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1. Introduction

Electrochemical supercapacitors (ESs), especially electrochemical double-layer capacitors (EDLCs) with high power densities, high coulombic efficiencies (CEs), and long-term cycling lives, are considered to be one of the best potential high-power energy storage devices.¹⁻⁴ As is well established, EDLCs generally store electrochemical energy based on the fast accumulation of charge in the double layer formed at the sur-/ interfaces of the porous carbon electrodes with large specific surface area (SSA).^{1,3,4} Carbon materials, such as activated carbon,⁵ multi-modal porous carbon,⁶ hollow carbon spheres,⁷

Sustainable rose multiflora derived nitrogen/ oxygen-enriched micro-/mesoporous carbon as a low-cost competitive electrode towards highperformance electrochemical supercapacitors†

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Cost-efficient carbonaceous materials have been utilized extensively for advanced electrochemical supercapacitors. However, modest gravimetric/volumetric capacitances are the insuperable bottleneck in their practical applications. Herein, we develop a simple yet scalable method to fabricate low-cost micro-/mesoporous N/O-enriched carbon (NOC-K) by using natural rose multiflora as a precursor with KOH activation. The biomass-derived NOC-K is endowed with a large surface area of \sim 1646.7 m² g⁻¹, micro-/mesoporosity with \sim 61.3% microporosity, high surface wettability, and a high content of N (\sim 1.2 at%)/O (~26.7 at%) species. When evaluated as an electroactive material for supercapacitors, the NOC-K electrode (5 mg cm⁻²) yields large gravimetric/volumetric specific capacitances of \sim 340.0 F g⁻¹ (\sim 238.0 F cm⁻³) at 0.5 A g⁻¹, and even \sim 200.0 F g⁻¹ (\sim 140.0 F cm⁻³) at 5.0 A g⁻¹, a low capacitance decay of \sim 4.2% after 8200 consecutive cycles, and a striking specific energy of \sim 8.3 W h kg⁻¹ in aqueous KOH electrolyte, benefiting from its intrinsic structural and compositional superiorities. Moreover, a remarkable specific energy of \sim 52.6 W h kg⁻¹ and \sim 96.6% capacitance retention over 6500 cycles for the NOC-K based symmetric cell are obtained with the organic electrolyte. More promisingly, the competitive NOC-K demonstrates enormous potential towards advanced supercapacitors both with aqueous and organic electrolytes as a sustainable electrode candidate. PAPER

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modified activated carbon aerogel,⁸ carbon nanotubes⁹ and so on, have been comprehensively explored as the electrode materials for ESs. Unfortunately, the modest gravimetric/ volumetric capacitances and low specific energy (SE) of the EDLCs still inherently hinder their practical widespread applications.2,3,10 Thus, numerous efforts have been made to design and explore various carbon materials with optimum pore structures and surface compositions/properties, with the aim to enhance their electrochemical behaviors particularly at high rates.

In general, the superior specific capacitances (SCs) of carbon materials are mainly determined by their large SSA, hierarchical micro-/mesopores, and high-surface heteroatom content (such as O, N, B, P, $etc.$).^{3,4,10-18} To begin with, the high SSA can guarantee massive electroactive sites for fast and efficient charge accommodation.11–²⁰ Besides, bi-modal micro-/ mesopores greatly favor for convenient ion diffusion into the electrode bulk by providing ion highways, thus ensuring highpower property, and additionally contribute to large electroactive SSA meanwhile.^{3,18,20} In addition, heteroatom doping can partially form N- and O-based surface functional groups, which improves the surface wettability of carbon electrodes for effective charge storage.^{11,21-23} Notedly, the polar functional groups

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[†] Electronic supplementary information (ESI) available: Electronic supporting information corresponding XRD pattern, charge–discharge plots, electrochemical comparisons with other carbon-based electrodes, EIS simulated results, and cycling performance of the symmetric devices within the potential range from 0.0 to 2.5 or 3.0 V. See DOI: 10.1039/c8ra00858b

