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# Multifunctional fluorine–nitrogen modified carbon quantum dots for enhanced solid polymer electrolytes toward solid-state lithium batteries

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Solid polymer electrolytes (SPEs) are pivotal for developing all-solid-state lithium batteries due to their enhanced safety, but their practical application is hindered by low ionic conductivity. Herein, fluorine–nitrogen–modified carbon quantum dots (FNCDs) are designed and prepared as a multifunctional filler to overcome this limitation. The FNCDs provide abundant sites for Li<sup>+</sup> transport and immobilize anions (TFSI<sup>-</sup>), promoting Li salt dissociation. Moreover, the fluorinated surface ensures excellent compatibility with the fluorinated polymer matrix (PVDF–HFP), facilitating favorable interfacial Li<sup>+</sup> hopping channels. Consequently, the FNCD–incorporated SPE (FNCDs–SPE) film achieves a high ionic conductivity of  $9.59 \times 10^{-4} \text{ S cm}^{-1}$  and a Li<sup>+</sup> transference number of 0.77. It also exhibits enhanced mechanical strength (3.53 MPa) and elasticity (249.2% elongation). When paired with LiFePO<sub>4</sub>, LiCoO<sub>2</sub>, or NCM811 cathodes, the corresponding solid-state cells demonstrate exceptional cycling stability. This work highlights the great potential of zero-dimensional fluorinated carbon materials in developing high-performance SPEs.

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## 1 Introduction

With the growing demand for high safety and high energy density in electric vehicles and renewable energy storage, conventional lithium-ion batteries, which rely on flammable liquid electrolytes and graphite anodes, have become increasingly inadequate. In traditional lithium-ion batteries, the flammable liquid electrolytes not only pose significant thermal runaway risks and serious safety hazards but also exhibit restricted compatibility with high-energy lithium metal anodes.<sup>1,2</sup> Consequently, solid-state lithium metal batteries (SSLMBs) have garnered considerable research interest in recent years.<sup>3–5</sup> The electrolyte, as a critical component, plays a pivotal role in realizing battery systems with high safety and energy density. Although solid-state electrolytes (SSEs) can mitigate safety concerns, they face intrinsic challenges including lower ionic conductivity, poor interfacial contact, stringent fabrication requirements, and higher cost compared to their liquid counterparts.<sup>6–9</sup> Overcoming these challenges is therefore imperative for the advancement of SSLMBs.

In response to these challenges, extensive research efforts have been devoted to developing various SSE material systems. The mainstream SSE material systems primarily include inorganic solid electrolytes (ISEs), organic solid

polymer electrolytes (SPEs), and composite polymer electrolytes (CPEs). While ISEs (*e.g.*, oxides, sulfides, and halides) typically exhibit high ionic conductivity ( $10^{-4}$  to  $10^{-2} \text{ S cm}^{-1}$ ), they often suffer from poor interfacial contact, sensitivity to moisture/oxygen in the air, high cost, and poor processability, which severely hinder their practical application.<sup>10,11</sup> In contrast, SPEs have emerged as promising candidates for practical large-scale production owing to their excellent flexibility, good interfacial compatibility, facile processability, and cost-effectiveness.<sup>12–14</sup> However, commonly studied SPEs based on polymers like polyethylene oxide (PEO), poly(vinylidene fluoride) (PVDF), polyacrylonitrile (PAN), and poly(methyl methacrylate) (PMMA) generally suffer from low ionic conductivity (typically  $10^{-7}$  to  $10^{-5} \text{ S cm}^{-1}$ ) and narrow electrochemical stability windows, which constitute the core challenges for their development.<sup>15,16</sup> Nevertheless, significant improvements in both ionic conductivity and electrochemical stability have been achieved through strategies such as polymer matrix modification and the incorporation of inorganic fillers. For instance, Cao *et al.* synthesized cellulose phthalate (CP) *via* homogeneous esterification, and the corresponding CP-based SSE achieved a high ionic conductivity of  $1.09 \text{ mS cm}^{-1}$  and a Li<sup>+</sup> transference number of 0.81 at room temperature. This enhancement is attributed to the introduced phthalate groups, which enable multiple coordination with Li<sup>+</sup> and establish a stable conduction network.<sup>17</sup> Similarly, Ahn *et al.* developed a composite electrolyte (PLiZ) by incorporating ZIF-67 and an ionic liquid (ILE) into a PEO matrix, which

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achieved an ionic conductivity of  $1.19 \times 10^{-4} \text{ S cm}^{-1}$  and a  $\text{Li}^+$  transference number of 0.8 at room temperature. The fillers shortened  $\text{Li}^+$  transport pathways and immobilized anions, thereby enhancing battery performance.<sup>18</sup> These examples underscore the considerable potential of composite strategies for advancing SPEs.

In pristine (unfilled) SPEs, lithium-ion transport primarily relies on the segmental motion of polymer chains. Ionic conduction occurs predominantly in the amorphous regions, whereas it is highly restricted in the crystalline regions due to the tightly packed and ordered polymer chains.<sup>19</sup> Therefore, rational polymer structural design is crucial for facilitating efficient  $\text{Li}^+$  transport. Poly(vinylidene fluoride-co-hexafluoropropylene) (PVDF-HFP) is a copolymer formed with vinylidene fluoride (VDF) and hexafluoropropylene (HFP) monomers. The VDF segments provide a high dielectric constant to promote lithium salt dissociation and confer mechanical robustness, while the HFP segments introduce structural defects that disrupt the regularity of PVDF chains and increase the amorphous phase content. This synergy creates a favorable environment for  $\text{Li}^+$  transport.<sup>20,21</sup> Beyond polymer design, the incorporation of inorganic fillers is equally critical for enhancing SPE performance. Carbon quantum dots (CDs), a class of zero-dimensional carbon nanomaterials featuring abundant and tunable surface functional groups, have attracted significant research attention.<sup>22–25</sup> Their surfaces are typically rich in functional groups such as hydroxyl (–OH), carboxyl (–COOH), and amino (–NH<sub>2</sub>). These groups facilitate uniform dispersion within polymer matrices, interact with polymer chains to suppress crystallinity, and provide additional sites for ion transport. Furthermore, the rich surface chemistry of CDs can positively influence electrolyte mechanical properties and interfacial stability with electrodes. Therefore, the rational design and modification of inorganic fillers like CDs are of great importance for constructing fast and stable  $\text{Li}^+$  transport channels and optimizing interfacial compatibility within polymer electrolyte systems.

