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Pyromellitic dianhydride-based polyimide anodes for sodium-ion batteries



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ABSTRACT

Organic redox-reactive polymers have garnered great attention as a promising alternative for conventional transition-metal compounds in sodium-ion batteries (NIBs) due to their low cost, structural flexibility and diverse structure. Among this class of materials, polyimides with high mechanical strength, excellent thermal stability and high density of electroactive functional groups have shown promise as low-cost electrode materials for NIBs. Herein, a simple hydrothermal method was used to synthesise pyromellitic dianhydride-based polyimides $[C_{16}H_6O_4N_2]_n$. The polyimides consisting of interconnected nanosheets with a microflower-like morphology were tested as an NIB anode. The polyimide electrode exhibited a stable discharge capacity of 125 mAh g⁻¹ at a current density of 25 mAg⁻¹ at the 100th cycle. At a high current density of 2 Ag^{-1} , the electrode delivered a discharge capacity of 43 mAh g⁻¹. The capacity contribution of this polyimide electrode mainly occurred below 1.5 V making it suitable as an organic NIB anode. The mechanism of sodiation and desodiation during discharge and charge was studied using Fourier transform infrared spectroscopy, in which this polyimide experienced two-step enolisation reaction with reversible insertion of two sodium ions during the redox electrochemical reaction.

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1. Introduction

Sodium-ion batteries (NIBs) have attracted a great attention as an alternative to lithium-ion batteries (LIBs) due to the limited natural occurrence of lithium [1-3], higher abundance and lower cost of sodium resources (the cost of Na₂CO₃ is only 3% of that of Li₂CO₃) [4,5] as well as the similar electrochemical properties of sodium and lithium (sodium is only 0.3 V more positive than lithium) [6].

Although many cathode materials suitable for NIBs have been identified, the choice for anode materials is however very limited [7–12]. The most commonly used anode in LIBs, graphite, is unfortunately not suitable for NIBs because it hardly allows sodium intercalate into the space between the graphene layers, and the intercalation is electrochemically irreversible [13]. The current leading anode candidate, hard carbon, can exhibit stable cycling

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performance but poor rate capability, due to its rigid tortuous nanometric structure [14]. Transition-metal inorganic compounds are extensively studied as anode materials for NIBs. However, due to their limited mineral resources, concerns over recyclability and environmental issues warrant cheap and alternative electrode materials. Also, large sodium ions inserting into the rigid lattices can cause multi-step phase transitions, which can lead to low utilisation of capacity and/or sluggish kinetics [15].

Organic materials stand out in the energy storage field as promising alternatives to inorganic compounds because of their flexibility, easy processability and structural and compositional diversity [16,17]. For example, polyimides with high mechanical strength, excellent thermal stability and high density of electroactive functional groups are promising redox-active electrode materials for rechargeable batteries [18,19]. Aromatic carbonylderivative polyimides, containing a dianhydride core as the aromatic backbone, have been commonly used as cathodes for NIBs [20–22]. However, recently it was found that when altering the aromatic core from perylene 3,4,9,10-tetracarboxylic dianhydride

(PTCDA) to 1,4,5,8-naphthalenetetracarboxylic dianhydride (NTCDA) and to pyromellitic dianhydride (PMDA), the average discharge voltage became progressively lower [23]. For the PMDA-based polyimide, the average discharge voltage was only approximately 1.73 V, indicating its potential suitability as an anode material for NIBs due to its relatively low voltage (<2 V vs. Na/Na⁺) [13].

Current research on organic electrodes is mainly concerned with the molecular design of different organic materials. In fact, structure and morphology also play an important role in the electrochemical performance of the organic electrodes. Recently it was found that disodium terephthalate with nanosheet-like morphology exhibited much improved electrochemical properties than the bulk spheroidal morphology [1]. However, such studies on the effects of structure and morphology of a certain organic material are very infrequently reported.

Here, we report the electrochemical behaviour of a PMDA-based polyimide as an organic anode in sodium-ion batteries. The effects of the structure and morphology on the electrochemical performance of polyimides were investigated by controlling the synthesis temperature and reagent concentrations. The mechanism of sodium storage in the organic electrode material indicates a two-step enolisation reaction with reversible insertion of two sodium ions. The nontoxic polyimide is cheap and easy to synthesise, possibly fulfilling the needs for large-scale battery applications.