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upon the carbon surface exhibit benign affinity to solvents with the same polarity, especially in the ubiquitous aqueous systems.²⁴ Of particular note, the N doping also can change the intrinsic conductivity of carbon materials themselves by adjusting the giving/receiving capacity of the electrons in the C atom, which makes them even better electronic conductivity than their phase-pure counterparts, thanks to the high electronegativity, small atomic size, and additional free electrons of the N itself.^{3,13,20,21} Therefore, the nitrogen species have attracted the huge interest of many researchers around the word. It is noted that two strategies are generally explored to incorporate the elemental N into the porous carbon materials: posttreatment of porous carbon with extra N-based sources (such as, ammonia, urea, amoine, and so on) and in situ doping by directly pyrolyzing various N-containing precursors. Commonly, the former only introduces N-based functional groups on the surface of the carbon materials, while the latter can easily realize the homogeneous heteroatom doping into the bulk with controlled chemistry, rather than just on the surface of carbon electrodes.10 **Exception** Access Article Continuous Contract Continuous Control Cont

Recently, biomass materials, different from those artificial materials, have drawn enormous attentions, owing to their connate advantages including richness in nature, low cost, renewability and environmental friendlyness. Carbon materials derived from various biomasses not only possess unique special morphologies, but also can be in situ doped with the heteroatoms originally existing in these biomass precursors. Consequently, this type of carbon materials inherently owns higher market potential due to the lower cost than commercial carbon nanotubes and graphene.²⁵ Nowadays, lots of pioneering works have been reported by using the human hair, 14 dead leaves, 16 shiitake,¹⁸ albumen,²⁰ coconut shell,²⁶ corn stover,²⁷ banana peel,²⁸ cashmere,⁶ and others as the precursors.

With aforementioned considerations in mind, herein, we proposed that a facile yet efficient synthesis of N/O cofunctionalized carbons with bi-modal micro-/mesopores by using rose multiflora as the precursor. After KOH activation process at 800 °C, the as-resulted carbon electrode was obtained with high-content nitrogen (\sim 1.2 at%) and oxygen (\sim 26.7 at%) functionally doping, micro-/meso-porous channels with microporosity of \sim 61.3%, large SSA (\sim 1646.7 m² g⁻¹) and appealing surface wettability. When evaluated as an electrode material for ESs, the KOH-activated nitrogen/oxygen-enriched carbon (denoted as NOC-K) electrode exhibited remarkable electrochemical capacitances both in aqueous KOH solution and organic tetraethylammonium tetrafluoroborate/propylene carbonate (TEABF₄/PC) electrolyte, benefiting from its unique structural and compositional superiorities.

2. Experimental

$2.1.1.$ Synthesis of the NOC-K α

The waste rose multiflora (Anhui province, P. R. China) was thoroughly washed with de-ionized water and dried at 60 $^{\circ}$ C in an electric oven. The cleaned rose multiflora flower was first pyrolyzed in a tubular furnace at a temperature of 500 $^{\circ}$ C for 2 h with a heating rate of 5 °C min⁻¹ in a high-purity N₂

atmosphere. During following KOH-activated process, the asobtained black powder was further mixed with the activation agent of KOH with a KOH/pre-pyrolyzed carbon ratio of 3 : 1 (w/w). Afterwards, the mixture was activated under N_2 flow at 800 °C for 2 h in a tubular furnace with a heating rate of $3 °C min⁻¹$. After the completion of the activation, the sample was allowed to cool to room temperature. After purified in diluent HCl solution, rinsed with de-ionized water until neutral, and dried at 60 \degree C, the resulting black NOC-K product was obtained. For comparison, the carbon prepared with the absence of KOH was designed as NOC, while other parameters were kept the same as those for the NOC-K.

Crystalline phases of the samples are examined by powder X-ray diffraction (XRD) (UltimaIV, Rigaku, Japan) using Cu Ka source over a 2θ range of 10–80°. Morphologies and structures of the resulting samples were examined by field-emission scanning electron microscopy (FESEM) (JEOL-6300F, 15 kV), transmission electron microscopy (TEM), and high-resolution TEM (HRTEM) (FEI, TECANI-20). N₂ adsorption/desorption was determined by Brunauer–Emmett–Teller (BET) measurement by using an Autosorb-iQ surface area analyzer. X-ray photoelectron spectroscopy (XPS) was measured on a Thermo ESCALAB 250 Xi X-ray photoelectron spectrometer with Al ka excitation source $(hv = 1486.6 \text{ eV})$. The Raman data was recorded by laser Raman (T6400, Jobionyzon Corp., France). The contact angle measurement was performed with Contact Angle System OCA.