In this work, we developed a high-performance, flexible solid polymer electrolyte film (denoted as FNCDs-SPE) by incorporating fluorine–nitrogen modified carbon quantum dots (FNCDs) into a PVDF-HFP matrix. The abundant surface functional groups on FNCDs enable their excellent dispersion within the polymer matrix and prevent performance degradation caused by agglomeration. Simultaneously, they disrupt the ordered arrangement of PVDF-HFP chains, thereby increasing the amorphous region content, which is beneficial for ion transport. Notably, the fluorine- and oxygen/nitrogen-containing functional groups on the FNCDs can engage in Lewis acid–base interactions with  $\text{Li}^+$ , promoting lithium salt dissociation and providing additional hopping sites for  $\text{Li}^+$  transport. The resulting rapid ion conduction facilitates uniform lithium deposition/stripping, suppresses lithium dendrite growth, and enhances the interfacial stability. Moreover, FNCDs act as physical cross-linking points to entangle and interact with PVDF-HFP

molecular chains, which significantly improves the mechanical robustness of the electrolyte. As a result, FNCDs-SPE exhibits a high ionic conductivity of  $9.59 \times 10^{-4} \text{ S cm}^{-1}$  and a tensile strength of 3.53 MPa, which are competitive among state-of-the-art polymer-based electrolytes. The Li||Li symmetric cells assembled with FNCDs-SPE demonstrate stable cycling for over 2000 hours. Full cells incorporating FNCDs-SPE paired with commercial cathodes, including  $\text{LiFePO}_4$  (LFP),  $\text{LiCoO}_2$  (LCO), and  $\text{LiNi}_{0.8}\text{Co}_{0.1}\text{Mn}_{0.1}\text{O}_2$  (NCM811), all demonstrate outstanding capacity retention and rate capability. This study provides valuable insights into the design of high-performance SPEs through functional nanofiller engineering and paves a way toward practical high-safety and high-energy-density solid-state batteries.

## 2 Experimental

### 2.1 Synthesis of NCDs and FNCDs

Nitrogen-doped carbon quantum dots (NCDs) were synthesized *via* a solvothermal method. Specifically, 1 g of anhydrous citric acid and 0.8 g of *m*-phenylenediamine were dissolved in 10 mL of formamide under magnetic stirring until a homogeneous solution was obtained. The mixture was then transferred to a Teflon-lined stainless-steel autoclave and heated at 160 °C for 4 h. After the autoclave had cooled to room temperature, the resulting brownish-black solution was mixed with 30 mL of ethanol. The mixture was then centrifuged, and the precipitate was collected and washed with ethanol several times to remove impurities. This washing cycle was repeated three times. The purified precipitate was re-dispersed in deionized water, dialyzed against fresh deionized water for 36 h, and finally lyophilized to yield the NCDs as a solid powder.

FNCDs were obtained *via* a one-step fluorination of the as-synthesized NCDs. Typically, 250 mg of NCDs were dispersed in 19 mL of ethanol by ultrasonication for 30 min. The dispersion was then transferred to a Teflon-lined autoclave, followed by the addition of 1 mL of a 40 wt% hydrofluoric acid (HF) aqueous solution. The autoclave was sealed and maintained at 160 °C for 12 h to facilitate the fluorination reaction. After cooling to room temperature, the crude product was transferred to a Teflon beaker. To neutralize any residual HF, 25 mL of a saturated sodium bicarbonate ( $\text{NaHCO}_3$ ) aqueous solution was added slowly, followed by the addition of 100 mL of deionized water. The mixture was then centrifuged, and the collected solid was washed repeatedly with deionized water until the supernatant reached neutral pH. The final product (FNCDs) was obtained by lyophilization.

### 2.2 Preparation of FNCDs-based electrolyte films

The FNCDs-SPE films were fabricated using a solution casting technique. First, 1 g of PVDF-HFP (average  $M_w = \sim 600\,000$ ) and 1 g of bis(trifluoromethanesulfonyl)imide lithium salt (LiTFSI) were dissolved in 7 mL of *N,N*-dimethylformamide (DMF) in a glass vial under magnetic



stirring at 80 °C for 20 min. Subsequently, a predetermined amount of the ionic liquid plasticizer 1-allyl-2,3-dimethylimidazolium chloride (AmimCl) and 50 mg of FNCD powder were added to the above solution. The mixture was continuously stirred at 80 °C for 12 h to obtain a homogeneous and viscous casting solution. The casting solution was then poured onto a leveled polytetrafluoroethylene (PTFE) dish and allowed to rest for initial solvent evaporation. The dish was then transferred to a vacuum oven and dried at 80 °C for 24 h to thoroughly remove the DMF solvent, yielding a freestanding FNCDs-SPE film. The film was finally cut into disks of desired dimensions for subsequent measurements and cell assembly. For comparison, control electrolyte films, denoted as NCDs-SPE and P-SPE (pristine SPE), were prepared following the identical procedure but by replacing the 50 mg of FNCDs with an equal mass of NCDs or by omitting the carbon quantum dot filler entirely, respectively.

