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Structured for success: conjugated polymer binders with tailored composition and architecture for lithium-ion batteries

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Lithium-ion batteries (LIBs) are the leading energy storage technology, yet enhancing their energy density and cycle life remains critical. Significant progress has been made in high-capacity anodes and high-voltage cathodes, but their performance is hindered by electrode degradation, where it is related to the behaviors of binders at the surface and interface. Conventional non-conductive binders like poly(vinylidene difluoride) (PVDF), combined with conductive additives, often fail to maintain electrical pathways under repeated volume changes. Alternatively, conjugated polymer binders have emerged as a superior alternative, simultaneously offering intrinsic conductivity, mechanical flexibility, and strong adhesion through π -conjugated backbones and functional groups. Their tunable molecular structure enables efficient electron/ion transport while mitigating electrode cracking. Additionally, the development of hierarchically ordered nanostructures in conjugated polymer binder can further enhance their electrochemical performance. This review examines the design principles of conjugated polymer binders, focusing on molecular engineering and nanostructural control to optimize their performance in high-loading electrodes, such as silicon-based anodes. By addressing key challenges in binder functionality, these advanced materials pave the way for next-generation high-energy-density LIBs.

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Broader context

Here, we explore a crucial yet often underappreciated component in lithium-ion battery (LIB) development: electrode binders. While significant advancements have been made in high-capacity anodes (e.g., silicon) and high-voltage cathodes, electrode degradation continues to limit the realization of next-generation batteries with high energy densities, long-term cycling stability and stable calendar life. Traditional non-conductive binders often fail to accommodate the severe volume fluctuations and interface instabilities during cycling, leading to loss of electrical connectivity and to worsened parasitic reactions. In contrast, conjugated conductive polymer binders present a promising path forward. This review critically evaluates recent progress in their molecular design and nano-structural engineering. Distinct from broader polymer binder reviews, our minireview concentrates specifically on conjugated systems, with an emphasis on the development of ordered molecular architectures and well-defined nanostructures. These features are shown to play a pivotal role in improving electrochemical performance by forming robust and conductive polymer networks.

1. Introduction

Lithium-ion batteries (LIBs) have become the predominant energy storage technology, powering a wide range of applications ranging from consumer electronics, electric vehicles to grid storage.¹ To address the increasing need for sustainable energy storage, it is essential to improve the energy density of electrodes. In the past 20 years, substantial advancements have been achieved in the development of high-capacity anodes² (e.g., silicon-based materials) and high-voltage

cathodes^{3–5} (e.g., nickel-rich layered oxides (NMC) and lithium-rich layered oxides), offering promising solutions to LIBs with higher energy density. Nevertheless, the efficiency and cycle life of these batteries are strongly limited by the properties of electrode materials and their interfaces,⁶ where binders play an essential role in maintaining structural stability, materials integrity and electrical connectivity of electrodes.^{5,7,8}

In traditional electrode fabrication, insulating binders like poly(vinylidene difluoride) (PVDF) are commonly combined with conductive carbon additives, such as acetylene black.⁹ However, this configuration often struggles to preserve the required conductive pathways in the electrode, especially when subjected to substantial volume changes and pulverization during cycling.¹⁰ PVDF itself has negligible intrinsic conductivity, and it blocks ion/electron pathways in electrodes, par-

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ticularly in thick or high-loading designs (e.g., silicon anodes).¹¹ Meanwhile, acetylene black particles rely on physical contact for electron percolation, which could be easily disrupted by volume changes, therefore isolating active material particles. In this scenario, PVDF cannot rebuild the electrical contacts after fracture, due to its lack of self-healing or dynamic bonding.¹² In addition, the expansion and contraction of the high-capacity alloying type anode particles generate immense mechanical stress, causing the typically brittle solid electrolyte interphase (SEI) to crack and rupture. This rupture exposes fresh, highly reactive lithiated silicon surfaces to the electrolyte, triggering a destructive and continuous SEI reformation. Studies have shown that with a passive binder like PVDF, this process results in the formation of a thick, non-uniform, and mechanically unstable SEI dominated by organic carbonate species, which is particularly prone to failure.¹³ As a result, the continuous parasitic reactions at the electrode–electrolyte interfaces reduced the calendar life of the anodes through two main degradation modes: the irreversible consumption of the finite lithium inventory and a gradual increase in cell impedance.¹⁴

Conjugated polymer binders have emerged as a promising alternative, combining intrinsic electronic conductivity, mechanical resilience, and superior adhesion to electrode materials. Their unique π -conjugated backbones (or side chains) allow electron delocalization, creating uninterrupted conductive networks that enable efficient electron transport across the entire electrode structure (Fig. 1A).^{15,16} The flexible molecular nature balances rigidity and elasticity, reducing mechanical stress and preventing electrode cracking or pulverization, thereby enhancing cycle life. Conjugated polymers also form strong intermolecular interactions (e.g., π - π stacking,

hydrogen bonding, or electrostatic interactions) with active materials, preventing particle detachment during cycling.¹⁷ The conjugated polymer binder also function as a powerful coating, which limits the direct contact between the bulk electrolyte and the highly reactive anode surface, inherently suppressing the extent of parasitic side reactions.¹⁸ Additionally, functionalization with polar groups (e.g., hydroxyl, ether, carboxyl or (trifluoromethylsulfonyl)imide (TFSI) groups) can simultaneously enhance lithium-ion transport.^{19,20} It was reported that conjugated polymer binder preferentially interact with the electrolyte salt (e.g., LiPF_6), which accelerates the decomposition of the salts to form SEI based on LiF and Li_xPF_y .^{21,22} Those thinner, inorganic-rich SEI are highly desirable, known for its high interfacial stability, mechanical robustness, and excellent passivation properties.

To improve the durability and performance of high-loading electrodes (such as Si and SiO_x), conjugated polymer binders must be engineered with appropriate organic moieties and functional groups within a carefully designed framework. Fine-tuning the polymer's molecular arrangement significantly affects key properties, including electronic structure, electrolyte absorption, mechanical response, and ease of processing.^{23,24} In addition to the chemical nature of individual polymer molecules, the macroscopic packing behavior of the polymers also plays a crucial role. For example, it has been found that hierarchically ordered structures (HOS) with well-defined nanocrystalline morphologies in the binder can enhance both electronic and ionic conductivity (Fig. 1B).²² This review will systematically explore the chemical design principles of conjugated polymer binders, focusing on the relationship between their molecular structures and their functionality in lithium-ion batteries. Special emphasis is placed

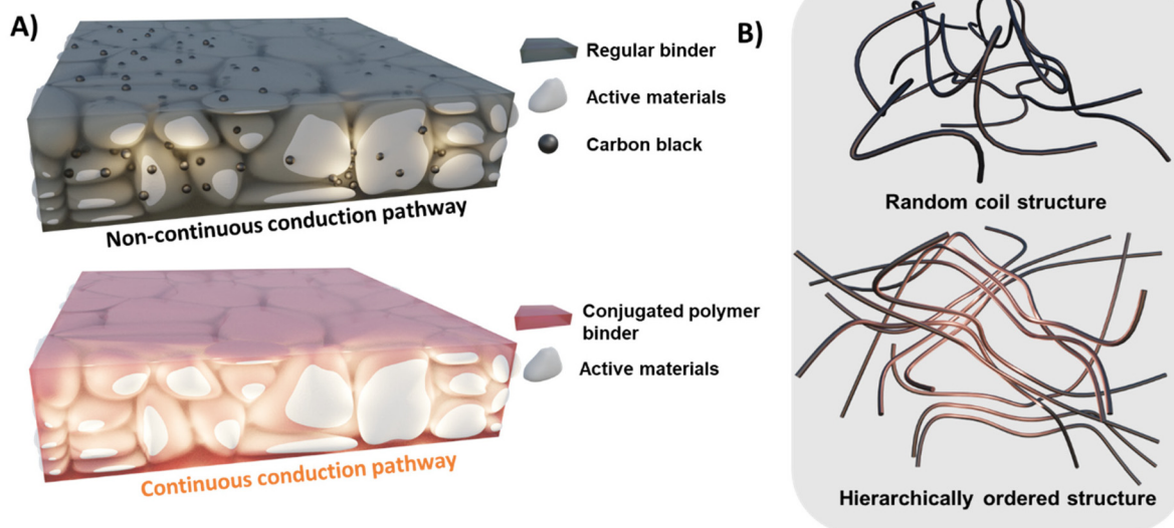


Fig. 1 (A) Illustration of the classical electrode design (based on PVDF and carbon black) featuring non-continuous conduction pathway, as well as the electrode with conjugated polymer binder featuring continuous conduction pathway. (B) Illustration of conjugated polymer with random coil structure or hierarchically ordered structure with improved crystallinity.



on strategies for constructing well-defined nanostructures to enhance their electrochemical performance.