Cyclic voltammogram (CV) at different scanning rates, chronopotentiometry (CP) tests at various current densities and electrochemical impedance spectroscopy (EIS) were carried out on an IVIUM electrochemical workstation (The Netherlands). A three-electrode electrochemical configuration, where a Pt plate (1 cm^2) and saturated calomel electrode (SCE) were used as the counter electrode and reference electrode, was applied for electrochemical evaluation of the as-obtained carbons in 6 M KOH solution at room temperature. The working electrodes were prepared by loading a slurry containing NOC-K or NOC, acetylene black and polytetrafluoroethylene in a weight ratio of 5 : 2 : 1. The mass loading of electroactive material on each electrode is 5 mg for three-electrode evaluation. And twoelectrode symmetric cell with two identical electrodes were assembled into coin cells (CR2032) with 6 M KOH as the electrolyte.

In addition, a symmetric cell was also electrochemically characterized in 1 M TEABF₄/PC electrolyte. The electrode was prepared by mixing NOC-K or NOC, acetylene black, polyvinylidene difluoride (PVDF) binder $(8:1:1, w/w/w)$ dispersed in the N-methylpyrrolidinone (NMP). The slurries coated on the Al and steel foils were employed as positive and negative electrodes. After dried in vacuum at 120 $^{\circ}$ C for 24 h, the electrodes were separated with glass fiber (GF/D, Whatman) in CR2032 coin-type cells for electrochemical measurement in 1 M TEABF₄/ PC electrolyte.

For the three-electrode systems or symmetric ESs, the mass capacitance $(F g^{-1})$ and volumetric specific capacitance (VSC) $(F cm^{-3})$ derived from galvanostatic tests can be calculated by the following equations:

$$
SC = \frac{It}{\Delta V} \tag{1}
$$

$$
VSC = \frac{1}{\rho} SC = \frac{1}{V_{\text{total}} + \frac{1}{\rho_{\text{carbon}}}} SC
$$
 (2)

where the I is the discharge current (A g^{-1}), Δt is the discharge time (s), ΔV is the potential window (V), ρ is the density of the NOC-K or NOC, V_{total} is the total pore volume of the NOC-K or NOC, and $\rho_{\rm carbon}$ is the true density of the bulk carbon $(2 \text{ g cm}^{-3}).$

For the symmetric cell, the SE and specific power (SP) were calculated as follows:

$$
SE = \frac{1}{2} SC\Delta V^2
$$
 (3)

$$
SP = \frac{SE}{t}
$$
 (4)

where SC, ΔV and t are the mass capacitance based on two identical electrodes, discharge time, and potential window of the symmetric device, respectively.

3. Results and discussion

3.1. Physicochemical characterizations

As schematically illustrated in Fig. 1, two procedures, involving pre-carbonization of the rose multiflora with a short flowering

phase of just one month, and subsequent KOH-activation process, were carried out for the efficient and scalable fabrication of the NOC-K material. Compared with the common template synthesis strategies, the KOH-activation of the biomass, as a simple and effective method, is particularly suitable for practical industrial production. As identified by the XRD measurement (Fig. S1, ESI†), two broadened diffraction peaks appearing at $2\theta = 24$ and 43° are related to the (002) and (100) facets of the typical carbon material.

Fig. 2a shows typical FESEM image of the hierarchical NOC-K product, and the sample presents loose morphology with interconnected hierarchical pores. TEM characterizations (Fig. 2b-d) were also carried out to further examine the specific microstructures of the sample more clearly. As shown in Fig. 2b, the discernable mesoporous structure can be presented for the NOC-K. Further high-magnification TEM (Fig. 2c) and HRTEM (Fig. 2d) observations show staggered and interpenetrated micropores dominating in the NOC-K sample along with several scattered mesopores, which would contribute to even more electroactive sites, and provide convenient and continuous channel meanwhile for the smooth entry and migration of electrolyte ions. As for the KOH activation, the expected products including $CO₂$ and $H₂O$ formed in the activation process, and the removal of intercalated metallic K and K-based compounds by following washing both can result in the generation of well-developed pore network in the nano-scale and/or sub-nanometer range observed for the NOC-K specimen. Corresponding energy dispersive spectrometry (EDS) elementary mapping analysis of the NOC-K sample, as evidenced in Fig. 2e-h, visually confirms the homogeneous coexistence and uniform distribution of the C, N and O species in the whole NOC-K sample. Paper

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Fig. 1 Schematic illustration of the fabricating procedure for the NOC-K with the rose multiflora as a starting material.

Fig. 2 FESEM (a), TEM ((b) and (c)) and HRTEM (d) images of the NOC-K product, corresponding selected TEM area (e) for elemental ((f), C; (g), N and (h), O) mapping images.