2. Experimental section

2.1. Synthesis of pyromellitic dianhydride-based polyimides

2.5mmol p-phenylenediamine (PPD) was dissolved in 30.0 mL N,N-dimethylformamide (DMF) followed by adding equimolar PMDA under stirring at room temperature. The mixture was then transferred into a Teflon lined autoclave and hydrothermally treated at 150, 180 or 210 °C for 10 h. The precipitate was separated by centrifugation, washed with DMF and ethanol several times, and dried under vacuum at 80 °C for 24 h. The as-synthesised samples are defined as PI-150-2.5, PI-180-2.5 and PI-210-2.5, where PI stands for polyimide, 150, 180 and 210 indicate the synthesis temperature, and 2.5 is the concentration of the PPD and PMDA added in millimoles per litre. The as-synthesised samples PI-150-2.5, PI-180-2.5 and PI-210-2.5 were heated at 100, 200 and 300 $^{\circ}\text{C}$ for 1 h under vacuum, respectively for post curing (PC). Samples thus obtained are named by adding the prefix PC before a sample, for example, PC-PI-150-2.5. Other polyimide samples were also synthesised with different concentrations of PPD and PMDA using the same method as described above under the hydrothermal treatment temperature of 210 °C. Samples thus obtained are designated as PI-210-1.25, PI-210-2.5 and PI-210-5.

2.2. Characterisation

Fourier transform infrared (FTIR) spectra of powder samples were recorded on an attenuated total reflectance FTIR (ATR-FTIR) spectrometer (Nicolet 5700, Thermo Electron). 13 C magic angle spinning nuclear magnetic resonance (MAS NMR) spectra were acquired on a Bruker Avance III spectrometer at a 13 C Larmor frequency of 75.468 MHz. All the experiments were carried out with a Bruker 4 mm MAS probe and zirconia rotor spinning at 5 kHz. A CPMAS pulse sequence was used with 100 kHz decoupling and a 3 s relaxation delay. X-ray diffraction (XRD) measurements were performed on an X-ray diffractometer (Bruker D8 Advance, Bruker) with Cu K α radiation (λ = 1.54056 Å). The morphology of the samples was characterised using a field emission scanning electron microscope (FESEM) (ISM-7100F, IEOL). Nitrogen adsorption-

desorption measurements were carried out on a surface area and pore size analyser (TriStar II 3020, Micromeritics) at 77 K. The specific surface area ($S_{\rm BET}$) and pore size distribution were analysed by the Brunauer-Emmett-Teller (BET) and Barrett-Joyner-Halenda (BJH) methods, respectively. The total pore volume ($V_{\rm t}$) was estimated at the relative pressure of 0.99, micropore volume ($V_{\rm mic}$) was obtained using the t-plot method, and the mesopore volume ($V_{\rm meso}$) was calculated from the difference between $V_{\rm t}$ and $V_{\rm mic}$.

2.3. Electrochemical measurements

Polyimide, carbon black, and poly(vinylidene difluoride) (PVDF) were dispersed in N-methyl pyrrolidone (NMP) at a mass ratio of 6:3:1 to form a slurry, which was cast onto a copper foil current collector using a doctor blade. The electrode was dried at $60\,^{\circ}$ C under vacuum for 12 h. A coin cell was assembled in an Argon-filled glovebox using the polyimide electrode as the working electrode, pure sodium foil as the counter electrode, glass fibre as the separator and $1\,\text{mol}\,L^{-1}\,\text{NaClO}_4$ in equal volume of ethylene carbonate (EC) and propylene carbonate (PC) mixed with $0.3\,\text{wt}\,\%$ fluoroethylene carbonate (FEC) as the electrolyte.