### 2.3 Material characterization and electrochemical measurements

The morphology, chemical structure, and physicochemical properties of the synthesized carbon quantum dots (NCDs and FNCDs) and the solid polymer electrolyte films were thoroughly investigated. The electrochemical performance, including ionic conductivity, lithium-ion transference number, electrochemical stability window, and cycling stability in both symmetric Li||Li cells and full cells (with LFP, LCO, and NCM811 cathodes), was systematically evaluated. Detailed descriptions of the characterization techniques, measurement procedures, and specific instrument parameters are provided in the SI.

## 3 Results and discussion

NCDs and FNCDs are synthesized through a simple hydrothermal process. Constant high temperature provides a favorable environment for the carbonization, polymerization and cross-linking of citric acid (CA) and *m*-phenylenediamine (*m*-PD), leading to the formation of zero-dimensional carbon nanomaterials. The surface of NCDs is rich in C=O and -OH functional groups, which provides active sites for subsequent fluorination (Fig. S1). The formation of C-F bonds occurs *via* insertion of HF into the defect sites of NCDs, forming reactive intermediates that subsequently undergo addition to C=C bonds or substitution with oxygen-containing groups, yielding covalently bonded fluorine on the carbon dot surface.<sup>26</sup> As illustrated in Fig. 1a, the solid polymer electrolyte is fabricated using a solution casting method. The FNCDs are incorporated into the PVDF-HFP matrix, where they act as cross-linking sites and are uniformly dispersed among the polymer chains. This uniform dispersion effectively suppresses polymer crystallization, thereby enhancing the mechanical strength and toughness of the electrolyte. Meanwhile, an ionic liquid (IL) is introduced as a

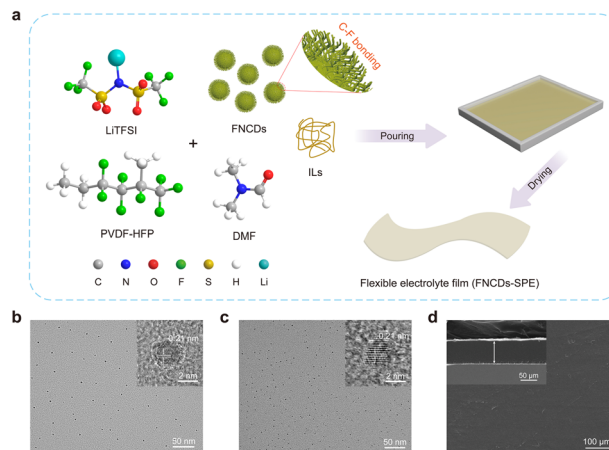


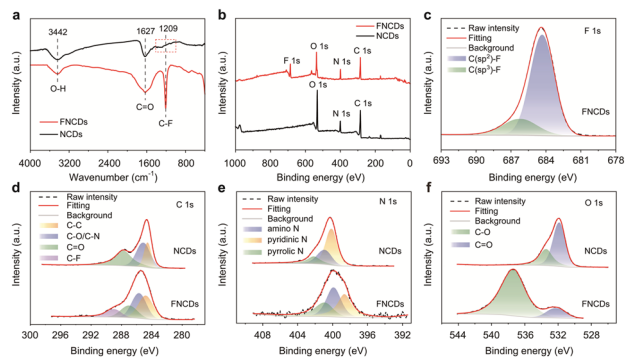
Fig. 1 (a) Schematic illustration of the synthetic route of FNCDs-SPE. TEM images of (b) NCDs and (c) FNCDs. (d) SEM image of the FNCDs-SPE film. The insets of (b) and (c) are the HRTEM images of NCDs and FNCDs, respectively. The inset of (d) is the cross-sectional SEM image of the FNCDs-SPE film.

plasticizer.<sup>17</sup> After solvent evaporation, a light-yellow, flexible freestanding film, denoted as FNCDs-SPE, is obtained.

The morphologies and structures of NCDs and FNCDs were investigated by transmission electron microscopy (TEM). The TEM images (Fig. 1b and c) reveal that both NCDs and FNCDs are uniformly dispersed and exhibit a quasi-spherical morphology. Statistical analysis of over 100 nanoparticles (Fig. S2) indicates that both samples have a similar average particle size of approximately 2.5 nm. The lattice fringe spacing measured from high-resolution TEM (HRTEM) images (insets of Fig. 1b and c) is  $\sim 0.21$  nm, corresponding to the (100) plane of graphite.<sup>27</sup> The morphology of the polymer electrolyte film was examined by scanning electron microscopy (SEM). As shown in Fig. 1d, the surface of the FNCDs-SPE film is smooth and dense without visible cracks or pores, which is further confirmed by its cross-sectional view (inset of Fig. 1d). The thickness of the electrolyte film is approximately 67  $\mu\text{m}$ . In contrast, the surfaces of the control NCDs-SPE and P-SPE films are comparatively rough (Fig. S3). Their cross-sectional images reveal the presence of some pores and inhomogeneous regions, which could result in uneven  $\text{Li}^+$  flux and localized high current density, ultimately impairing battery performance. The dense and homogeneous structure of FNCDs-SPE is conducive to rapid and uniform  $\text{Li}^+$  transport. This improved uniformity is attributed to the strong interactions, such as hydrogen bonding, between the FNCDs and the polymer chains.

The surface chemical structures of NCDs and FNCDs were characterized by Fourier transform infrared spectroscopy (FT-IR), nuclear magnetic resonance (NMR) spectroscopy and X-ray photoelectron spectroscopy (XPS). The FT-IR spectra (Fig. 2a) display several broad absorption bands from 500 to 4000  $\text{cm}^{-1}$ . The peaks at 3442 and 1627  $\text{cm}^{-1}$  correspond to the stretching vibrations of O-H and





**Fig. 2** (a) FT-IR spectra and (b) XPS survey spectra of NCDs and FNCDs. (c) High-resolution F 1s XPS spectrum of FNCDs. High-resolution (d) C 1s, (e) N 1s and (f) O 1s XPS spectra of NCDs and FNCDs.