To better understand the unique bi-modal porous structure and SSA parameters, we measured N_2 sorption isotherms of the NOC-K and NOC samples, as comparatively described in Fig. 3. Obviously, typical adsorption–desorption isotherms of the two, as plotted in Fig. 3a, are close to the combination form of the type I and IV according to the classic International Union of Pure and Applied Chemistry (IUPAC) definition. This distinct feature suggests the co-existence of the micro-/mesopores both in the NOC-K and NOC products, which can be well supported by their mesopore size distribution data (Fig. 3b) derived from

the adsorption branches by the Barrett–Joyner–Halenda method, and the micropore size distribution (the inset in Fig. 3b) estimated by the Horvanth–Kawazone method. According to the calculation from the adsorption branch, the BET SSA of the NOC-K is \sim 1646.7 m² g⁻¹, which is approximately two times that of the NOC (\sim 826.6 m² g⁻¹). And the total pore volume are estimated as \sim 0.39 and \sim 0.84 cm³ g⁻¹ for the NOC and NOC-K, respectively, as determined from the N_2 adsorption amount at $P/P_0 = 0.97$. According to the ratio of micropore volume to total pore volume, $\sim 62.4\%$ and $\sim 61.3\%$ of

Fig. 3 N₂ adsorption–desorption isotherms (a), mesopore size distribution (b) and micropore size distribution (the inset in panel b) for the NOC and NOC-K samples.

the porosity for the NOC and NOC-K come from the micropores. As a result, the average pore sizes of the NOC and NOC-K can be derived as \sim 2.1 and \sim 2.3 nm, respectively. As noted, the NOC-K product obtains high SSA and well-developed porosity with even higher proportion mesoporosity, which are of vital significance to enhance the electrolyte accessibility to the microporous surface for efficient charge storage.

Fig. 4a demonstrates the Raman spectra of the NOC and NOC-K. Typically, both of the samples mainly show two distinct peaks located at around 1350 cm⁻¹ (D-band) and 1590 cm⁻¹ (Gband), respectively. As is known to all, the D-band corresponds to defect sites or disordered sp²-hybridized carbon atoms of graphite, and the G-band is related to the phonon mode inplane vibration of sp²-bonded carbon atoms.²⁹ The I_G/I_D values of the NOC and NOC-K are \sim 0.85 and \sim 0.65, respectively. The reduced I_G/I_D value of the NOC-K sample suggests more defective structure and lower graphitization owing to the subsequent high-temperature KOH activation.

Fig. 4b–f collectively summarize the high-resolution elemental XPS spectra of the NOC and NOC-K, and corresponding profiles fitted by using the Gaussian fitting method. Clearly, three elements including elemental carbon, oxygen and nitrogen are all discernable in the two products. Typically, the C

1s spectra for the two, as evidenced in Fig. 4b, can be deconvoluted into following five peaks including $C=C$ (C-I, \sim 284.6 eV), C–C (C-II, \sim 285.0 eV), C–N/C–O (C-III, \sim 285.6 eV), C=O (C-IV, \sim 286.6), and O–C=O (C-V, \sim 288.4) bonds.³⁰ Fig. 4c shows the deconvolutions of the O 1s spectra to identify the surface oxygen-containing functional groups upon the NOC and NOC-K products. Specifically, the three peaks at binding energies of \sim 532.6, \sim 533.8 and \sim 535.1 eV correspond to the carbonyl $(C=O)$ $(O-I)$, hydroxyl $(C-O)$ $(O-II)$, and carbonxyl $(O=$ C–O) (O-III), respectively.³¹ The N 1s spectra of the samples (Fig. 4d) can be fitted well into four peaks centered at \sim 397.8, \sim 399.5, \sim 400.6 and \sim 402.7 eV, respectively, which are assigned to four types of nitrogen functional group, namely pyridinic (N-I), pyrrolic and/or pyridonic (N-II), quaternary nitrogen (N-III) and oxidized N-oxide species (N-IV).^{3,8,14,15} Compared with the NOC, total elemental nitrogen and carbon contents of the NOC-K sample are largely increased up to \sim 1.2 at% and \sim 72.1 at% from \sim 0.5 at% and \sim 66.2 at%, respectively, while the percentage of elemental oxygen is reduced from \sim 33.3 at% to \sim 26.7 at%. One especially note that the N-III and N-IV in the NOC-K (Fig. 4f), favoring for the rapid electronic transfer through the carbon electrodes,^{32,33} are estimated as \sim 0.36 and \sim 0.26 at%, higher than those for the NOC (N-III, ~ 0.21 at%; N-IV, ~ 0.12 at%) (Fig. 3e). Paper

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Fig. 4 Raman spectra (a) and high-resolution elemental ((b), C 1s; (c), O 1s; (d), N 1s) XPS spectra and fitted data for the resulting NOC and NOC-K products; Relative content profiles of different types of carbon, nitrogen and oxygen atoms in the NOC (e) and NOC-K (f).