Galvanostatic discharge and charge cycling were conducted on a battery tester (CT3008, Neware) at room temperature. Cyclic voltammetry (CV) measurements were carried out on an electrochemical workstation (CHI 660D, Chen Hua Instrument). To study the mechanism of charging, the charged or discharged electrodes were obtained from dissembled coin cells composed of polyimide as the positive electrode and sodium metal as the negative electrode, washed with anhydrous dimethyl carbonate (DMC) and then dried naturally in the glove box for characterisations. The theoretical capacity was calculated based on a two-electron transfer redox process for each formula unit using the following equation: [16]

$$\begin{split} C\Big(\text{mAh g}^{-1}\Big) &= \frac{n \times F\big(\text{C mol}^{-1}\big)}{M_{W}(\text{g mol}^{-1}\big)} = \frac{n \times 96485(\text{C})}{M_{W}(\text{g})} \\ &= \frac{n \times 96485(\text{As})}{M_{W}(\text{g})} \\ &= \frac{n \times 96485 \times 1000/3600(\text{mAh})}{M_{W}(\text{g})} \\ &= \frac{26801 \times n}{M_{W}} \Big(\text{mAh g}^{-1}\Big) \end{split} \tag{1}$$

C, n, F and M_w are the theoretical specific capacity, the transferred electron number in each formula unit (here n=2), the Faraday constant and the molecular weight of the formula unit, respectively.

3. Results and discussion

Fig. 1a shows the FTIR spectra of the as-synthesised samples PI-150-2.5, PI-180-2.5 and PI-210-2.5 prepared at different hydrothermal temperatures. All characteristic absorbance signals of the imide group can be seen. The peaks at 1782, 1720 and 723 cm⁻¹ can be assigned to the asymmetric stretching vibration (v_{as}), symmetric stretching vibration (v_{as}) and bending vibration (δ) of the imide group C=O, respectively. The peak at 1380 cm⁻¹ can be assigned to the stretching vibration of the imide C-N group. The FTIR results are in good agreement with previously reported data [23], demonstrating the successful synthesis of the polyimide.

Traditionally, a post-cure step is necessary to ensure full ring closure in the synthesis of polyimides [24–26]. However, the hydrothermal synthesis method in this work resulted in polyimide

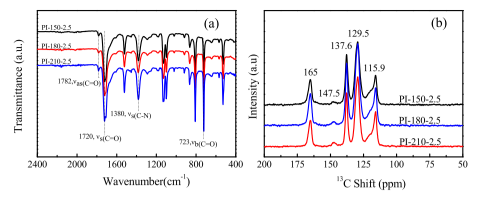


Fig. 1. (a) FTIR spectra and (b) ¹³C NMR spectra of PI-150-2.5, PI-180-2.5 and PI-210-2.5.

without post-curing, as was proven by the FTIR data shown in Fig. S1. It can be seen that the FTIR spectra of post-cured samples are identical to that of the as-synthesised samples, confirming complete imidisation of the polyimide obtained from the hydrothermal synthesis method. The successful synthesis of polyimides was further confirmed by the ¹³C MAS NMR results shown in Fig. 1b. The peak at 165 ppm corresponds to the imide carbonyl carbon, the small peak at 147.5 ppm to the aromatic carbon connected to terminal amino group, the peak at 137.6 ppm to the non-protonated aromatic carbon of the bensimide group, and the peak at 129.5 ppm to protonated aromatic bensimide carbon, respectively. The peak at 115.9 ppm can be assigned to the N-substituted aromatic carbon and the protonated aromatic carbon of the N-substituted phenyl group. The NMR spectra match well with previously published results [27].

Fig. 2a shows the XRD patterns of the polyimide samples synthesised at different temperatures. It can be seen that the crystal-linity was increased at higher synthesis temperature as seen from the increased peak intensities. This is because higher improved hydrothermal synthesis temperature increases the system pressure, favouring crystallisation of the polyimides [28]. The average

crystallite dimension or size of the coherent crystalline domain and lattice imperfections can be determined using the Williamson and Hall equation: [29,30]

$$\beta \cos\theta = 4\varepsilon \sin\theta + 0.9\lambda/D$$

where β is the corrected integral width, ε is the microstrain ($\Delta d/d$, d: lattice-plane spacing), and D is the average size of crystallite domain. The above equation was applied to the six relatively strong peaks at approximately 14.2° , 19.4° , 21.2° , 27.7° , 30.5° and 36.2° two theta. By plotting $\beta\cos\theta$ against $\sin\theta$, an intercept $(0.9\lambda/D)$ and a slope (4ε) that correlate with the average size of crystallite domain and microstrain, respectively, were obtained. As can be seen from Fig. 2b-d, the plots gave a similar intercept, but very different slopes. The plot of the sample synthesised at 210° C showed a smaller slope than the sample synthesised at 150° C, indicating a smaller microstrain.