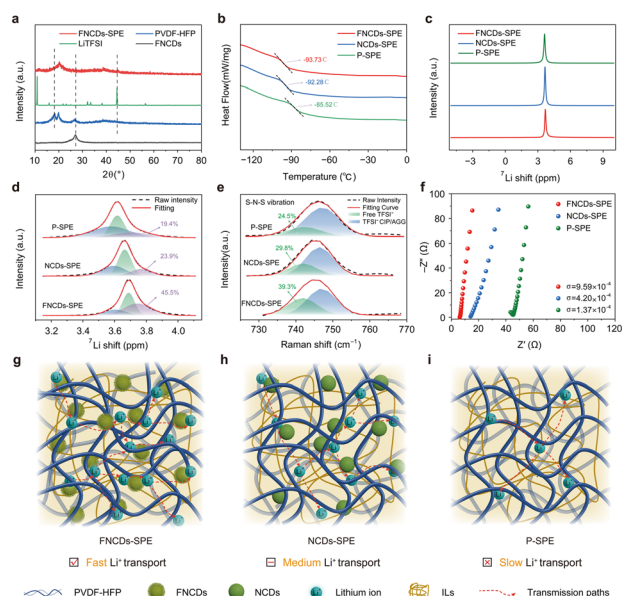
C=O bonds, respectively. Notably, a distinct absorption peak at  $1209\text{ cm}^{-1}$  appears in the spectrum of FNCDs, which is assigned to the C-F stretching vibration.<sup>28</sup> On the contrary, this peak is absent in the spectrum of NCDs, confirming the successful introduction of fluorine atoms onto the carbon dot surface. To further investigate the chemical state of fluorine in FNCDs, we performed solid-state  $^{19}\text{F}$  NMR analysis on the FNCDs sample. The  $^{19}\text{F}$  NMR spectrum (Fig. S4) exhibits main signals in the range of  $-140$  to  $-170$  ppm, characteristic of covalent C-F bonds such as  $\text{C}(\text{sp}^3)\text{-F}$  and  $\text{C}(\text{sp}^2)\text{-F}$  bonds.<sup>29</sup> No detectable signal is observed at around  $-224$  ppm (characteristic of ionic fluoride, *e.g.*, NaF),<sup>30</sup> confirming the absence of residual fluoride salts. These results confirm that fluorine in FNCDs exists primarily as C-F bonds. Furthermore, XPS analysis was employed to probe the surface composition and bonding states. The XPS survey spectrum (Fig. 2b) confirms that FNCDs are composed of C, N, O, and F. The absence of F 1s peak in the survey spectrum of NCDs is consistent with the FT-IR results. The high-resolution F 1s XPS spectrum of FNCDs (Fig. 2c) shows two peaks at 684.3 and 686.3 eV, corresponding to the  $\text{C}(\text{sp}^2)\text{-F}$  and  $\text{C}(\text{sp}^3)\text{-F}$  bonds, respectively.<sup>28</sup>

The integrated areas of the F 1s XPS peaks indicate that the contents of  $\text{C}(\text{sp}^3)\text{-F}$  bonds and  $\text{C}(\text{sp}^2)\text{-F}$  bonds are 21.84% and 78.16%, respectively. The  $\text{C}(\text{sp}^3)\text{-F}$  bonds primarily originate from the fluorination substitution of oxygen-containing functional groups (such as  $-\text{OH}$  and  $-\text{COOH}$ ) located at the edges and defect sites of the carbon dots, which are the most chemically active regions on the carbon dot surface.<sup>31</sup> Their strong electron-withdrawing effect can more effectively decouple  $\text{Li}^+\text{-TFSI}^-$  interactions and promote lithium salt dissociation. The  $\text{C}(\text{sp}^2)\text{-F}$  bonds are mainly situated within the carbon skeleton plane and contribute to modulating the electronic structure and stability of the carbon dots.<sup>32</sup>

Deconvolution of the high-resolution C 1s spectrum of NCDs (Fig. 2d) reveals peaks at 284.8, 285.4 and 288.0 eV, attributable to C-C, C-O/C-N and C=O bonds, respectively.

In contrast, the C 1s spectrum of FNCDs shows an additional distinct peak at 289.1 eV, which is assigned to the C-F bond.<sup>23</sup> The high-resolution N 1s spectra (Fig. 2e) indicate that nitrogen in both NCDs and FNCDs exists primarily in the forms of amino N (400.9 and 399.9 eV), pyridinic N (400.2 and 398.6 eV) and pyrrolic N (402.1 and 401.0 eV). The high-resolution O 1s spectra (Fig. 2f) are deconvoluted into two components, corresponding to the C=O and C-O bonds. For NCDs, these peaks are located at 531.9 and 533.45 eV, respectively. After fluorination, the respective binding energies of FNCDs shift to 532.3 and 537.4 eV.<sup>33</sup> Following fluorination, the intensity of the C=O component decreases significantly, while the C-O component increases and shifts to a higher binding energy. This suggests the presence of multiple types of C-F bonds including  $\text{C}(\text{sp}^2)\text{-F}$  and  $\text{C}(\text{sp}^3)\text{-F}$  on the carbon dot skeleton. The notable increase in binding energy is attributed to the strong electron-withdrawing inductive effect of fluorine atoms, which reduces the electron cloud density around both the carbon and oxygen atoms.