Fig. 5 Contact angles of the NOC (a) and the NOC-K (b) electrodes.

There is no doubt that the unique feature of the NOC-K is greatly favorable for its enhanced electrochemical energy storage capacity, particularly at high rates, and to some extent, can make up for the lower graphitization of the NOC-K after KOH activation for enhanced rapid charge-storage ability.

Besides, the surface N-/O-containing functional groups play a significant role in enhancing the hydrophilicity and wettability of the carbon electrodes. To further examine the wettability of the NOC and NOC-K electrodes, the contact angle measurement was conducted, as shown in Fig. 5. Visually, the contact angle of the NOC with the 6 M KOH aqueous solution is measured as \sim 71.6°, which is greatly larger than that of the NOC-K $(\sim 47^{\circ})$. It confirms the better wettability of the NOC-K sample with the aqueous KOH, which makes the electrolyte easily impregnate and reach the inner pore surface as large as possible for higher electrochemical efficiency of the total surface area towards enhanced electrochemical behaviors.

$\mathbf{1}$. Electronical properties in a queous KoH electrolyte

Due to large SSA and unique micro-/mesopores and surface wettability, along with the heteroatom doping in the NOC-K, the

NOC-K sample can be highly expected to achieve outstanding electrochemical properties for advanced ESs as a highperformance electrode. Electrochemical performance of the NOC and NOC-K electrodes are first investigated detailedly in a three-electrode configure. Fig. 6a comparatively exhibits the typical CV curves of the NOC and NOC-K electrodes within the potential range from -1.0 to 0.0 V (vs. SCE) at a scan rate of 10 mV s^{-1} in 6 M KOH aqueous solution. The nearly rectangular CV shape observed here for the two evidences their typical electrochemical double-layer capacitance in nature. It is particularly worthy of mentioning that the electrochemical response current of the NOC-K is obviously increased when compared with that of the NOC electrode, which means that the NOC-K can deliver much higher energy-storage capacity. Fig. 6b displays the CP plots of the NOC and NOC-K at a current density of 1 A g^{-1} within the potential window ranged from -1.0 to 0.0 V (vs. SCE) in 6 M KOH electrolyte. Almost triangular appearance with linear charge–discharge plots suggests their good electrochemical supercapacitance. According to the data in Fig. 6b, the mass capacitance of the NOC-K with a loading of 5 mg cm^{-2} can be calculated as \sim 281.6 F g⁻¹, far higher than that for the NOC (\sim 88.6 F g⁻¹) at the same current density of 1 A g⁻¹ (*i.e.*, 5 mA **EXALGEMENTES ARTICLE ARTICLE CONSULTERATION** PUBLISHED CONSULTER CONSULTER THE SUCCESS ARTICLE IS THE SUCCESS ARTICLE ARTICL

Fig. 6 CV curves (a), CP plots (b), SC and VSC as a function of current density (c), and EIS spectra and fitting files (d) of the NOC and NOC-K. The inset in panel (d) for the enlarged high-medium frequency region (the lower) and corresponding equivalent circuit model (the upper).

Fig. 7 CV curves (a), CP profiles (b), SC vs. current density (c) for the NOC-K based symmetric device. Typical CV plots ((d), 100 mV s⁻¹) and CP data ((e), 1 A g⁻¹) of the symmetric devices with various loadings of NOC-K as indicated. Cycling behaviors and CE data of the NOC-K based symmetric device (f). The inset in panel (e) for the SCs as a function of total mass loading in one symmetric device.