2. Structural design of conjugated polymer binders and manipulation of π -conjugation and π - π stacking

2.1. Electronic conductivity modulation

Adjusting the chemical composition of the polymer backbone is crucial for modifying its electronic structure to enable practical use as a multifunctional binder in lithium-ion batteries. The choice of the repeating monomer units that constitute the conjugated backbone is the most direct way to define the polymer's inherent electronic characteristics, including its highest occupied molecular orbital (HOMO), lowest unoccupied molecular orbital (LUMO) energy levels and band gap. Different aromatic or heteroaromatic systems (*e.g.*, thiophenes, fluorenes, phenylenes, pyrroles, quinones) offer distinct electronic profiles. Importantly, undoped conjugated polymers typically have low conductivity (10^{-10} to 10^{-5} S cm⁻¹), which can be substantially increased by magnitudes *via* electrochemical or chemical doping.²⁵ This suggests that the intrinsic characteristics of the polymer backbone critically influence the level of conductivity attainable in the material.¹⁷

A critical consideration, often overlooked in literature, is that this electronic conductivity is not universally persistent. It manifests only when the polymer is held within its specific electrochemical doping window. This necessitates a crucial alignment between the polymer's redox potential and the electrode's actual operating potential. For instance, a polymer designed for a cathode binder must be able to sustain a stable p-doped (oxidized) state at high operating potentials (*e.g.*, 3.0–4.5 V *vs.* Li/Li⁺). Conversely, for a binder in a low-potential anode (such as silicon, operating <1.0 V *vs.* Li/Li⁺), the polymer is only functionally conductive if it can be electrochemically reduced (n-doped) and remain in that conductive state at these low potentials.

Towards p-doping, incorporation of protonic acid is one of the most widely used methods. Certain conjugated polymers such as polyaniline (PANI) allow protonation of the imine nitrogen, converting the emeraldine base (semiconducting) form to the emeraldine salt (conducting) form.²⁶ This process involves quinoid-benzenoid resonance stabilization, making the charged state very stable. Different organic and inorganic acids such as polyacrylic acid,²⁷ anthranilic acid,²⁸ phytic acid^{29,30} and sulfuric acid³¹ 5-sulfo isophthalic acid³² have been demonstrated as excellent dopants for polyaniline, enabling practical use as binder for various anodes and cathodes. Additionally, conjugated polymer binder could also undergo electrochemical doping during the battery operation, towards promoted conductivity.³³ For example, the electron-donating nature of the oxygen atoms in the ethylenedioxy bridge of poly(3,4-ethylenedioxythiophene) (PEDOT) lowers the overall band gap of the polymer and, crucially, reduces its ox-

idation potential. This makes PEDOT easier to p-dope. Thompson *et al.* quantified the electronic conductivity evolution of a PEDOT derivative, dihexyl-substituted poly(3,4-propylenedioxythiophene) (PProDOT-Hx2) as a function of p-doping level. They observed that the conductivity increased from $\sim 10^{-5}$ S cm⁻¹ to 10^{-1} S cm⁻¹ at ~ 3.6 V, coinciding with the complete conversion of polarons to bipolarons (Fig. 2A).²⁵ Higgins *et al.*, came up with an *in situ* secondary doping of PEDOT: polystyrene sulfonate (PEDOT:PSS) with small amounts of formic acid (5 and 10 wt%) for Si anodes. Upon doping the electrical conductivity close to 4.2 S cm⁻¹ was observed with Si nanoparticles electrodes. The system delivered a specific capacity of 3865 mA h g⁻¹ at 1 A g⁻¹ and could retain 75% after 100 cycles. Furthermore, at higher electrode thickness of 1.5 mg cm⁻², a high areal capacity of 3 mA h cm⁻² was observed for the 10 wt% formic acid doped PEDOT:PSS/Si electrodes.³⁴

Unlike p-doping, n-doping presents greater challenges because most π -conjugated systems are inherently electron-rich, leaving few classical conjugated polymers naturally suited for efficient n-doping.⁷ Effective n-doping generally requires polymers with low electron affinity to facilitate electron transfer from the dopant. Typical conjugated polymers based on polythiophene and polypyrrole derivatives have been applied as anode binder with some n-doping potential, such as poly[3-(potassium-4-butanoate)thiophene] (PPBT),^{35,36} poly[3-(2-ethylhexyl)thiophene-*alt*-2,5-terephthalic acid sodium salt] (PT1T).³⁷ However, to enhance n-doping performance, introducing more electron-withdrawing heteroatoms (*e.g.*, N, O, or S) into the backbone is suggested as it can help delocalize electron density and improve electron-accepting capability.³⁸ Jiang *et al.* developed a highly conductive n-type polymer binder for silicon anode, using polyoxadiazoles (PODs).^{39,40} The strong electron-withdrawing nature of the oxadiazole ring endows PODs with low LUMO energy levels, facilitating n-doping and enabling superior electron transport properties (Fig. 2B). As a result, the capacity of silicon microparticles (4200 mAh g⁻¹) can be almost entirely released during the half-cell testing. Polyisoidigo is another important class of n-type conducting polymer which works well at lower operating potentials and hence was employed as conducting additive cum binder for Si@C core shell anodes. The attractive mechanical and electrochemical properties of polyisoidigo makes it a promising n-type binder for Si anodes. The Si@C electrodes with polyisoidigo binder displayed high-capacity of 1400 mA h g⁻¹ and maintains a stable performance over 500 cycles with average coulombic efficiency of 99.56%.⁴¹ Similarly, Matsumi *et al.* reported a bis(imino)acenaphthene quinone (BIAN)-type conjugated polymer, the delocalized lone pair of electrons on the nitrogen atoms enabled low-lying LUMO level of the binder (-3.69 eV), endowing its great potential to be n-doped.⁴²

The same group designed and synthesized BIAN-fluorene copolymer binder for graphite anode to enhance the electronic conductivity and to prevent the continuous degradation of SEI. The copolymer has lower lying LUMO level than that of ethylene carbonate thus preventing continuous degradation of elec-



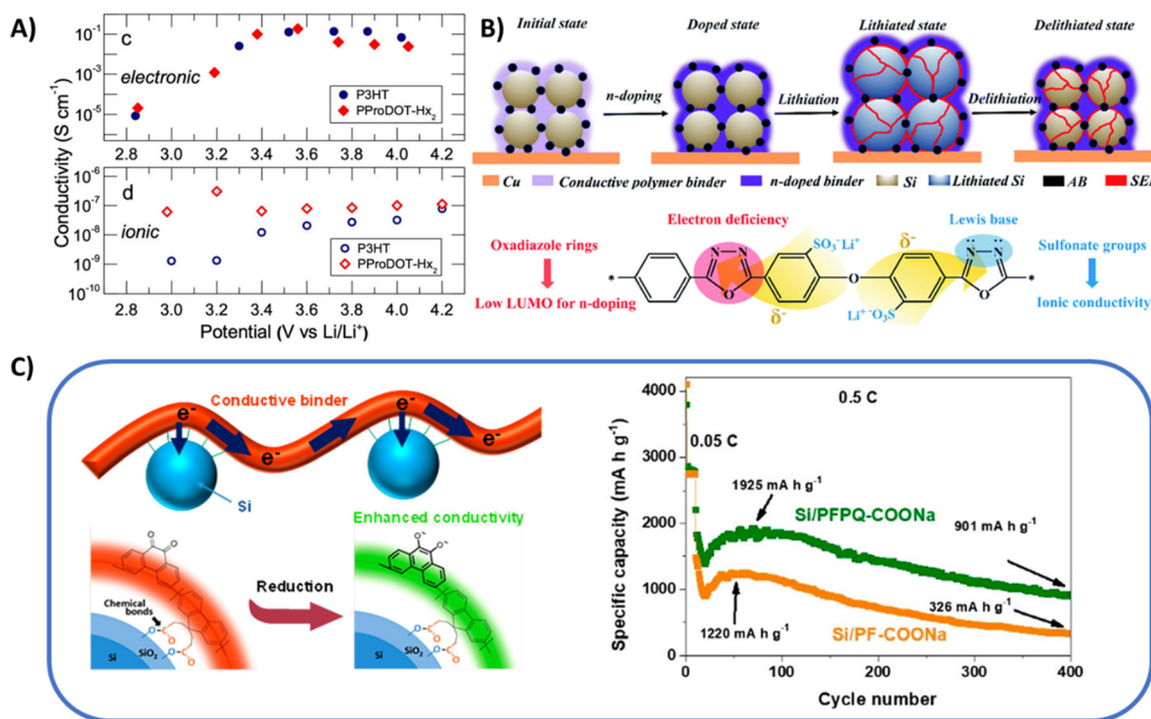


Fig. 2 (A) Electronic and ionic conductivities of poly(3-hexylthiophene) (P3HT) and PProDOT-Hx₂ at different p-doping state. Reprinted by permission from American Chemical Society, ref. 25, copyright 2020. (B) n-type conjugated polymer binder that maintains the electrical and mechanical integrity of the Si microparticle anode during charge and discharge cycles, as well as chemical structure of aromatic polyoxadiazole lithium sulfonate. Reprinted by permission from The Royal Society of Chemistry: ref. 39, copyright 2021. (C) Schematic illustration of the electrochemical behavior of PFPQ-COONa binder in Si anode, and cycling performance of Si/PF-COONa and Si/PFPQ-COONa at 0.5 C. For all measurements above, the areal loading of the active material was approximately 1.2 mg cm⁻². Reprinted by permission from American Chemical Society, ref. 43, copyright 2018.

trolyte during charge–discharge process. Furthermore, the HOMO and LUMO energy levels are in a position where the polymer gets doped well before the reduction of carbonate solvents of the electrolyte thus maintaining electronic conductivity. On the other hand, PVDF has LUMO energy level lying well above that of ethylene carbonate thus resulting in continuous degradation of EC and breakdown of SEI in each charge–discharge cycle. Thus, the BIAN-fluorene copolymer displayed better electrochemical performance compared to PVDF for graphite anode.⁴⁴