The crystallinity and component interactions within the FNCDs-SPE film were investigated by X-ray diffraction (XRD). As shown in Fig. 3a, the characteristic diffraction peaks belonging to FNCDs and LiTFSI are absent in the XRD pattern of FNCDs-SPE, indicating the formation of a predominantly amorphous structure. Furthermore, the diffraction peak of pure PVDF-HFP at  $18.2^\circ$  disappears in the FNCDs-SPE pattern. This can be ascribed to the incorporated FNCDs disrupting the conformational rearrangement and packing of the polymer chains, thereby significantly reducing



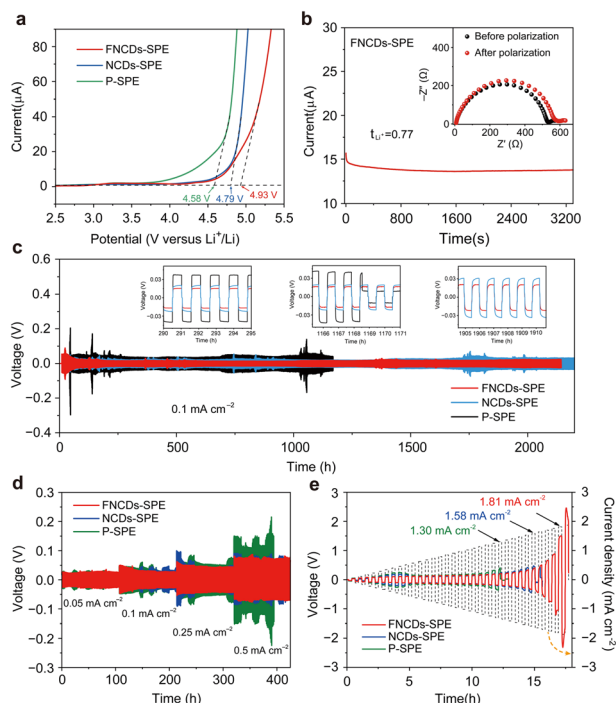
**Fig. 3** (a) XRD patterns of the FNCDs-SPE film and the related components. (b) DSC curves, (c)  $^7\text{Li}$  solid-state NMR spectra, (d) the corresponding fitting peaks in the  $^7\text{Li}$  NMR spectra and (e) Raman spectra of FNCDs-SPE, NCDs-SPE and P-SPE films. (f) Nyquist plots in stainless steel (SS) symmetric cells of FNCDs-SPE, NCDs-SPE and P-SPE. (g) Schematic illustrations of possible  $\text{Li}^+$  transmission paths in FNCDs-SPE, (h) NCDs-SPE and (i) P-SPE.



its crystallinity. A comparison of the XRD patterns of FNCDs-SPE, NCDs-SPE, and P-SPE films (Fig. S5) further confirms that FNCDs are most effective in promoting amorphization of the polymer matrix, which is favorable for segmental motion and  $\text{Li}^+$  transport. Correspondingly, the FNCDs-SPE film exhibits excellent flexibility and mechanical strength (Fig. S6), which are crucial for suppressing lithium dendrite penetration. Quantitative tensile tests (Fig. S7 and Table S1) reveal that the FNCDs-SPE film possesses the highest elongation at break (249.2%). Although its tensile strength (3.54 MPa) is lower than that of NCDs-SPE (5.77 MPa), it remains substantially higher than that of P-SPE (1.58 MPa), demonstrating a good balance between toughness and strength. The thermal stability of these three electrolyte films was examined by thermogravimetric analysis (TGA). TGA curves (Fig. S8) show that all electrolyte films exhibit comparable thermal stability, undergoing a two-stage weight loss corresponding to residual solvent evaporation followed by decomposition of the lithium salt and polymer.<sup>21</sup>

Differential scanning calorimetry (DSC) results (Fig. 3b) show that the FNCDs-SPE film has the lowest glass transition temperature ( $T_g = -93.73$  °C) compared to NCDs-SPE ( $-92.28$  °C) and P-SPE ( $-85.52$  °C), confirming enhanced chain mobility, which facilitates  $\text{Li}^+$  transport.<sup>34,35</sup> To investigate the local lithium chemical environments, solid-state  $^7\text{Li}$  NMR spectroscopy was performed. Compared with P-SPE, the  $^7\text{Li}$  NMR signal of carbon dot-filled SPEs shifts to a higher field (Fig. 3c), indicating a weaker interaction between  $\text{Li}^+$  and the PVDF-HFP matrix after filling NCDs or FNCDs. By fitting the  $^7\text{Li}$  NMR spectra, distinct  $\text{Li}^+$  environments are identified (Fig. 3d). The peak at higher field (shaded in purple) is typically associated with mobile  $\text{Li}^+$ , which contributes more effectively to ionic conduction. This mobile  $\text{Li}^+$  fraction reaches 45.5% in FNCDs-SPE, significantly higher than those in NCDs-SPE (23.9%) and P-SPE (19.4%). In contrast, the lower-field components correspond to  $\text{Li}^+$  trapped by oxygen or fluorine in PVDF-HFP, which are less conducive to migration.<sup>36</sup> These results confirm that FNCDs facilitate the release of mobile  $\text{Li}^+$ . Raman spectroscopy was employed to probe the changes in Li-TFSI interactions within the polymer electrolytes (Fig. 3e). The characteristic band near  $745\text{ cm}^{-1}$ , associated with the S–N–S deformation of the TFSI<sup>−</sup> anion, is deconvoluted to quantify the populations of free TFSI<sup>−</sup> ( $741.6\text{ cm}^{-1}$ ), contact ion pairs (CIPs), and ion aggregates (AGGs).<sup>37,38</sup> The FNCDs-SPE film shows the highest proportion of free TFSI<sup>−</sup> (39.3%), indicating the most effective promotion of LiTFSI dissociation. This suggests that more  $\text{Li}^+$  ions are liberated to coordinate with and migrate along the FNCDs and polymer chains, resulting in higher ionic conductivity. In contrast, a higher proportion of  $\text{Li}^+$  in the NCDs-SPE and P-SPE films exists as CIPs or AGGs, which impedes cationic mobility. This analysis is also supported by FT-IR spectra (Fig. S9), where the S=O stretching vibration of TFSI<sup>−</sup> shifts to lower wavenumbers in the composites, signifying changes in its chemical environment.<sup>17</sup>