 $\rm cm^{-2}$). Moreover, a capacitance as large as ${\sim}340.0 \; \rm F\,g^{-1}$ can be delivered by the NOC-K at 0.5 A $\rm g^{-1}$, while the NOC just delivers a capacitance of \sim 199.8 F g^{-1} , as derived from the CP plots (Fig. S2, ESI†). More impressively, when the current rates are further up to 2.0, 3.0, 4.0 and 5.0 A $\rm{g}^{-1},$ the SCs of the NPC-K in 6 M KOH still remain \sim 251.6, \sim 233.4, \sim 218.4 and \sim 200.0 F $\rm g^{-1}$, respectively, as demonstrated in Fig. 6c, that is, a SC retention of \sim 58.8% can be obtained with the current increasing from 0.5 to 5 A $\rm g^{-1}$. More encouragingly, the SCs of our fabricated NOC-K are much higher than other biomass-based carbon electrodes even with less mass loadings (Table S1, ESI†). As a sharp contrast, the NOC only shows a SC of 13.6 F g^{-1} at the current density of 5 A $\rm g^{-1}$, corresponding to a SC degradation of 93.2% when the current is up to 5 from 0.5 A $\rm g^{-1}.$ As analyzed above, the NOC-K displays superior rate behaviors especially for highrate operation. Considering the high density of \sim 1.1 g cm⁻³ for the NOC, a VSC of \sim 219.8 F cm⁻³ can be obtained accordingly, as seen in Fig. 6c, which is somewhat less than that of the NOC-K (\sim 238.0 F cm⁻³) with a density of \sim 0.7 g cm⁻³ at the same rate. Anyway, the VSCs for the NOC and NOC-K electrodes still can be compared with and/or higher than those of the porous carbon layer/graphene hybrids $(\sim]212 \text{ F cm}^{-3})$,³⁴ carbon aerogels $({\sim}123 \text{ F} \text{ cm}^{-3})^{35}$ apricot shell-based porous carbon $(\sim 171 \text{ F cm}^{-3})$,³⁶ poly(vinylidene chloride)-derived porous carbon $({\sim}214$ F cm⁻³),³⁷ and CO₂-activated ordered mesoporous carbons $({\sim}53 \text{ F cm}^{-3})$,³⁸ and so on. One should especially note

that the NOC-K provides a VSC as high as \sim 140.0 F cm⁻³, which is more than eight times higher than that of the NOC (\sim 15.0 F cm⁻³) at the same case of 5 A g⁻¹, as shown in Fig. 6c.

For further in-depth investigation into the specific electrochemical processes for the NOC and NOC-K electrodes, EIS tests were performed in the frequency from $0.01-10^5$ Hz with an AC signal amplitude of 5 mV at a potential of -0.3 V (vs. SCE). Typical Nyquist plots and corresponding fitted files with the equivalent circuit model (the upper inset) for the two are collected in Fig. 6d, and the tting parameters are summarized (Table S2, ESI†). The intersections of the plots at the X-axis reasonably represent the solution resistance (R_s) , which to a certain extent is related to the intrinsic resistance of electroactive materials itself. As observed from the enlarged highmedium frequency region (the lower inset in Fig. 6d), the R_s of the NOC-K can be estimated as \sim 0.4 ohm, somewhat smaller than that of the NOC (\sim 0.6 ohm). Furthermore, the semicircle, whose diameter stands for the charge transfer resistance (R_{ct}) , can be notable in Fig. 6d. The little semicircle of the NOC-K means its small R_{ct} (~1.6 ohm), and the R_{ct} is approximated to \sim 6.4 ohm for the NOC. Evidently, the smaller $R_{\rm ct}$ and $R_{\rm s}$ values of the NOC-K electrode should well account for its superior power behaviors, as discussed in Fig. 6c, compared with those for the NOC electrode.

In order to investigate the practical application of the NOC-K as potential electrodes for ESs, a symmetric device was

assembled by two identical NOC-K electrodes with total mass loading of 6 mg $\rm cm^{-2}$, along with the 6 M KOH as aqueous electrolyte. Fig. 7a shows the CV curves of the NOC-K based symmetric device with various sweep rates ranged from 50 to 500 mV s^{-1} in the potential window with a upper voltage of 1.0 V. Obviously, the symmetric device presents representative supercapacitive performance with quasi-rectangular shape voltammetry characteristics with respect to the zero-line and rapid current responses on voltage reversal at each end voltage, even at a high scanning rate of 500 mV s $^{-1}\!,$ suggesting its outstanding electrochemical properties in KOH. Fig. 7b plots the CP data of the symmetric supercapacitor at different current densities from 0.5 to 10.0 A $\rm{g^{-1}}.$ It can be seen that all these charge–discharge curves show a standard regular triangle, further supporting its excellent double layer capacitance characteristics. According to the data in Fig. 7b, the SCs of the device can be quantified, and collected in Fig. 7c. Remarkably, the symmetric cell yields high SCs of \sim 59.6, \sim 56.8, \sim 54.1, \sim 51.6, \sim 50.5, \sim 49.1 and \sim 44.1 F g⁻¹, at the current densities of 0.5, 1.0, 2.0, 3.0, 4.0, 5.0 and 10.0 A $\mathrm{g}^{-1},$ respectively, that is, just a SC degradation of \sim 74.0% is obtained with the current rate increasing from 0.5 to 10 A $\rm g^{-1},$ strongly suggesting its appealing rate behaviors. As is well known, electrochemical capacitances of any cell are hugely **Excharage Article is the state of the published on 02** March 2018. The state of the published on 02 March 2018. The state of the published on 1/6 March 2018. Downloaded to the state of the state of the state of the state