Another effective strategy to enhance n-doping performance involves integrating donor/acceptor moieties, such as electron-withdrawing carbonyl groups, into the polymer backbone. This approach is particularly practical since polycondensation synthesis naturally allows for the incorporation of multiple monomer types. This is exemplified by the series work by Liu *et al.*, as introducing fluorenone units into the polyfluorene backbone allows the polymer to undergo cathodic doping in the reducing lithium environment, enhancing its overall electrical conductivity by magnitudes.^{23,45,46} Pan *et al.* reported a similar design by introducing 10% of phenanthraquinone (PQ) into polyfluorene backbone (Fig. 2C). The produced copolymer (PFPQ-COONa) is successfully applied as binder for Si anode, delivering 76.1% capacity retention after 200 cycles (0.2 C).⁴³

While the extended conjugated backbone primarily dictates the fundamental electronic characteristics of conjugated polymers, the moieties located on the polymer side chains also plays an important role in influencing the polymer's electrochemical characteristics. Sizable side chains can cause steric interference, hinder tight packing and weakening intermolecular interactions among polymer chains. This may prevent the development of planar backbone conformations, lowering crystallinity and, as a result, diminishing electrical conductivity. Additionally, the bulky side chains can induce torsional strain along the polymer backbone, disrupting its planarity.⁴⁷ Such structural distortion may hinder π -electron delocalization, which is essential for effective charge transport in conjugated polymer binders.⁴⁸ For example, poly(9,9-dioctylfluorene-*co*-fluorenone-*co*-methylbenzoic ester) (PFM), which features dioctyl side chains, exhibited an electronic conductivity of only 10⁻⁶ S cm⁻¹.²² In contrast, a polyspirobifluorene-based binder with similar backbone but more compact side chain structure achieved a significantly higher conductivity of 4.7 × 10⁻³ S cm⁻¹.⁴⁹

Interestingly, bulky side chains do not always hinder the electronic conductivity of polymer binders, as they can also be conjugated and enhance long-range polaron mobility. For instance, Liu *et al.* developed a class of conductive polymers



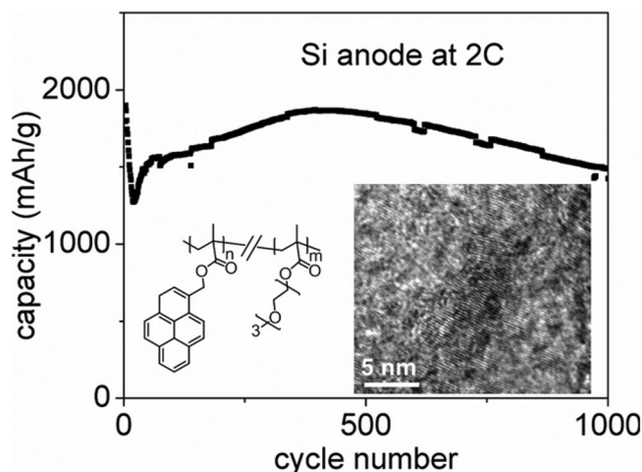


Fig. 3 Structure of the side chain conducting poly(1-pyrenemethyl methacrylate-co-triethylene oxide methyl ether methacrylate) (PPyE), high-resolution transmission electron microscopy (HRTEM) image of PPyE polymer, and charge (delithiation) capacities of PPyE-based Si electrodes at 2 C rate. Reprinted by permission from American Chemical Society, ref. 50, copyright 2015.

with π - π stacking side chains to facilitate electron transport (Fig. 3). By incorporating pyrene-based moieties *via* copolymerization, they achieved polymer binders with electron mobilities up to $8.5 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. These binders preserved electrode integrity and maintained stable Si interfaces over 1000 cycles, enabling high-capacity batteries with superior rate performance, thanks to their self-assembled nanostructures.

2.2. Polymer flexibility modulation

Rational design of conjugated polymer backbone is important for tuning their flexibility, as inherent rigidity of some classical conjugated polymers (*e.g.*, polyacetylene) limits their use as binders, and they are not resistant to cycle-induced large strain. Incorporating flexible segments (*e.g.*, introducing methyl benzoate ester blocks into the polyfluorene backbone²³) addresses this brittleness, enabling stretchability and deformability. Despite these advantages, this modification comes at the cost of disrupted conjugation and hindered chain packing, ultimately compromising the polymer's crystallinity and electrical conductivity. Thompson *et al.* investigated conjugated break spacers (CBS) in PProDOT cathode binders for Li-ion batteries.⁵¹ Increased CBS content and spacer length reduced lamellar peak intensity, consequently lowering charge carrier mobility. These results aligned with rate capability tests of the lithium nickel cobalt aluminum oxide (NCA) cathodes with different PProDOT binder, where electronic conductivity (over mechanical properties) governed performance and polymers with higher conductivity retained more capacity (Fig. 4A).

The ideal case is to establish 2D/3D-conjugated conductive network with favorable flexibility without compromising the continuous π -conjugation. In a systematic comparison of polyaniline-based binders with different architectures including star-like (s-PANI), cross-linked (c-PANI), and linear (l-PANI)

(Fig. 4B), the silicon anode with s-PANI binder delivered a reversible capacity of 1776 mAh g^{-1} after 100 cycles at 500 mA g^{-1} , significantly outperforming systems using c-PANI or l-PANI. It was found s-PANI/Si exhibited a much lower electrode thickness expansion (114%) than that of c-PANI/Si (286%).⁵² Similarly, Seferos *et al.* reported a 2D-conjugated polysulfonamide as binder for organic lithium-ion batteries.⁵³ The findings indicate that the linear (1D) oligosulfonamide binder, lacking branching structures, exhibits weak mechanical properties and fails to ensure long-term stability in perylene-tetracarboxylic dianhydride-based composite electrodes. In contrast, the 2D-conjugated polysulfonamide binder demonstrated no delamination or discoloration during cycling. Additionally, in rate performance tests, the electrode with the 2D-conjugated binder retained twice the capacity compared to the one with the 1D oligosulfonamide binder.

2.3. Polymer solvation modulation

A key requirement for scalable electrode manufacturing is satisfactory slurry processibility, which is governed by rheology. This presents a unique challenge for conjugated polymer binders. Without solubilizing side chains, their rigid conjugated backbones cause significant intermolecular aggregation, leading to dramatically high and uncontrolled viscosity.⁵⁴ This problem cannot be solved by simply reducing the binder content, as is common in conventional non-conductive binder systems; here, the conjugated polymer dominates the conductive network, and maintaining its concentration is critical for electrode conductivity.

Therefore, molecular engineering *via* "internal plasticization" is the preferred pathway. The side chains are the primary tool for tuning intermolecular interactions and solubility of conjugated polymers. The target is to achieve an ideal shear-thinning profile: a high viscosity or yield stress at low shear rates (at rest) to suspend components and prevent active material settling, followed by a sharp decrease in viscosity under the high shear rates of coating.⁵⁵ These side chains achieve this dual function in two ways. First, long, bulky, or branched side chains provide crucial steric hindrance, shielding the rigid backbones and preventing them from stacking into aggregates, especially as they align under shear flow. Second, these same flexible side chains enhance the degree of polymer entanglement at rest. This entanglement creates a high initial (zero-shear) viscosity, which in turn leads to a more pronounced shear-thinning effect as these entanglements break and the chains align under shear.

So far, the most commonly applied side chain is still the alkyl groups based on hydrocarbons. Liu *et al.* inaugurally developed the polyfluorene-type polymers with tailored side chain structures as conductive binders for silicon anode applications.⁴⁵ The polyfluorene polymers containing octyl side chains enhance the polymer's solvation in non-polar solvents, therefore the conductive polymer matrix is well-suited for the lithium-ion slurry manufacturing process. The cost-effective silicon/polymer composite anodes demonstrate exceptional cycling performance, delivering approximately 2100 mAh g^{-1}



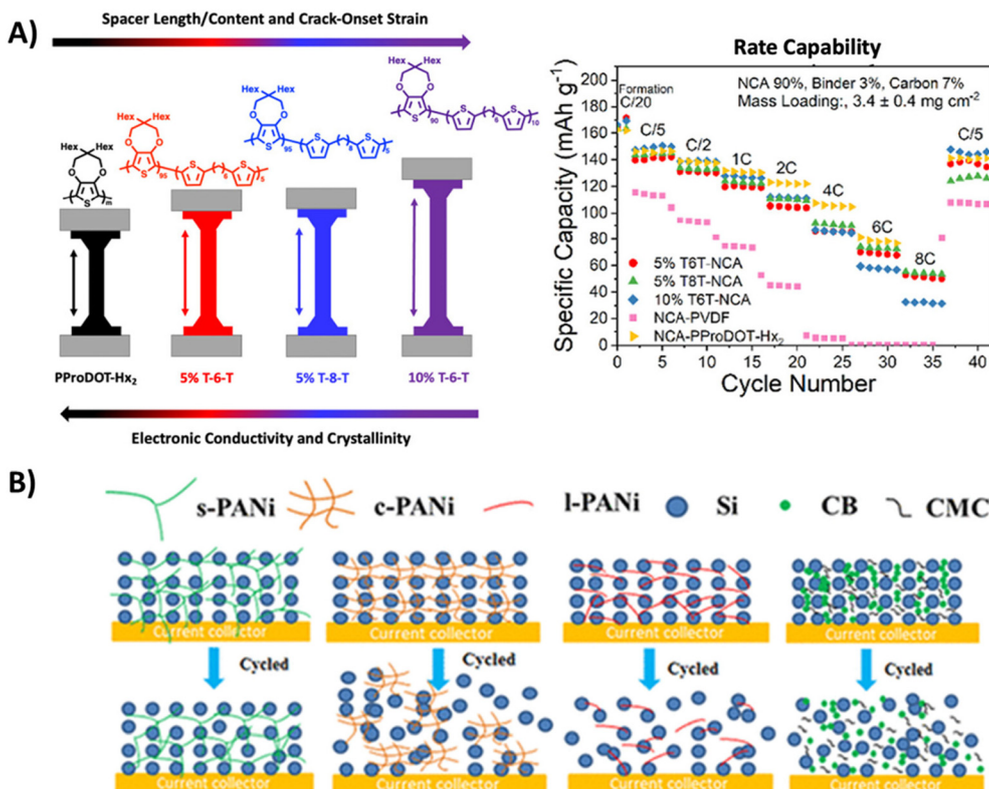


Fig. 4 (A) Left: Relationship between the spacer length, crack-onset strain, electronic conductivity and crystallinity of the conjugated polymer with different conjugation break spacer. Right: Rate capability testing for the Li-NCA-5% T-6-T, Li-NCA-5% T-8-T, Li-NCA-10% T-6-T, Li-NCA-PProDOT-Hx₂, and Li-NCA-PVDF cells with a cathode mass composition of 90% NCA, 3.5% Super P, 3.5% CNT, and 3% binder. Two formation cycles at C/20 were carried out before testing. Reprinted by permission from American Chemical Society, ref. 51, copyright 2024. (B) Schematic illustrations of cycle-induced structural changes of s-PANI/Si, c-PANI/Si, l-PANI/Si, and carboxymethyl cellulose (CMC) electrode systems. Reprinted by permission from American Chemical Society, ref. 52, copyright 2020.