Electrochemical impedance spectroscopy (EIS) reveals that the FNCDs-SPE electrolyte has the lowest bulk resistance (Fig. 3f), leading to the highest ionic conductivity of  $9.59 \times 10^{-4}\text{ S cm}^{-1}$ , as calculated from Table S2 and eqn (S1) in the SI. This value is significantly higher than those of NCDs-SPE ( $4.20 \times 10^{-4}\text{ S cm}^{-1}$ ) and P-SPE ( $1.37 \times 10^{-4}\text{ S cm}^{-1}$ ). A schematic illustration of the proposed  $\text{Li}^+$  transport mechanisms in the three electrolytes is presented in Fig. 3g–i. The superior conductivity of FNCDs-SPE originates from two synergistic effects. First, the uniformly dispersed FNCDs act as physical cross-linkers, disrupting polymer chain order and expanding the amorphous domain for faster segmental motion. Second, the abundant surface functional groups (*e.g.*, C–F, C–N, –OH, and –COOH) on FNCDs engage in Lewis acid–base interactions with  $\text{Li}^+$ , providing numerous additional hopping sites for ion transport. Specifically, the fluorine atoms with a strong electron-withdrawing effect weaken the coupling between  $\text{Li}^+$  and TFSI<sup>−</sup>, thereby promoting salt dissociation,<sup>39</sup> while the nitrogen-containing groups enhance the lithiophilicity of the filler, reducing the nucleation overpotential for  $\text{Li}^+$  deposition.<sup>40</sup> These combined effects establish continuous and efficient conduction pathways, leading to the observed enhancement in ionic conductivity and  $\text{Li}^+$  transference number.

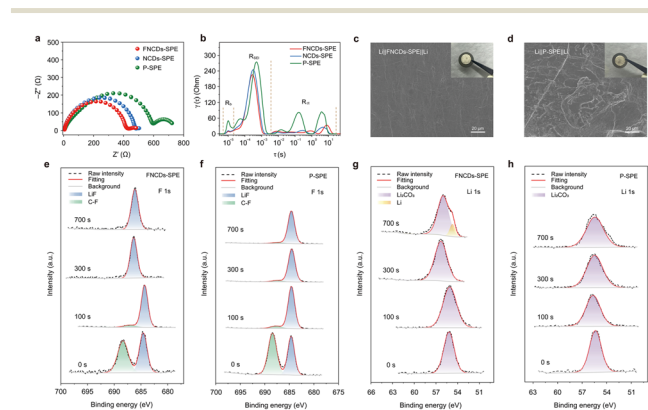


**Fig. 4** (a) LSV curves in Li||Li symmetrical cells of FNCDs-SPE, NCDs-SPE and P-SPE. (b) Polarization curve of the Li||FNCDs-SPE||Li cell (inset: the impedance diagram before and after polarization). (c) Galvanostatic cycling curves in Li||Li symmetrical cells of FNCDs-SPE, NCDs-SPE and P-SPE at a current density of  $0.1\text{ mA cm}^{-2}$  (inset: local magnification curves). (d) The cycling curves in Li||Li symmetrical cells of FNCDs-SPE, NCDs-SPE and P-SPE at different current densities. (e) Critical current density (CCD) curves in Li||Li symmetrical cells of FNCDs-SPE, NCDs-SPE and P-SPE.



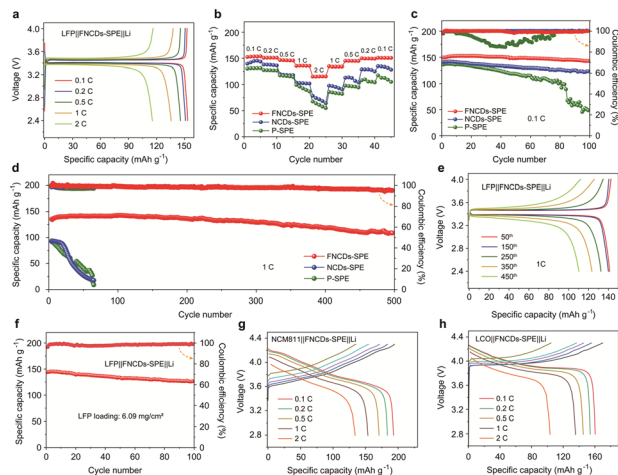
The electrochemical performance of the electrolytes was further investigated using Li||Li symmetric cells. Linear sweep voltammetry (LSV) results (Fig. 4a) demonstrate that the FNCDs-SPE electrolyte possesses the widest electrochemical stability window (up to 4.93 V vs. Li/Li<sup>+</sup>), compared to NCDs-SPE (4.79 V) and P-SPE (4.58 V), indicating its compatibility with high-voltage cathodes. The chronoamperometry tests (Fig. 4b) show that the FNCDs-SPE electrolyte also achieves the highest Li<sup>+</sup> transference number ( $t_{Li^+} = 0.77$ ), confirming that ionic conduction is predominantly cationic.<sup>41</sup> In contrast, the NCDs-SPE and P-SPE electrolytes exhibit lower  $t_{Li^+}$  values of 0.66 and 0.41, respectively (Fig. S10 and Table S3). Galvanostatic cycling tests (Fig. 4c) of these three cells at a current density of 0.1 mA cm<sup>-2</sup> demonstrate stable cycling for over 2000 h. The Li||FNCDs-SPE||Li cell exhibits the lowest polarization voltage (~20 mV) and the most stable long-term performance, indicating uniform lithium deposition and stripping behavior. By contrast, the Li||P-SPE||Li cell shows inferior cycling stability, experiencing a premature short circuit after 1168 h, which is likely attributable to lithium dendrite penetration through the electrolyte film. Even when the current density is increased from 0.05 to 0.5 mA cm<sup>-2</sup>, the Li||FNCDs-SPE||Li cell maintains stable cycling with a relatively low polarization voltage (Fig. 4d). This enhanced stability is ascribed to the abundant functional groups on the FNCDs, which help homogenize the Li<sup>+</sup> flux at the electrode interface, thereby guiding uniform lithium deposition/stripping and suppressing dendrite growth. The critical current density (CCD), which defines the maximum sustainable current before cell failure, was also evaluated.<sup>42,43</sup> As shown in Fig. 4e, the CCD value of the Li||FNCDs-SPE||Li cell (1.81 mA cm<sup>-2</sup>) is significantly higher than those of the Li||NCDs-SPE||Li (1.58 mA cm<sup>-2</sup>) and Li||P-SPE||Li cells (1.30 mA cm<sup>-2</sup>), highlighting the beneficial role of the functional fillers.