[28] Fig. 3 shows the FESEM images of samples PI-210-1.25, PI-210-2.5 and PI-210-5. All samples are in the form of micrometresized particles consisting of nanosheets as the primary particles. In addition, the primary particles became smaller as the reagent

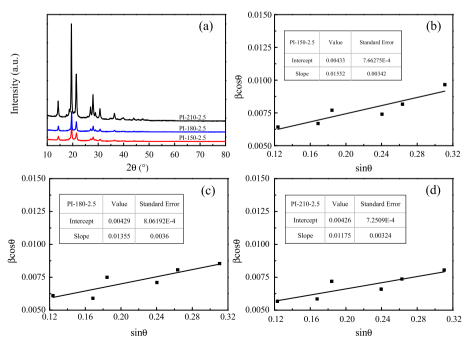


Fig. 2. (a) XRD patterns and (b-d) plots of the Williamson and Hall equation of as-synthesised samples (b) Pl-150-2.5, (c) Pl-180-2.5 and (d) Pl-210-2.5.

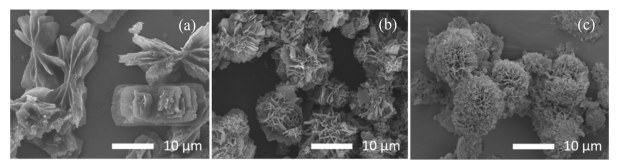


Fig. 3. FESEM images of (a) PI-210-1.25, (b) PI-210-2.5 and (c) PI-210-5.

concentration increased, which is ascribed to increased nucleation density during the polymerisation reaction [31]. The PI-210-1.25 exhibited two-dimensional (2D) lamellar morphology (Fig. 3a), and the PI-210-2.5 and PI-210-5 synthesised at higher concentrations presented three-dimensional (3D) interconnected microflower-like morphology composed from 2D nanosheets (Fig. 3b and c). The formation of the 3D microflowers is a result of the growth mechanism of spherulites [32,33]. At low concentrations, the 2D lamellar structures were formed by polymer chains orienting perpendicularly to the nanosheet plane. At higher concentrations, fully developed 3D microflower-like structures were formed by the self-assembly of additional nanosheets via unidirectional growth and low angle branching. In energy storage systems, 3D structures have advantages over 2D structures, such as high contact area, novel physical properties, and structural stability during cycling [34-36]. Here, the 2D polyimide as the primary building block can be exposed as electroactive sites to sodium ions. while the 3D polyimide can present a shorter pathway for diffusion of sodium ions due to the interconnected nanosheet structure.

Fig. S2 shows the nitrogen adsorption-desorption isotherms and the corresponding pore size distributions of PI-210-1.25, PI-210-2.5 and PI-210-5 synthesised using different concentrations of the reagents. As shown in Fig. S2a, all of the samples presented a type IV isotherm with a hysteresis loop, indicating the existence of the mesopores. The pore size distributions in Fig. S2b also confirmed that the samples had a mesoporous structure, and the sample PI-210-2.5 occupied a very wide mesopore distribution ranging from 4.5 nm to 40 nm. It can be seen from Table S1 that mesopores comprised a major proportion of the pore structure in all samples. Although PI-210-5 had the highest specific surface area, PI-210-2.5 occupied the largest mesopore volume of 0.2552 cm³ g⁻¹. A large surface area in mesoporous materials can enhance the contact area between electrode materials and the electrolyte. A mesoporous structure can not only promote the infiltration of electrolyte to shorten the diffusion pathways for sodium ions, but also provide a large effective working interface for electrochemical reactions to decrease the electrochemical polarisation and improve the capacity utilisation and rate capability of the electrode [37].