for silicon and 1400 mAh g⁻¹ for the full electrode, even after 650 cycles—all without the need for conductive additives.

Compared to the hydrophobic side chains, the hydrophilic side chains (e.g., oligo ethylene glycol, carboxylic acid) could also enhance the polymer processability. More importantly, they also create ion-permeable pathways and improves ionic conductivity.²³ It was demonstrated that the enhanced polarity of the PEFM polymer binder could result in a 3-fold electrolyte uptake improvement, compare to the PFM binder without oligo(ethylene glycol) side chains (Fig. 5A). Thompson *et al.* conducted a systematic study on the role of oligoether side chains in controlling solvent-induced swelling and electrolyte absorption.⁵⁶ They designed a series of binders derived from dihexyl-substituted poly(3,4-propylenedioxythiophene) (PProDOT-Hx₂), in which 5% to 35% of the hexyl groups were replaced with oligoether chains (Fig. 5B). Their results showed that as the oligoether content increased from 5% to 35%, the binder's swelling ratio rose from 41% to 97%. Simultaneously, the ionic conductivity improved fourfold, reaching a maximum value of 4×10^{-7} S cm⁻¹. However, it should be noted that the combination of hexyl and oligoether side chains increased structural disorder in the binder, leading to a decline in electronic conductivity. The binder with 5% oligoether exhibited

the highest electronic conductivity (1.1 S cm⁻¹), while the 35% oligoether variant showed the lowest (0.14 S cm⁻¹). Grazing-incidence wide-angle X-ray scattering (GIWAXS) analysis confirmed this trend, revealing reduced crystallinity in the doped polymer binders with higher oligoether fraction. As an optimal compromise, the (75 : 25) PProDOT binder that incorporating 25% oligo(ethylene glycol) side chains demonstrated the best rate capability and fastest charge/discharge kinetics in symmetric testing.

3. Structural reordering to enhance conjugated polymer binder performance

3.1 Side chain cleavage

A more specialized and innovative strategy involves designing conjugated polymer binders with labile side chains. After slurry-based processing step, these specific side chains are intentionally cleaved and removed from the main conjugated backbone. The removal of these bulky solubilizing groups reduces steric hindrance and allows the conjugated backbones



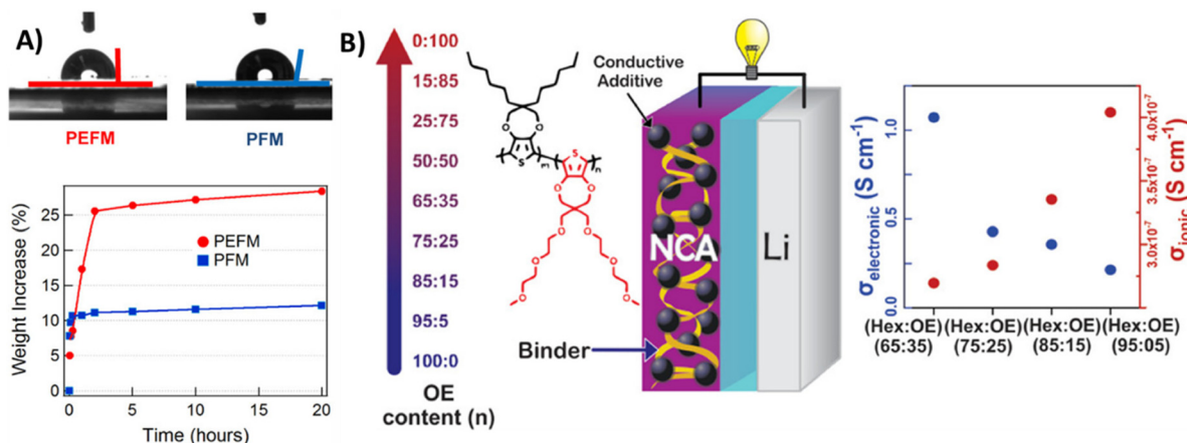


Fig. 5 (A) Water contact angles on the surfaces of PEFM and PFM films, and the swelling tests of PEFM and PFM polymer film in the 1 M LiPF₆ ethylene carbonate (EC)/diethyl carbonate (DEC) (1 : 1) electrolyte. Reprinted by permission from American Chemical Society, ref. 23, copyright 2013. (B) Electronic and ionic conductivities of the PProDOT copolymer series with different amount of oligoether side chains. Reprinted by permission from American Chemical Society, ref. 56, copyright 2022.

to pack more closely. This enhanced proximity facilitates stronger intermolecular π - π stacking, leading to a significant improvement in electronic conductivity.⁵⁷ This method cleverly addresses the common trade-off where side chains necessary for good solubility (and thus processability) can impede optimal backbone packing for conductivity. For instance, Ponder *et al.* developed a carefully designed PProDOT-based polymer featuring cleavable 2-butyloctyl ester side chains.⁴⁷ Following standard solution processing, the polymer films were treated with a basic solution to hydrolyze the side chains, converting them into an insoluble, solvent-resistant material with tight π - π stacking. This structural rearrangement led to a dramatic rise in electronic conductivity, from 6 S cm⁻¹ to 60 S cm⁻¹.

However, the side chain cleavage process does not always require intricate molecular design. Liu *et al.* proposed that thermal-induced side chain cleavage could be a broadly applicable method, as demonstrated with polyfluorene-based polymer binder. Polyfluorene exhibits high charge mobility, but only in a well-ordered β -phase conformation.^{58,59} Unfortunately, this phase usually exists as isolated domains within a predominantly amorphous and insulating film. To optimize the chain packing for creating a percolating network of highly conductive phase, their study showed that heating poly(9,9-dioctylfluorene-*co*-fluorenone-*co*-methylbenzoic ester) (PFM) to 500 °C selectively removed its octyl side chains while preserving the polymer backbone (Fig. 6A and B).²² This transformation facilitated the development of hierarchically ordered structures (HOS) (Fig. 6C), significantly enhancing conductivity from 10⁻⁶ S cm⁻¹ to 0.1 S cm⁻¹. Notably, the HOS design also enabled rapid lithium-ion diffusion, comparable to that in graphite. Unlike conventional ion transport mechanisms, which rely on segmental polymer motion or solvated ions, the HOS conductive polymer couples lithium-ion movement with negative polaron (electron) diffusion, establishing it

as a novel soft material for efficient ion transport. As a demonstration of its potential as a lithium-ion battery binder, the conductive HOS-PFM facilitated stable cycling and excellent capacity retention in high-loading silicon-based anodes (4.5 mAh cm⁻²).

Subsequently, this approach was adapted for practical applications using the water-soluble polyfluorene binder poly(2,7-9,9-di(oxy-2,5,8-trioxadecane))fluorene) (PFO). Thermal treatment at 400 °C removed the oligo(ethylene glycol) side chains, and wide-angle X-ray scattering (WAXS) and X-ray diffraction (XRD) analyses revealed enhanced π - π stacking of the polymer backbone. The interchain spacing (*d*-spacing) decreased from 4.3 Å to 3.4 Å, which is close to double of the van der Waals radius of carbon and approaching ideal molecular packing. The resulting HOS-PFO displayed a six-order-of-magnitude increase in conductivity compared to pristine PFO, enabling stable cycling of high-loading SiO_x anodes. Notably, the SiO_x-HOS-PFO electrodes achieved ~86% capacity retention over 200 cycles at 0.33 C, even without conductive carbon additives.⁶⁰ Furthermore, HRTEM analysis demonstrated that the HOS-PFO maintained its crystallinity even after repeated cycling, highlighting the exceptional stability of the HOS frameworks (Fig. 6D and E).