dependent upon the mass loadings of the electroactive material. Fig. 7d contrastively shows the CV curves at a sweep rate of 100 mV s $^{-1}$ for the three symmetric capacitors with the total mass loadings of 6, 8 and 10 mg, respectively. Evidently, higher electrochemical current responses and larger integrated areas under the I–E curves are observed with the mass loading increasing. As calculated from the CP plots in Fig. 7e, the SCs of the symmetric cells are still maintained as \sim 50.8 and \sim 50.5 F g⁻¹, which are \sim 85.2% and \sim 84.7% of the SC for the symmetric cell with the mass loading of 6 mg, when the total mass loading is up to 8 and 10 mg. The excellent SC independence of the NOC-K upon the mass loading fully highlights its remarkable capability for versatile potential applications.

Fig. 7f illustrates the long-span electrochemical stability of the symmetric device with a mass loading of 6 mg. Strikingly, the high SC retention of \sim 95.8% over continuous 8200 cycles, *i.e.*, \sim 0.05% SC decay per 100 cycles, consistently verifies the excellent long-term cyclic stability of the NOC-K based symmetric cell at a current rate of 0.5 A $\rm g^{-1}$. Additionally, the CE of the cell is nearly always kept as high as \sim 100%, suggesting its highly electrochemical reversibility for efficient charge storage. This fully confirms the promising advantages of the NOC-K electrode for advanced ESs.

Fig. 8 Electrochemical evaluation of the NOC-K based symmetric device with an aprotic electrolyte of 1 M TEABF₄/PC: CV curves (a) and CP plots (b) with various upper voltage limits as indicated; CV curves at different sweep rates (c) and CP plots at various current rates (d) within the potential range from 0.0 to 2.8 V; SC as a function of current density (e) and cycling performance ((f), 0.5 A g⁻¹) with the upper potential limit of 2.8 V.

Fig. 9 Ragone plots for the NOC-K based symmetric devices with the TEABF₄/PC and KOH as the electrolytes (a), and digital images ((b) and (c)) for a light-emitting diode indicator powered by our symmetric cell with the TEABF₄/PC electrolyte. The red and black data in panel (a) are for aqueous and organic symmetric systems, respectively.