Fig. 4a shows the electrochemical impedance spectra of PI-150-2.5, PI-180-2.5 and PI-210-2.5. It can be seen that polyimides synthesised at 210 °C show smaller diameter of the semicircle in the medium frequency region than that of the other samples, indicating lower charge transfer resistance. The lower microstrain of PI-210-2.5 may enhance the ion transport in the material [38,39], leading to improved charge transfer at the interface between the electrode and electrolyte [40]. Fig. 4b shows the cycling performance of the PI-150-2.5, PI-180-2.5 and PI-210-2.5 at a current density of 25 mA g⁻¹ between 0.005 and 3 V. As expected, the sample PI-210-2.5 achieved higher capacity and better cycling performance than PI-150-2.5 and PI-180-2.5. The superior cycling

performance may be associated to the lower microstrain, which can cause less irreversible capacity loss [41,42]. It should be noted that all the polyimide electrodes delivered an extremely high initial discharge capacity, which can be ascribed to the decomposition of electrolyte and the formation of a solid electrolyte interphase (SEI) at the low potential range. As is shown in Fig. S3, the formation and growth of SEI layer were evident from the increased size of the Nyquist semicircle of the cycled electrode in three-electrode cells in the initial few cycles [43]. After the 5th cycle, the resistance almost stabilised.

Fig. 4c shows the cycling performance of PI-210-1.25, PI-210-2.5 and PI-210-5 at a current density of 25 mA g^{-1} between 0.005 and 3 V. It can be seen that PI-210-2.5 delivered the highest capacity of 125 mAh g⁻¹ at the 100th cycle, corresponding to approximately 68% of the theoretical capacity of 185 mAh g⁻¹ according to Equation (1). The coulombic efficiency of PI-210-2.5 reached 80% at the 10th cycle and remained nearly 95% at the 100th cycle as shown in Fig. S4. The weight ratio of carbon black for the preparation of organic electrodes is usually 30 wt% or even higher [1,6,7,15,20-22]. Considering 30 wt% of carbon black used during the polyimide electrode preparation, cycling performance of carbon black was also investigated at the same condition as shown in Fig. S5, which indicates that the capacity contribution $(\sim 11 \text{ mAh g}^{-1})$ from carbon black is negligible. Fig. 4d presents the rate performance of PI-210-1.25, PI-210-2.5 and PI-210-5 at different current densities. PI-210-2.5 achieved the highest capacity under the same current density, showing the best rate performance. Initial discharge capacities of 124, 90, 71, 59, 50, 43 mAh $\rm g^{-1}$ were achieved at current densities of 50, 100, 200, 500, 1000, $2000 \, \text{mA g}^{-1}$, respectively. It should be pointed out that at a current density of 2000 mA g⁻¹, PI-210-2.5 delivered a discharge capacity of $43 \, \text{mAh} \, \text{g}^{-1}$. In addition, a reversible discharge capacity of $123 \, \text{mAh} \, \text{g}^{-1}$ can be recovered after 5 additional cycles when the current density was reduced to $25 \, \text{mA g}^{-1}$, demonstrating a good rate capability. The good rate performance of PI-210-2.5 can be explained by its open macroscopic structure exposing the electrode to the electrolyte, and mesoporous structure for enhanced electrolyte transport as well as effective interconnected structure.

Selected discharge and charge profiles of PI-210-2.5 at a current density of 25 mA g⁻¹ are shown in Fig. 4e. It can be seen that the capacity stabilised after the 10th cycle. The discharge and charge profiles from the second cycle appeared very similar with sloping curves due to fast redox reactions [22,44,45]. Additionally, it can be found that the capacity was mainly from a discharge voltage below 1.5 V, indicating the suitability of this polyimide electrode as an anode for NIBs. Fig. 4f displays the CV curves of the polyimide electrodes between 0.005 and 3 V vs. Na/Na⁺ at the scan rate of 0.2 mV s⁻¹. During the first cathodic scan, a strong broadened and split reduction peak between 0.5 and 1.5 V was observed. This peak became diminished in the next cycle. As a result, this peak may be

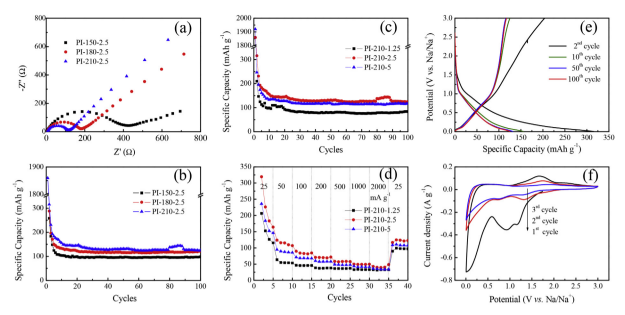


Fig. 4. (a) Nyquist plots of PI-150-2.5, PI-180-2.5 and PI-210-2.5 for the fresh cells, (b) cycling performance for PI-150-2.5, PI-180-2.5 and PI-210-2.5 synthesised under different temperatures, (c) cycling performance for PI-210-1.25, PI-210-2.5 and PI-210-5 synthesised at different reagent concentrations, (d) rate performance for PI-210-1.25, PI-210-2.5 and PI-210-5 from the second discharge capacity, (e) selected discharge and charge profiles for PI-210-2.5, and (f) CV curves at a scan rate of 0.2 mV s⁻¹ for PI-210-2.5.