3.2 Introduction of secondary polymer

Polymer blending can promote the reorganization of conjugated polymer, leading to well-ordered nanostructures with enhanced electrochemical performance. This approach has been extensively used in organic electronic device fabrication. For instance, combining conjugated polymer with secondary insulating polymer can greatly improve the π - π stacking of the conjugated polymers and therefore enhance the charge carrier mobility, owing to the nanoconfinement effect.^{61,62} This design principle also applies to the formulation of conjugated polymer-based binder for lithium-ion batteries. A well-studied



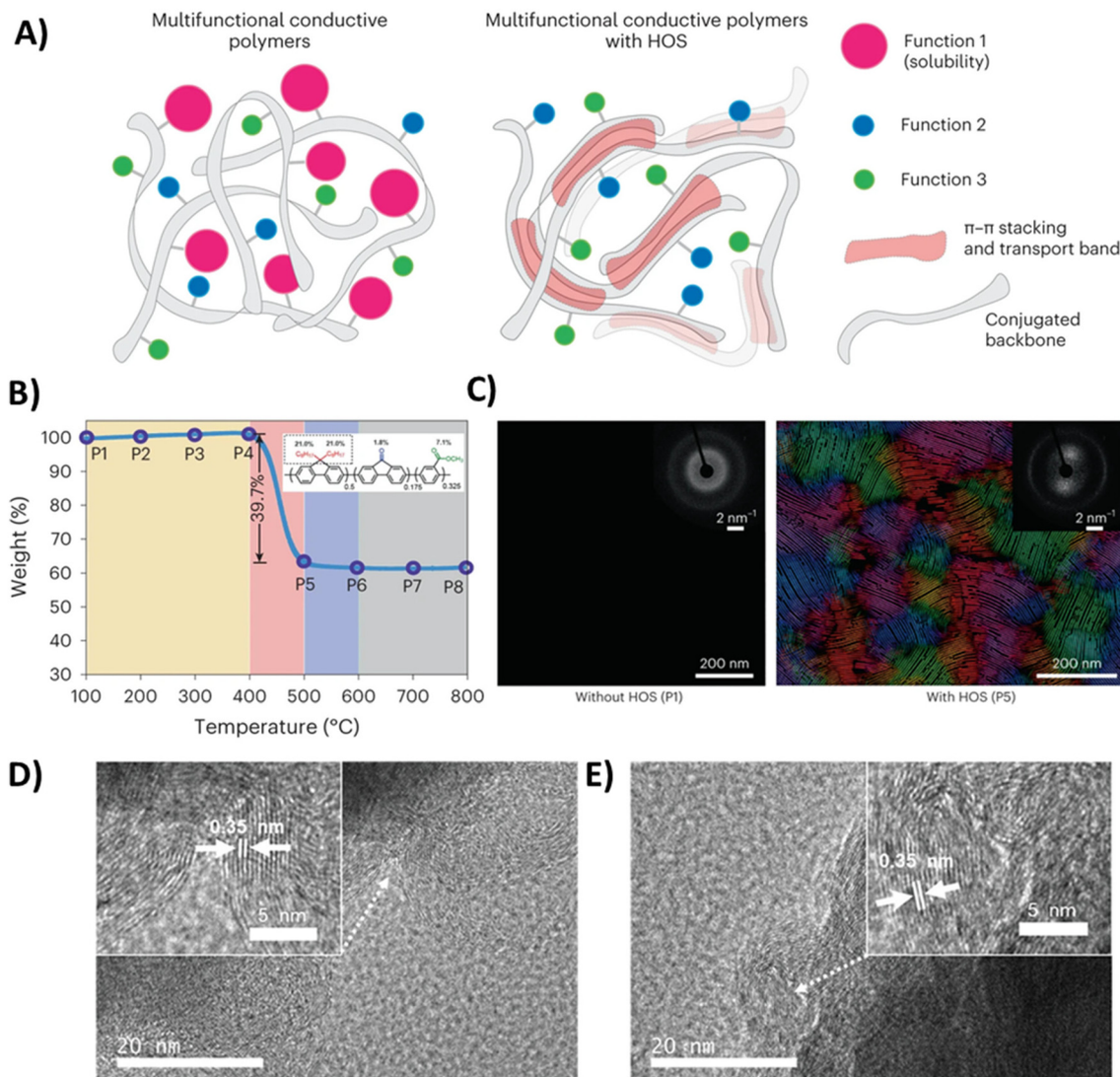


Fig. 6 (A) Schematic illustration of polymer chain arrangements in multifunctional conductive polymers and multifunctional conductive polymers with HOS. Reprinted by permission from Springer Nature, ref. 22, copyright 2023. (B) Thermalgravimetric analysis of PFM. Samples are named P1–P8 based on the highest thermal processing temperature (100 to 800 $^{\circ}\text{C}$). Tan shading: thermally stable region for PFM; pink shading: alkyl side-chain cleavage; purple shading: losing carbonyl groups; grey shading: carbonization region. Reprinted by permission from Springer Nature, ref. 22, copyright 2023. (C) Flow line maps tracking the polymer main chains and examples of diffraction patterns (insets, weighted averages over 3×3 pixels) of PFM and HOS-PFM acquired by 4D-STEM. The color wheel represents the orientation of the main chain of the polymer. Reprinted by permission from Springer Nature, ref. 22, copyright 2023. (D and E) High-resolution TEM image of SiO_x -PFO-aq-400 before cycling (D) and after being cycled 100 times (E). Reprinted by permission from Wiley, ref. 60, copyright 2023.

case involves blending PEO (polyethylene oxide) with PEDOT:PSS as a polymer binder. Beyond boosting ionic conductivity, polyethylene glycol (PEG) also enhances PEDOT chain ordering, leading to improved electronic conductivity.⁶³ Kang *et al.* demonstrated that the hydrogen bonding between PEG and PSS promotes phase separation, leading to a more linear and densely packed PEDOT conformation with increased crystallinity (Fig. 7A), as verified by GIWAXS analysis (Fig. 7B).⁶⁴ At an optimal PEDOT:PSS/PEG ratio of 1 : 0.3, the composite binder exhibited a 72-fold increase in electronic conductivity alongside improved mechanical properties. When applied in a silicon anode, this binder enabled a high-rate capacity of

500 mAh g^{-1} at 2 C and full capacity recovery to $\sim 2300 \text{ mAh g}^{-1}$ at 0.1 C, showcasing excellent cycling stability. Compatible with PEDOT:PSS, CMC serves as an additional water-soluble polymer option beyond PEG. An *et al.* documented analogous conductivity enhancements using CMC, attributing them to charge-screening effects that cause phase separation and reorientation of PEDOT chains.⁶⁵ Similarly, Song *et al.*, developed a mixed ion/electron conducting binder for anodes by introducing ionically conducting PEO branches to an electronically conducting polythiophene with a redox active pendant. The ionically conducting branches of PEO were controlled to understand the contributions of electronic and ionic conductivity to



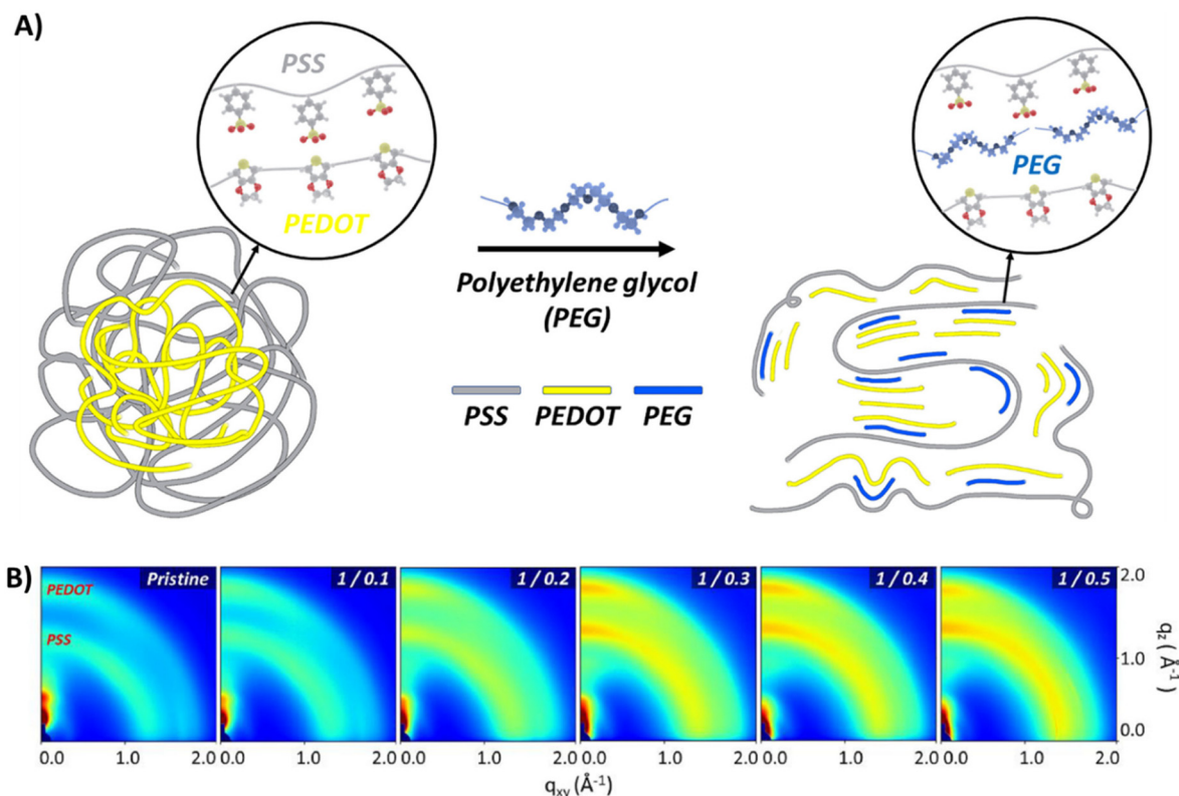


Fig. 7 (A) Proposed mechanism of PEG in PEDOT:PSS. (B) 2D GIWAXS patterns of PEDOT:PSS films according to the PEG blending ratio. Reprinted by permission from Elsevier, ref. 64, copyright 2024.