3.3. Electrochemical properties in organic TEABF₄/PC electrolyte electrolyte

As we all know, the use of organic electrolytes is in high demand for practical application because of their even larger working potential window for high-performance ESs when compared to aqueous systems, which would further renders a large SE. We next test the coin-type symmetric supercapacitor with an organic TEABF4/PC as the electrolyte. Fig. 8a presents the CV curves of the symmetric device within the different upper potential limits from 2.5 to 3.1 V. Noteworthily, when the voltage is higher than 2.8 V, there exists a noticeable strong peak beyond 2.8 V, maybe corresponding to the electrolyte decomposition and/or some gas evolution. As a result, the CE of the symmetric device decreases from 99.7% (2.5 V) to 97.3% (3.1 V), as estimated from the CP plots in Fig. 8b. Furthermore, the SCs within the different voltage ranges just change from \sim 44 to \sim 45 F g⁻¹. But serious SC degradation of \sim 24.1% is observed after 6500 cycles within the potential window from 0.0 to 3.0 V (Fig. S4, ESI†). Therefore, we choose the upper voltage of 2.8 V for the following detailed characteristics. As seen in Fig. 8c, all these CV curves of the device exhibit nearly perfect rectangular shape at various rates from 100 to 1000 mV s^{-1} in the electrochemical window from 0.0 to 2.8 V, and still keep the geometrical rectangle even at a super-high rate of 1000 mV $\mathrm{s}^{-1},$ revealing the remarkable rate behaviors of the cell with the organic electrolyte. Typical CP plots at various current rates from 0.5 to 10 A g^{-1} , as demonstrated in Fig. 8d, show symmetric triangle with nearly linear potential-time behaviors in the potential range from 0.0 to 2.8 V, further verifying the desirable electrochemical capacitances of the symmetric cell. Accordingly, the SCs of the symmetric supercapacitor, as plotted in Fig. 8e, can be calculated as \sim 48.3, \sim 45.6, \sim 42.1, \sim 40.1, \sim 39.1, \sim 37.9 and \sim 35.7 F g⁻¹ at the current densities of 0.5, 1.0, 2.0, 3.0, 4.0, 5.0 and 10.0 A g^{-1} , respectively. Encouragingly, \sim 73.4% of the capacitance still can be retained with the current rate increasing from 0.5 to 10.0 A g^{-1} . This means the superior capacitive behaviors of the cell, which is of great importance in practical ESs, particularly under high-power operation conditions. Fig. 8f demonstrates the stable capacitance retention of NOC-K based symmetric supercapacitor over 6500 consecutive charge–discharge cycles with the upper voltage limit of 2.8 V at a current density of 0.5 A g^{-1} . Remarkably, the device achieves a SC retention of \sim 96.6% at the end of the 6500th cycle, which corresponds to a small SC degradation of $\sim 0.05\%$ per 100 charge–discharge cycles over the prolonged cycles. Of note, it is much closed to the SC decay of \sim 97.7% for the case with the upper voltage of 2.5 V (Fig. S5, ESI†). Besides this, the electrochemical CE is mainly kept around 100% during these consecutive cycles.

Ragone plots of the NOC-K based symmetric devices with the TEABF4/PC and aqueous KOH electrolytes are further depicted in Fig. 9a. As shown in Fig. 9a, the aqueous symmetric cell with 6 M KOH as electrolyte just yields a maximal SE of \sim 8.3 W h kg⁻¹ (*i.e.*, \sim 2.7 W h L⁻¹) along with a SP of 250 W kg⁻¹, and at the case of 5 kW kg⁻¹, only a SE of \sim 6.1 W h kg⁻¹, corresponding to a volumetric SE of \sim 2.0 W h L⁻¹, can be rendered. Appealingly, the gravimetric SE for the aqueous cell can be compared to, and/or even higher than others reported before,5,39,40,42,43,48 as plotted in Fig. 9a. As for the organic device, the highest SE of the cell with 1 M TEABF $_4$ /PC electrolyte has been obtained as large as \sim 52.6 W h kg⁻¹ (\sim 2.6 W h L⁻¹) at a SP of \sim 700 W kg⁻¹. And attractively, the device still can achieve a SE of \sim 38.9 W h kg⁻¹ (\sim 1.9 W h L⁻¹) at a higher SP of \sim 14.0 kW kg⁻¹, which is much larger than those for other carbonbased symmetric systems,^{5,41-49} as summarized in Fig. 9a. Moreover, the symmetric device with the TEABF4/PC electrolyte can efficiently power a light-emitting diode indicator, and maintain it for around 90 s, as visually demonstrated in Fig. 9b and c.

4. Conclusions

In summary, in the work, we adopted a simple yet industrializing synthetic strategy to fabricate low-cost N/O-enriched carbon (NOC-K) with bi-modal micro-/mesoporosity by using the natural rose multiflora as a precursor with the general KOH

activation. Physicochemical characterizations demonstrated the rose multiflora-derived NOC-K possessed a large SSA, hierarchical micro-/mesoporosity, high surface wettability, and high-content elemental N and O. It was benefiting from these intrinsic structural and compositional advantages that the resultant NOC-K electrode with a mass loading of 5 mg cm^{-2} yielded large gravimetric/volumetric SCs of \sim 340.0 F g^{-1} (\sim 238.0 F cm⁻³) at a current rate of 0.5 A g⁻¹, and a SE of \sim 8.3 W h kg⁻¹ at 250 W kg⁻¹, when evaluated as electroactive material in 6 M KOH aqueous solution for ESs. More competitively, the NOC-K based symmetric cell with 1 M TEABF4/PC organic electrolyte delivered an exciting SE of \sim 52.6 W h kg⁻¹ and excellent cycling performance with $\sim 0.05\%$ SC decay per 100 cycles over 6500 charge–discharge cycles. Therefore, we strongly believe that the as-fabricated NOC-K holds enormous potential for advanced ESs, and even high-performance Na/Liion batteries as a promising electrode material. **BSC Advances**

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Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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