To gain deeper insights into the superior cycling stability of the FNCDs-SPE-based cells, the electrode/electrolyte interface was systematically investigated using a combination of electrochemical and post-mortem analyses. First, electrochemical impedance spectroscopy (EIS) was performed on the cycled Li||Li symmetric cells, and the data were deconvoluted using the distribution of relaxation times (DRT) method to decouple the various interfacial contributions. As shown in Fig. 5a, the Li||FNCDs-SPE||Li cell exhibits the smallest overall impedance (413 Ω). The corresponding DRT spectra (Fig. 5b) effectively separate the frequency-dependent processes: the time constant ( $\tau$ ) at around 10<sup>-5</sup> s corresponds to the ohmic resistance ( $R_b$ ) from the electrolyte and electrode components; processes with  $\tau$  ranging from 10<sup>-4</sup> to 10<sup>-2</sup> s are attributed to the SEI layer resistance ( $R_{SEI}$ ); and the  $\tau$  range of 10<sup>-5</sup> to 10<sup>1</sup> s corresponds to the charge transfer resistance ( $R_{ct}$ ).<sup>44,45</sup> Throughout the entire spectrum, the FNCDs-SPE electrolyte consistently exhibits the lowest values for both  $R_{SEI}$  and  $R_{ct}$ , indicating more stable interfacial reaction kinetics and suppressed side reactions during cycling.<sup>46</sup> This enhanced interfacial stability is directly reflected in the lithium deposition morphology. Post-mortem SEM images (Fig. 5c and d) after 100 cycles at 0.25 mA cm<sup>-2</sup> reveal a significantly smoother and more uniform lithium metal surface in the Li||FNCDs-SPE||Li cell compared to the rough, dendritic surface observed in the Li||P-SPE||Li cell, providing direct visual evidence of effective dendrite suppression.<sup>47</sup> To further understand the origin of this uniform deposition, XPS depth profiling was conducted on the cycled lithium electrodes to probe the chemical composition and thickness of the SEI layer. As shown in Fig. 5e and f and S11, the F 1s spectra reveal the presence of C-F bonds and LiF, which are typical decomposition products originating from the PVDF-HFP matrix and LiTFSI salt.<sup>48</sup> Notably, when FNCDs-SPE is used, the intensities of these decomposition components are consistently lower throughout the depth profile compared to those observed with P-SPE. This indicates that parasitic reactions between the electrolyte and lithium metal are significantly mitigated with the incorporation of FNCDs. The Li 1s spectra (Fig. 5g and h) further corroborate this finding: after 700 s of sputtering, a distinct Li metal signal at 54.5 eV appears for the Li||FNCDs-SPE||Li cell, whereas no such signal is detected for the Li||P-SPE||Li cell. This confirms the formation of a thinner SEI layer when FNCDs-SPE is employed, which is beneficial for rapid Li<sup>+</sup> transport and reduced interfacial resistance.<sup>49</sup> Additionally, the O 1s and S 2p spectra (Fig. S12) show lower contents of Li<sub>2</sub>CO<sub>3</sub>, residual LiTFSI, and Li<sub>2</sub>S in the FNCDs-SPE case, suggesting suppressed oxidative decomposition at the interface and the formation of a more uniform and robust SEI layer.<sup>21</sup> These combined results demonstrate that the FNCDs filler not only facilitates rapid ion conduction but also plays a critical role in constructing a stable and thin SEI, which collectively contribute to the exceptional cycling stability of the FNCDs-SPE-based cells.



**Fig. 5** (a) Nyquist plots of Li||Li symmetric cells with FNCDs-SPE, NCDs-SPE, and P-SPE after 100 cycles and (b) corresponding DRT analysis of the EIS spectra. SEM images of the lithium metal surface after 100 cycles in the (c) Li||FNCDs-SPE||Li cell and (d) Li||P-SPE||Li cell. XPS depth profiles of the F 1s and Li 1s spectra of (e and g) FNCDs-SPE and (f and h) P-SPE.





**Fig. 6** (a) The charge/discharge curves of the LFP||FNCDs-SPE||Li cell at different current densities. (b) The rate performance at 0.1–2C and (c) cycling performance at 0.1C of LFP||FNCDs-SPE||Li, LFP||NCDs-SPE||Li and LFP||P-SPE||Li cells. (d) The long-term cycling performance of LFP||FNCDs-SPE||Li, LFP||NCDs-SPE||Li and LFP||P-SPE||Li cells at 1C. (e) The charge/discharge curves of the LFP||FNCDs-SPE||Li cell at 1C. (f) The cycling performance of the LFP||FNCDs-SPE||Li cell under an LFP loading of 6.09 mg cm<sup>-2</sup> at 0.1C. The charge/discharge curves of the (g) NCM811||FNCDs-SPE||Li cell and (h) LCO||FNCDs-SPE||Li cell at different rates.