LIB performance. This redox active binder when employed as a binder, the rate performance and the storage capacity was significantly improved when compared to that of PVDF. The polymers with simple octyl side chains were also prepared as a control which displayed poor electrolyte wettability and hence lower capacity than that of PEO comprising polythiophene binder.

Building upon the concept of controlling phase separation through physical blending, a more delicate strategy involves engineering the nanostructure *via* chemical synthesis to create multifunctional composite binders. A prominent example is the core-shell aqueous conductive binder (PABS:PEDOT) for sulfur cathode, recently reported by Wang *et al.*⁶⁶ They first synthesized a sulfonate-containing elastic copolymer (PABS) as the “core” *via* emulsion polymerization, followed by *in situ* polymerization to coat it with a conjugated polymer “shell” of PEDOT. The novelty of this binder architecture originates from the dual nature of the core-shell interaction, which combines electrostatic attraction with physical incompatibility. Consequently, during the drying process, the system undergoes bi-continuous mesoscopic phase separation towards highly controlled nanostructure with high conductivity (up to 32.5 S m⁻¹). This approach stands in sharp contrast to simple physical blending of PABS and PSS:PEDOT, which, as their control experiment showed, leads to macroscopic agglomeration and poor conductivity (2.2 S m⁻¹). Additionally, the PABS

component provides chemical functionality, as its abundant ester and cyanide groups effectively capture lithium polysulfides. This suppression of the shuttle effect was validated in full-cell tests, where the battery with PABS:PEDOT maintained 81.6% of its specific capacity after 450 cycles at 0.5 C. Recently, a cross-scale stabilization binder based on electronically conductive PPy was used for μ Si anodes to stabilize the SEI and electrode's stability against volume expansion. The cross scale stabilized binder comprised rigid polyacrylic acid (PAA), flexible poly(vinyl alcohol) (PVA) and conductive PPy with citric acid crosslinker. The rigid-flexible network of PAA-PVA with covalent and flexible hydrogen bonds provides pathway for the stress dissipation during cycling while the CA and PPy crosslinks direct the LiF rich growth of SEI by preferentially binding with PF₆⁻ ions. This multifunctional binder enabled stable cycling of μ Si anodes showed high CE of 99.6% with excellent areal capacity and volumetric capacity of 5.9 mAh cm⁻², 2458 mAh cm⁻³ respectively. Furthermore, full cells with NMC811 showed 81% capacity retention after 100 cycles with N&P ratio of 1.15.⁶⁷

While conjugated polymers with a stacked architecture provide structural integrity, their limited elasticity can lead to electrode delamination under repeated volume changes. Introducing a functional secondary polymer represented a solution through mechanical reinforcement. This secondary polymer can form electrostatic interactions, hydrogen bonds,



or covalent crosslinks with the primary conjugated polymer, creating a resilient and conductive network that maintains electrode stability during cycling. Matsumi *et al.* developed an n-type self-healing polymer composite (P-BIAN/PAA) for silicon anodes, where P-BIAN dynamically interacts with PAA *via* electrostatic hydrogen bonding.⁶⁸ The nitrogen atoms in P-BIAN's di-imine backbone bond with PAA's carboxyl groups, creating a crosslinked, conductive network. This structure offers both mechanical flexibility to withstand silicon's volume expansion and sustained electronic conductivity across the electrode. Similarly, Taskin *et al.* developed a crosslinked polymer network by thermally inducing esterification between carboxylic acid-functionalized poly(fluorene phenylene) and PVA. When used as a binder for silicon anodes, this system demonstrated stable cycling performance, achieving a coulombic efficiency of 99.89% after 100 cycles.⁶⁹

It is critical to note that these introduced charged moieties are also beneficial for efficient chelating the electrode active materials as well as current collector. Kang *et al.* recently engineered an adhesive-conductive (AC-polymer) interlayer by redesigning the polyanion dopant for PEDOT, towards the application for Li metal powders (LMP) anode.⁷⁰ they synthesized a P(SS-co-AA) copolymer, chemically incorporating adhesive acrylic acid (AA) segments alongside the traditional styrene sulfonate (SS) units. This molecular-level integration achieved dual functionality, herein, the -COOH groups from the AA segment provided a 28% increase in adhesive strength, while the PEDOT chains maintained their highly ordered polymer stacking and high electrical conductivity (450 S cm⁻¹). By securing the interface, this AC-polymer interlayer suppressed LMP delamination, reduced the Li stripping overpotential by 60%, and enabled stable cycling in high-rate full cells. Building on this concept of using functional interlayers for metal anodes, Mei *et al.* explored a similar strategy, but one based on nitrogen-based lithiophilic groups instead of carboxylic acids.⁷¹ They synthesized and compared two hexaazanonaphthalene (HATN)-based conjugated polymers, s-HATN (linked by flexible sigma bonds) and b-HATN (linked by sp²-carbon bonds), as a coating on the Cu current collector. The study found that tailoring the intramolecular conjugation was critical, as the more rigid, fully conjugated b-HATN exhibited higher electrical conductivity and a more favorable mesoporous structure compared to the flexible s-HATN. This superior combination of conductivity and lithiophilicity in b-HATN led to a dramatically reduced Li nucleation overpotential (down to 33.2 mV) and suppressed dendrite growth, enabling stable full-cell cycling.

This strategy of combining an elastic matrix with a conductive polymer was also demonstrated in earlier foundational work by Milroy and Manthiram.⁷² They identified that while conjugated polymers like polypyrrole (PPy) are conductive, their inherent brittleness prevents their direct use in sulfur cathodes with large volume changes. To solve this, they synthesized the nanocomposite by *in situ* emulsion polymerization of pyrrole within an elastomeric polyurethane (PU) continuous phase. This process created a distinct nanostructure

where PPy nanoparticles formed an electrically percolating network within the insulating, but elastic PU matrix. This PPy-PU binder represents a nanoparticle-in-matrix percolation model, powerfully illustrating how nanostructural design can overcome the intrinsic flaws of single-component polymers.

Blending with an elastic secondary polymer is also an important strategy to improve ionic conductivity, compensating for the fact that the rigid conjugated chains of conductive polymers typically impede ion transport. Du *et al.* demonstrated this by fabricating a hybrid PEG/PANI binder for NMC811 cathodes.⁷³ The ion-conductive PEG and electron-conductive PANI network enhanced Li⁺ transfer and provided a physical barrier to protect the cathode from HF corrosion. This blending approach has also been applied to PEDOT:PSS, where mixing it with PEO created a dual-conductor for LFP and Li_xNi_{0.5}Mn_{1.5}O₄ cathodes, resulting in high electrical conductivity of 45 S cm⁻¹ and ionic conductivity of 10⁻⁴ S cm⁻¹.⁶³ Along this line, Ryder *et al.* systematically studied the electrical conductivity and mechanical properties of variations in alginate-PANI blends with different (10–20 weight%) PANI compositions. This blend when employed as binder for graphite anode improved the rate performance and CE of the cells.

3.3 Introduction of metal ions

The ability of metal ions to induce crosslinking is rooted in fundamental coordination chemistry. Polymer chains in advanced binders are often designed with specific functional groups that can act as Lewis bases (electron-pair donors). Metal cations, being positively charged, act as Lewis acids (electron-pair acceptors). Bonds like M²⁺-carboxylate or M³⁺-sulfonate can partially reform after fracture, enabling self-healing during electrode volume changes. A prime example is the crosslinking of the widely used conductive polymer PEDOT:PSS with multivalent cations such as Mg²⁺.⁷⁴ The PEDOT:PSS aqueous dispersion contains a large excess of PSS chains with sulfonate (-SO₃H) groups. The Mg²⁺ ion can coordinate with the deprotonated sulfonate anions (-SO₃⁻) on adjacent PSS chains, forming a stable -SO₃⁻...Mg²⁺...O₃S⁻ ionic bridge. This process links the PSS chains, and by extension the PEDOT chains they are entangled with, into a robust 3D network.

Meanwhile, it is important to consider the effects of metal ion dopants in electronic and ionic conductivity of the conjugated polymers. Jiang *et al.* conducted a systematic study on how crosslinking metal cations (Li⁺, Na⁺, Mg²⁺, Ca²⁺, and Sr²⁺) influence the conductivity of the n-type POD polymer binder (Fig. 8A).⁴⁰ Their findings revealed that larger ionic radii induce greater conformational disorder in polymer chains, enhancing electrolyte wettability and ionic conductivity. However, increased chain entanglement strengthens mechanical strength at the expense of elasticity, which is a critical trade-off for long-term cycling stability. Among the tested ions, Ca²⁺-POD demonstrated an optimal balance, effectively mitigating electrode volume expansion while preserving structural integrity and conductive pathways (Fig. 8B). Cells employing the Ca-POD binder delivered outstanding performance, retain-