due to the formation of a SEI layer along with the sodiation of the polyimide. During the reverse anodic scan, there was a broad peak between 1 and 2 V, corresponding to the desodiation of the sodiated polyimide. In the next scan, two cathodic peaks appeared at potentials of 0.75 and 1.25 V, likely corresponding to the sodiation of the carbonyls into radical anions and dianions [22]. These two close redox peaks with only a slightly positive shift can be also seen during subsequent scans, indicating a multiple and fast electron transfer reaction as well as good reversibility. The CV results were consistent with the discharge and charge observations.

Fig. 5 shows typical FTIR spectra of the PI-210-2.5 electrodes discharged to 0.005 V and recharged to 3 V. As discussed above, the signals at 1720 and 723 cm⁻¹ were attributed to v_s (imide C=O) and δ (imide C=O) of the active material polyimide. It was found that the peak of the δ (imide C=O) decreased in intensity when the electrode was discharged to 0.005 V, indicating reduction of the carbonyl groups. In addition, the peak due to v_s (imide C=O)

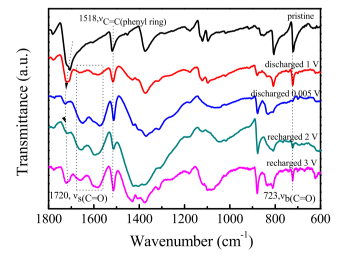


Fig. 5. FTIR spectra of the polyimide electrodes at different discharged and charged states.

gradually shifted from 1720 cm⁻¹ to 1726 cm⁻¹, due to formation of sodium enolate O-C-O-Na [22]. When the electrode was recharged to 3 V, the intensity of the δ (imide C=0) peak gradually recovered, demonstrating the recovery of the carbonyl groups due to desodiation. After discharging to 1 V where the CV curve shows a huge reduction peak, two new peaks at 1655 and 1588 cm⁻¹ appeared and remained after recharging, which may be due to the SEI layer [46]. Additionally, during the discharge and charge processes, the peak at 1518 cm⁻¹ attributed to the stretching vibration of C=C in phenyl rings was almost unchanged, suggesting that the unsaturated double carbon bond does not actively participate in sodium-ion storage. The FTIR results also match well with the CV observation (Fig. 4f) that there was sodiation of the carbonyls into radical anion and dianion during the redox reaction. When combining analysis of the CV results and the reversible dischargecharge capacities (125 mAh g^{-1} at the 100th cycle), the redox mechanism of the PMDA-based polyimide can be illustrated as a two-step enolisation reaction with reversible insertion of two sodium ions as shown in Scheme 1.

4. Conclusion

In this study, a one-pot hydrothermal method was used to prepare polyimides with controllable hierarchical mesoporous structure constructed from nanosheets. A microflower-like polyimide with high degree of crystallinity and large mesopore volume can be obtained by adjusting synthesis temperatures and reagent concentrations to achieve higher capacity and superior performance. As an anode material in sodium-ion batteries (NIBs), the PMDA-based polyimide experienced a two-step enolisation reaction with reversible insertion of two sodium ions during the redox electrochemical reaction. This polyimide electrode can be considered as promising for organic sodium-ion batteries with its capacity mainly contributed below 1.5 V and a high reversible capacity of 125 mAh g⁻¹ at the 100th cycle. This nontoxic polymer is inexpensive and simple to synthesise, and can therefore fulfil the needs for large-scale battery applications. Overall, this study provides a new method to improve the electrochemical performance of organic electrodes via adjusting the structure and morphology.

Scheme 1. Possible discharge and charge mechanism for the PMDA-based polyimide in a NIB.

Conflicts of interest

None.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.electacta.2018.01.208.

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