Full cells with the lithium metal anode and commercial cathodes were assembled to further evaluate the overall performance of the electrolyte films. The charge/discharge profiles of the LFP||FNCDs-SPE||Li cell (Fig. 6a) show a reversible specific capacity of 153.9 mAh g<sup>-1</sup> at 0.1C, which is higher than those of the LFP||NCDs-SPE||Li (145.2 mAh g<sup>-1</sup>) and LFP||P-SPE||Li cells (131.3 mAh g<sup>-1</sup>) (Fig. S13). At a higher rate of 2C, the LFP||FNCDs-SPE||Li cell exhibits the smallest capacity decay, retaining a capacity of 115.6 mAh g<sup>-1</sup>. The rate performance of the LFP||FNCDs-SPE||Li cell is shown in Fig. 6b. It delivers specific capacities of 153.9, 151.0, 145.8, 135.9, and 115.6 mAh g<sup>-1</sup> at 0.1, 0.2, 0.5, 1, and 2C, respectively. Notably, when the rate is switched back to 0.1C, the LFP||FNCDs-SPE||Li cell demonstrates better capacity recovery compared to the LFP||NCDs-SPE||Li and LFP||P-SPE||Li cells. The cycling performance at 0.1C is shown in Fig. 6c. After 100 cycles, the LFP||FNCDs-SPE||Li cell maintains a discharging capacity of 142.8 mAh g<sup>-1</sup> and a capacity retention of 95.6%, superior to that of the LFP||NCDs-SPE||Li cell (123.4 mAh g<sup>-1</sup> and 87.7%). The capacity of the LFP||P-SPE||Li cell dropped sharply after 80 cycles, indicating issues related to uneven lithium deposition/stripping. When cycled at a higher rate of 1C (Fig. 6d), the LFP||FNCDs-SPE||Li cell demonstrates excellent long-term stability, retaining 81.8% of its initial capacity after 500 cycles. In contrast, the capacities of the LFP||NCDs-SPE||Li and LFP||P-SPE||Li cells decline rapidly, indicating the outstanding interfacial stability of the FNCDs-SPE-based cell. The corresponding charge/discharge voltage profiles of the LFP||FNCDs-SPE||Li cell at different cycles are shown in Fig. 6e. Even with a higher active material loading of 6.09 mg

cm<sup>-2</sup> (Fig. 6f), the LFP||FNCDs-SPE||Li cell exhibits a discharging capacity of 126.8 mAh g<sup>-1</sup> and a capacity retention of 86.8% after 100 cycles at 0.1C, demonstrating the practical robustness of the FNCDs-SPE electrolyte.

To assess broader compatibility, the FNCDs-SPE was paired with other commercial high-voltage cathodes, such as NCM811 and LCO. As shown in Fig. 6g, the NCM811||FNCDs-SPE||Li cell delivers high reversible specific capacities of 192.9 mAh g<sup>-1</sup> at 0.1C and 134.0 mAh g<sup>-1</sup> at 2C, both substantially higher than those of the NCM811||NCDs-SPE||Li (171.6 and 90.7 mAh g<sup>-1</sup>) and NCM811||P-SPE||Li cells (142.1 and 85.8 mAh g<sup>-1</sup>) (Fig. S14). The NCM811||FNCDs-SPE||Li cell also shows the best cycling stability and rate performance among the three electrolytes (Fig. S15), attributable to the key role of FNCDs in facilitating ion transport and stabilizing the interface. Similarly, the LCO||FNCDs-SPE||Li cell (Fig. 6h) demonstrates superior cycling and rate performance compared to the LCO||NCDs-SPE||Li and LCO||P-SPE||Li cells (Fig. S16 and S17). It is noteworthy that the LCO cathode itself exhibits inferior cycling stability and rate capability retention compared to both NCM811 and LFP, which is inherently linked to the structural and chemical instability of LCO at high operating voltages. Compared with other recently reported polymer electrolytes, our FNCDs-SPE film shows competitive advantages in overall performance (Table S4).

These findings collectively demonstrate that the FNCDs-SPE electrolyte, with its rapid Li<sup>+</sup> conduction, robust mechanical properties, and excellent compatibility with various commercial cathodes, presents a promising strategy for developing high-performance, safe, and reliable solid-state batteries for practical applications.

## Conclusions

In summary, we develop fluorine–nitrogen modified carbon dots (FNCDs) and incorporated them as functional fillers to fabricate a flexible solid polymer electrolyte (FNCDs-SPE) for high-rate lithium-ion conduction. The ~2.5 nm FNCDs are uniformly dispersed within the PVDF-HFP matrix. Their abundant fluorine- and oxygen-containing surface functional groups interact with the polymer chains, effectively disrupting the chain packing and crystallinity, thereby increasing the amorphous phase content. This structural modification, coupled with the Lewis acid–base interactions between the FNCDs and Li<sup>+</sup> that provide additional hopping sites, collectively leads to a significant enhancement in ionic conductivity. The resulting rapid and uniform ion transport promotes stable lithium deposition/stripping and improves interfacial stability. Consequently, the FNCDs-SPE outperforms both the NCDs-SPE and P-SPE in long-term cycling and rate capability. Furthermore, the FNCDs-SPE demonstrates excellent compatibility with various high-voltage commercial cathodes (LFP, LCO, and NCM811), highlighting its great promise for practical applications. This work provides a valuable strategy for the design of high-



performance, safe, and reliable solid-state battery electrolytes through functional nanofiller engineering.

## Author contributions

Yang Wu: investigation, experiment, writing – original draft, and data curation. Wenbo Yue: writing – review and editing, supervision, project administration, and conceptualization.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The authors confirm that the data supporting the findings of this study are available within the article and its supplementary information (SI). Raw data that support the findings of this study are available from the corresponding author, upon reasonable request.

Supplementary information is available. See DOI: <https://doi.org/10.1039/d6lf00021e>.

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