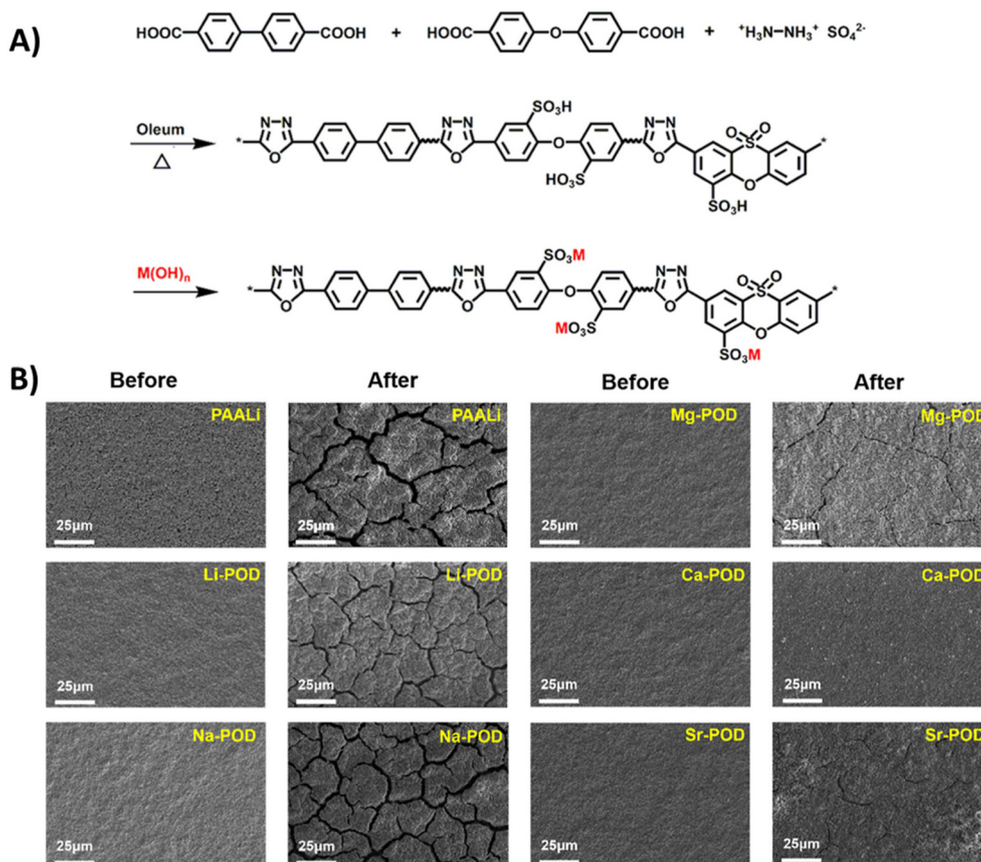


Fig. 8 (A) Synthesis procedure and structures of Mn^+ -PODs. (B) SEM images of the Si anodes with PAALi and Mn^+ -PODs binders before and after 100 cycles at 0.2 C. Reprinted by permission from American Chemical Society, ref. 40. Copyright 2023.

ing $1770.1 \text{ mAh g}^{-1}$ after 100 cycles at 0.2 C and achieving $1281.6 \text{ mAh g}^{-1}$ at 1 C.

This ion-cross-linking approach is also a powerful tool for modulating slurry viscosity, a critical parameter for fabricating high-loading electrodes. This was systematically investigated by Liu *et al.* for PEDOT:PSS binders in Si anodes.⁷⁵ They found that the viscosity of the binder solution could be precisely tuned by the choice of multivalent cation dopant, with the observed trend directly correlating to the cations' ionic potential (Φ) and softness parameters (Σ). Cations with a higher ionic potential, such as Sn^{4+} , exerted stronger electrostatic forces with the PSS chains, leading to a dramatic increase in viscosity, from 5.28 mPa s for pristine PEDOT:PSS to 60.5 mPa s for Sn^{4+} -cross-linked PEDOT:PSS. Importantly, the choice of ion also controlled the electrode morphology. The higher solubility of Sn^{4+} sulfite facilitated the formation of a smooth, homogeneously dispersed Sn^{4+} -cross-linked PEDOT:PSS film. In contrast, group 2 ions (*e.g.*, Sr^{2+} , Ba^{2+}) formed large, detrimental precipitates of insoluble sulfites. Specifically, a sheet structure with a diameter of $\sim 500 \text{ nm}$ appeared on the Sr^{2+} -cross-linked film, whereas square shaped particles appeared on the $\text{Ba}^{2+}/\text{Mg}^{2+}/\text{Ca}^{2+}$ -cross-linked binder film. This demonstrates that the rheological properties of conjugated polymer slurries can be rationally engineered *via* ion dopant selection

to achieve the desired viscosity for high-loading processing. The optimized Sn^{4+} -cross-linked binder, which combined high viscosity with uniformity, enabled Si anodes to retain $1876.4 \text{ mAh g}^{-1}$ after 100 cycles.

3.4 Introduction of small molecules

Small molecules are highly effective additives for conjugated polymer binders in lithium-ion batteries, enabling precise crosslinking control and multifunctional enhancement. Typical example include glycerol, which leverages its trifunctional hydroxyl groups to form robust covalent networks *via* esterification with acid groups on polymers like PEDOT:PSS.⁷⁶ Unreacted hydroxyl groups further establish dynamic hydrogen bonds, enabling self-healing during electrode volume changes. Simultaneously, glycerol plasticizes the binder by increasing chain mobility, effectively suppressing crack propagation while enhancing ionic transport and mechanical flexibility. Jiang *et al.* developed an innovative 3D crosslinked conductive poly-oxadiazole (POD) binder using glycerol (GL), effectively dissipating mechanical stress and accommodating volume changes in silicon particles (Fig. 9A).¹⁹ Notably, similar to the effects of metal ions, incorporating small-molecule crosslinkers can increase structural disorder and reduce the crystallinity of conjugated polymers. This reduction in crystallinity here however



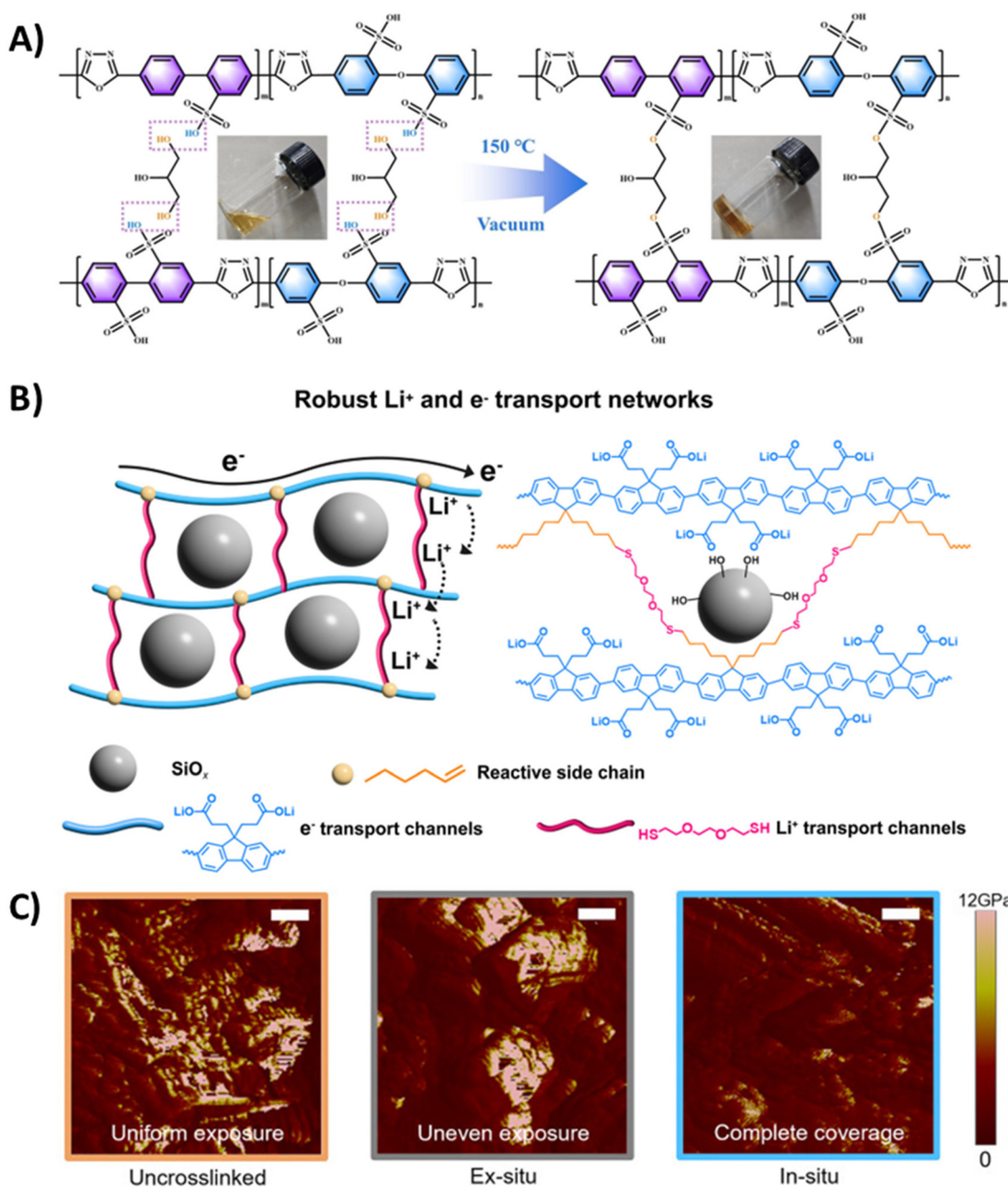


Fig. 9 (A) Schematic drawing of the crosslinked POD-c-GL binder formed by condensation between LPOD and GL. Reprinted by permission from Wiley, ref. 19, copyright 2024. (B) Architecting an elastic binding network to promote e⁻/Li⁺ transport via the proposed *in situ* photo-crosslinking strategy. (C) DMT modulus mappings of the above three binder electrodes. The scale bars represent 300 nm. Reprinted by permission from Cell Press, ref. 78, copyright 2025.

benefits ionic conductivity: POD-c-GL demonstrates higher ionic conductivity than POD across all tested temperatures. At 30 °C, for instance, crosslinking boosts the Li⁺ conductivity of POD-c-GL to $2.61 \times 10^{-4} \text{ S cm}^{-1}$ from $2.24 \times 10^{-4} \text{ S cm}^{-1}$. Nevertheless, a trade-off emerges in electronic conductivity. POD-c-GL exhibits an electronic conductivity of $5.84 \times 10^{-2} \text{ S cm}^{-1}$, slightly lower than non-crosslinked POD ($9.19 \times 10^{-2} \text{ S cm}^{-1}$). A further trade-off involves mechanical performance. Excessive crosslinking increases brittleness primarily due to

elevated local stress, while insufficient crosslinking compromises the material's structural integrity. Therefore, optimizing the crosslinking degree is critical.

