suggesting both EDLC and pseudocapacitive characteristics [82–84]. The GCD curves of these two samples displayed the typical features of pseudocapacitors, with high symmetry indicating good electrochemical reversibility and capacitance performance [85]. The discharging time of Py-FFC-CMP was longer than that of TPE-FFC—CMP [Figures S6(c) and S6(d)], indicating that the capacitance of the former was greater than that of the latter [Figure S6(e)]. Figure S6(e) presents the specific capacitances of TPE-FFC-CMP and Py-FFC-CMP, calculated from GCD curves using Eq. (S1). The capacitance of Py-FFC—CMP was relatively higher (5.07 F g^{-1}) than that of TPE-FFC—CMP (4.8 F g^{-1}) at a current density of 0.5 A g^{-1} , presumably because Py-FFC-CMP featured a higher surface area (50 m² ¹), higher pore volume (0.07 cm³ g⁻¹), better π -stacking, and a g^{-1} more ordered structure. We tested the durability of these two samples by cycling them over 2000 times at 10 A g^{-1} [Figure S6(f)]. TPE-FFC-CMP and Py-FFC-CMP both displayed good cycling stability [Figure S6(f)], with retentions of their original capacitances of 89.87% and 90.65%, respectively, after 2000 cycles. Figures S7(a) and S7(b) present the CV profiles of the TPE-FFC-CMP/CD-BZ and TPE-FFC-CMP/poly(CD-BZ) ICs, respectively. Their CV curves were rectangular, indicating that their capacitive responses originated mainly from EDLC (with minor pseudocapacitance), arising from the presence of electron-rich phenyl rings, electroactive ferrocene backbones, and heteroatoms (N and O atoms) [75–79]. In addition, the current density of the TPE-FFC-CMP/poly(CD-BZ) IC after ROP was higher than that of the TPE-FFC-CMP/CD-BZ IC. Figures S7(c) and S7(d) present the GCD curves of the TPE-FFC-CMP/CD-BZ and TPE-FFC-CMP/poly(CD-BZ) ICs, respectively. These GCD curves were triangular with a slight bend, suggesting the characteristics of both EDLC and pseudocapacity [78–81]. The discharging time of the TPE-FFC-CMP/poly(CD-BZ) IC was longer than that of the TPE-FFC-CMP/CD-BZ IC [Figures S7(c) and S7(d)], indicating that the capacitance of the former was greater than that of the latter [Figure S7(e)]. Figure S7(e) presents the specific capacitances of the TPE-FFC-CMP/CD-BZ and TPE-FFC-CMP/poly(CD-BZ) ICs. The capacitance of the TPE-FFC—CMP/poly(CD-BZ) IC after ROP (37.07 F g^{-1}) was much higher than that of the TPE-FFC-CMP/CD-BZ IC (7.53 F g^{-1}) at a current density of 0.5 A g^{-1} . We attribute this behavior to the high contents of N atoms and phenolic OH groups derived from the BZ units. These two TPE-FFC-CMP/CD-BZ and TPE-FFC-CMP/ poly(CD-BZ) ICs displayed excellent cycling stability [Figure S7(f)],

with retentions of their original capacitances of 94.3 and 96.03%, respectively, after 2000 cycles. Fig. 10(a) and 10(b) displays the CV curves of the Pv-FFC-CMP/CD-BZ and Pv-FFC-CMP/poly(CD-BZ) ICs, respectively. These curves were rectangular, indicating that the capacitive responses originated mainly from EDLC (with minor pseudocapacitance), arising from the presence of electron-rich phenyl rings, electroactive ferrocene backbones, and heteroatoms (N and O atoms) [75–79]. In addition, the current density of the Py-FFC-CMP/ poly(CD-BZ) IC after ROP was higher than that of the Py-FFC—CMP/ CD-BZ IC. Fig. 10(c) and 10(d) presents the GCD curves of the Py-FFC—CMP/CD-BZ and Py-FFC—CMP/poly(CD-BZ) ICs, respectively. These GCD curves were triangular shapes with a slight bend, suggesting the characteristics of both EDLC and pseudocapacity [79–81]. The discharging time of the Py-FFC-CMP/poly(CD-BZ) IC after ROP was longer than that of the Py-FFC-CMP/CD-BZ IC [Fig. 10(c) and 10 (d)], indicating that the capacitance of the former was larger than that of the latter [Fig. 10(e)]. Fig. 10(e) provides the specific capacitances of the Py-FFC-CMP/CD-BZ and Py-FFC-CMP/poly (CD-BZ) ICs. The capacitance of the Py-FFC-CMP/poly(CD-BZ) IC after ROP (46 F g^{-1}) was much higher than that of the Py-FFC-CMP/CD-BZ IC (10.15 F g⁻¹) at a current density of 0.5 A g⁻¹, presumably because of the high contents of N atoms and phenolic OH groups derived from the BZ units. These two Py-FFC-CMP/CD-BZ and Py-FFC-CMP/poly(CD-BZ) ICs displayed excellent cycling stability [Fig. 10(f)], with retentions of their original capacitances of 95.2% and 97.49%, respectively, after 2000 cycles. Table S2 compares the specific capacitances of these materials with those reported previously for other materials. Gratifyingly, the specific capacitance of the Py-FFC-CMP/poly(CD-BZ) IC after ROP was higher than those of previously reported covalent organic frameworks (COFs) and other CMPs [82,86-89].

4. Conclusions

Two FFC-CMP/CD-BZ CMP inclusion complexes were formed through supramolecular host-guest interactions between the benzoxazine-linked Cyclodextrin (CD-BZ) and ferrocene unit of FFC-CMP. The obtained FFC-CMP/CD-BZ ICs were examined by using FTIR, 2D NOESY ¹H NMR, solid-state ¹³C CP/MAS NMR, X-ray diffraction, and TGA analyses. After thermal curing polymerization (ROP), the Py-FFC/poly(CD-BZ) CMP ICs exhibited outstanding thermal stability (T_{d10} = 366 °C) with char yield of 62.1%, which could be attributed to the presence of intramolecular (OH-N) and intermolecular (OH-O) hydrogen bonding interactions. Interestingly, the Py-FFC-CMP/poly(CD-BZ) ICs revealed higher CO₂ uptake ability of 1.42 mmol g⁻¹and higher specific capacitance of 46 F g⁻¹at current density 0.5 A g⁻¹. In addition, we expect that these new porous materials could be used for other potential applications such as dyes adsorption, iodine capture, and photocatalytic H₂ evolution from water.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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12

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M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1–13

Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:10.1016/j.jtice.2021.10.010.

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13

M.M. Samy et al. / Journal of the Taiwan Institute of Chemical Engineers 00 (2021) 1-13

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