When employing small molecules as crosslinkers for conjugated polymer binders, the processing strategy is a critical consideration. While many studies have directly utilized pre-crosslinked polymer binders in slurry-based electrode fabrication,⁷⁷ the *in situ* crosslinking approach—where the conjugated polymer binder is crosslinked after electrode coating—is often



the preferred method. Yang *et al.* demonstrated this by employing a light-induced thiol-ene click reaction between a 3,6-dioxa-1,8-octanedithiol crosslinker and olefin-terminated side chains on a linear polyfluorene binder (PFDP-Li) to form a binder network (Fig. 9B).⁷⁸ *In situ* crosslinking was found to facilitate a homogeneous electron/ion transport network in SiO_x electrodes. Compared to SiO_x anodes with *ex situ* crosslinked PFDP-Li binders, those with *in situ* crosslinked PFDP-Li exhibited superior rate performance and lower charge transfer resistance, attributed to more uniform binder coverage. A more direct visualization was provided by Derjaguin–Muller–Toporov (DMT) modulus mapping. Both uncrosslinked PFDP-Li and *ex situ* crosslinked PFDP-Li electrodes displayed localized high-modulus regions (Fig. 9C), suggesting incomplete embedding of SiO_x particles within the polymer network, which leads to uneven internal stress during electrode volume changes. In contrast, the *in situ* crosslinked electrode showed consistent DMT modulus values matching the binder, confirming the formation of a uniform and fully integrated 3D elastic network. Along this direction, Pan *et al.*, developed an *in situ* crosslinking strategy with Si nanoparticles using PEDOT:PSS and citric acid crosslinker in IPA. This binder with multiple anchoring sites forms strong hydrogen bonding, chemical and physical crosslinks with Si nanoparticles. The linear PEDOT chains within the binder provides excellent electronic conductivity to the binder while the hydroxyl groups PSS promote Li ion conductivity. Furthermore, the citric acid crosslinker enables the formation of lithium citrate SEI layer on the Si anode which could buffer the volume changes during lithiation and delithiation process. This unique binder design could hold up to 90% Si particles and displayed excellent electrochemical performance metrics of 89% and 81% capacity retention after 2000 cycles at 0.1 and 0.2 A g⁻¹.⁷⁹

Similarly, Jiang *et al.*, developed a dual crosslinking of Si anodes by an esterification reaction between –SO₃H of PODs and –OH groups on Si particles and functionalized CNT-OH particles at 120 °C under vacuum. This crosslinked mixed ion/electron conducting binder provides mechanically robust network through multiple hydrogen bonds for the Si particles which help to maintain electrodes structural integrity and stress management during cycling. The Si anode with this binder showed a very high capacity of 3079 mA h g⁻¹ at 0.6 A g⁻¹ after 200 cycles and 2151 mA h g⁻¹ at 2 A g⁻¹ after 500 cycles.⁸⁰

4. Conclusion and perspective

Conjugated polymer binders have emerged as a transformative solution for next-generation lithium-ion batteries, effectively addressing three critical challenges: sufficient electron conductivity, smooth ion transport, and mechanical integrity. By rationally tailoring the polymer backbone and side-chain functionalities, their π -conjugation characteristics and solid-state chain organization can be precisely modulated to optimize n-type or p-type doping performance, thereby enabling various

applications in both anode and cathode systems. Moreover, the molecular architecture of these conjugated polymers critically determines their interfacial behavior within the electrode–electrolyte matrix, dictating their adhesion to active materials, accommodation of volumetric strain, as well as electrolyte solvation dynamics which is essential for enabling sufficient processability and facilitating efficient electron/ion transport.

A critical advantage driving the development of conjugated polymer binders is their potential to significantly improve practical, full-cell energy density. Conventional electrodes always require two inactive components including an insulating binder (like PVDF) and a separate conductive additive (like carbon black). These “dead weight” materials dilute the cell’s overall gravimetric and volumetric energy density. Because conjugated binders are multifunctional, they can partially or completely replace the conductive additive. Furthermore, these polymers are what enable the use of high-capacity materials like the Si anode. For example, a layered biphenyl-polyoxadiazole (CBPOD) binder enabled a high-performance Si anode, allowing an NMC811||Si full cell to exhibit a remarkable gravimetric energy density of 574 Wh kg⁻¹.⁸⁰

The performance of conjugated polymer binders is fundamentally governed by their molecular packing, particularly the π - π stacking that dictates both electrical and mechanical properties. As this review has detailed, strategies to tune this structural orderliness normally aim for a “Goldilocks” state, balancing processability, mechanical integrity, and mixed conductivity. However, this goal is frustrated by inherent trade-offs, such as sacrificing electronic conductivity for flexibility or impeding ion transport when optimizing π - π stacking. Consequently, the widely adapted design principle evolve from merely balancing these compromises to overcoming them. This generally necessitates the development of multifunctional polymers with hierarchical architectures, where nano structural control is leveraged to decouple conflicting properties. In such systems, mechanical robustness, electronic conductivity, and ionic transport can be engineered as distinct, yet synergistic, functions of the overall architecture. Various components, such as secondary polymers, small molecules, and ions, have demonstrated their ability to function as additives with conjugated polymer binders to achieve this goal.

While simplifying the design of binder systems remains an ideal goal, recent advances in understanding ion transport physics within polymer binders have opened new avenues for a paradigm shift. Notably, the discovery of ballistic ion transport in a densely packed, electron-conductive polyfluorene binder has also demonstrated Li⁺ conductivities one to two orders of magnitude higher than those of conventional PEO-based systems.⁸¹ This unique ion-conduction mechanism, being independent of segmental motion, breaks the fundamental trade-offs of conventional polymers. It enables a single material to be simultaneously mechanically robust, electron-conductive, and ion-conductive. This discovery moves binder design away from multi-component formulations and toward simpler, high-energy systems based on a single, multifunc-



tional polymer. The redox activity of the polymer binders also becomes crucial in enhancing the ion transport and diffusion kinetics in LIBs. Thus, employing different redox active moieties in the polymer backbone or side chains becomes important in not only improving the ion transport but also improves the storage capacity, adhesion and overall stability of the batteries.

To enable the practical adoption of conjugated polymer binders, scalable and efficient synthesis methods must also be developed. Currently, conjugated polymer production relies on energy-intensive organic synthesis and polymerization techniques, such as polycondensation of specialized aromatic or heterocyclic monomers (*e.g.*, fluorene, thiophene, or aniline).⁸² Processes like electrochemical polymerization or transition-metal-catalyzed coupling (*e.g.*, Stille, Kumada reactions⁸³) suffer from high energy demands and low yields. Furthermore, post-polymerization modifications, such as sulfonate group incorporation to balance ionic and electronic conductivity, add complexity to the synthesis. Future research should prioritize environmentally benign processing to facilitate commercialization. So far, the processing of most conjugated polymers require hazardous solvents (*e.g.*, chlorobenzene), complicating handling and disposal.⁸⁴ While traditional PVDF binders also use toxic solvents like NMP, efforts are underway to transition to aqueous alternatives. Additionally, optimizing the quality of slurry processing of conjugated polymer binder is critical, as conjugated polymers seldom display predictable rheology and they tend to aggregate in electrode formulations,⁸⁵ often necessitating surfactants or additives to ensure uniformity. By addressing these synthesis and processing challenges, conjugated polymer binders can unlock their full potential for next-generation battery applications.

Looking forward, the design principles of multifunctional conjugated binders are proving critical not only for conventional liquid-electrolyte systems but also for next-generation high-energy-density platforms like solid-state batteries (SSBs). In SSBs, creating a stable, multifunctional interface to address the poor solid–solid contacts and sluggish Li⁺ transport in the composite electrodes is a primary challenge. Li *et al.* demonstrated a dual conductive polymer network for LiCoO₂ cathode in SSB, formed by the molecular interaction of an ionic conductor (lithiated polyvinyl formal-derived single-ion conductor, LiPVFM), an electronic conductor (PEDOT:PSS), and a lithium salt (LiODFB). This composite network yielded both exceptionally high electronic conductivity (68.9 S cm⁻¹) and high Li⁺ ionic conductivity (2.76 × 10⁻⁴ S cm⁻¹).⁸⁶ The HOS-PFM polymer binder serves as another key example. Its inherent electronic and ionic conductivity enabled the fabrication of “additive-free” silicon anodes (60% Si, 40% HOS-PFM) for use in SSBs. In a full cell test, this “additive-free” anode demonstrated a 114% improvement in cycle life compared to electrodes using a conventional PVDF/PEO/carbon black formulation. Ultimately, this evolution of the binder from a passive glue to a dominant, multifunctional, and active component is the key to unlocking the next generation of high-energy-density batteries.

Conflicts of interest

The authors declare no conflict of interest.

Data availability

No primary research results, software or code have been included and no new data were generated or analyzed as part of this review.

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