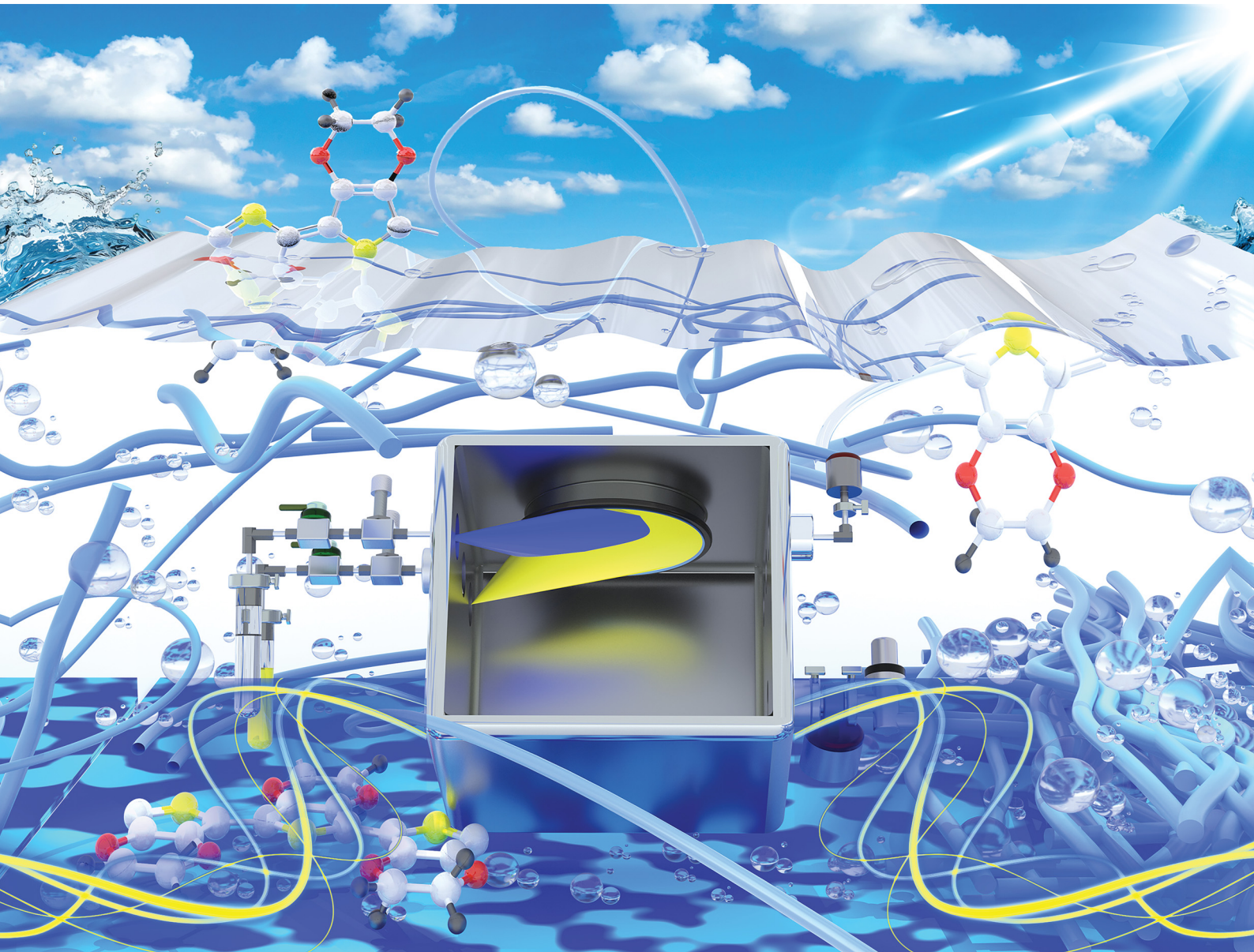


# Energy Advances

[rsc.li/energy-advances](https://rsc.li/energy-advances)



ISSN 2753-1457

**REVIEW ARTICLE**

Meysam Heydari Gharahcheshmeh and Kafil Chowdhury  
Fabrication methods, pseudocapacitance characteristics,  
and integration of conjugated conducting polymers in  
electrochemical energy storage devices

## REVIEW ARTICLE

[View Article Online](#)  
[View Journal](#) | [View Issue](#)



Cite this: *Energy Adv.*, 2024, 3, 2668

## Fabrication methods, pseudocapacitance characteristics, and integration of conjugated conducting polymers in electrochemical energy storage devices

Meysam Heydari Gharahcheshmeh  <sup>\*a</sup> and Kafil Chowdhury  <sup>ab</sup>

Among the diverse range of modern renewable energy storage technologies, electrochemical energy storage devices have been rapidly adopted across various applications owing to their superior characteristics, including high coulombic efficiency, elevated energy and power densities, scalability, modularity, and rapid response capabilities. Conjugated conducting polymers have recently attracted significant attention in electrochemical energy storage devices due to their unique pseudocapacitive behavior, hybrid ionic/electronic conduction, rapid doping/de-doping dynamics, bulk intercalation of ionic species, high specific capacity, and exceptional structural and thermal stability. Conducting polymers exhibit pseudocapacitance through reversible redox reactions coupled with doping/de-doping processes, facilitating the movement of counterion dopants and ionic species between the polymer matrix and the electrolyte. The size and nature of counterion dopants significantly influence the electrochemical performance of these polymers. Small counterion dopants like chloride enhance redox exchange with the electrolyte and broaden the electrochemical potential window, which is advantageous for electrochemical energy storage devices. The pseudocapacitive properties can be further enhanced by increasing the semi-crystalline characteristics and attaining longer polymer chains. This review article focuses on the fabrication methods, fundamental aspects of ionic and electrical conductivity, and pseudocapacitance characteristics of conjugated conducting polymers, as well as their applications in Li-ion batteries, supercapacitors, and redox flow batteries.

Received 14th August 2024,  
Accepted 14th October 2024

DOI: 10.1039/d4ya00504j

[rsc.li/energy-advances](http://rsc.li/energy-advances)

<sup>a</sup> Department of Mechanical Engineering, San Diego State University, 5500 Campanile Drive, San Diego, CA 92182, USA. E-mail: [mheydari@sdsu.edu](mailto:mheydari@sdsu.edu)

<sup>b</sup> Department of Mechanical and Aerospace Engineering, University of California at San Diego, 9500 Gilman Drive, La Jolla, CA 92093, USA



Meysam Heydari  
Gharahcheshmeh

functional materials, such as conducting polymers and superconductors, fabricated using chemical vapor deposition (CVD) for renewable energy and biocompatible device applications.

Dr Meysam Heydari Gharahcheshmeh is currently an Assistant Professor in the Department of Mechanical Engineering at San Diego State University (SDSU). He completed his postdoctoral training in the Department of Chemical Engineering at the Massachusetts Institute of Technology (MIT) in 2021 and received his PhD in Materials Science and Engineering from the University of Houston in 2017. His research focuses on texture and nanostructure engineering of advanced



Kafil Chowdhury

energy-related applications including solar cells, supercapacitors, and redox flow batteries.

Kafil Chowdhury is currently pursuing his PhD as a joint doctoral program student in the Department of Mechanical Engineering at San Diego State University (SDSU) and Department of Mechanical and Aerospace Engineering at the University of California, San Diego (UCSD). His research focuses on manufacturing conducting polymers using the chemical vapor deposition (CVD) method and their implementation in various renewable-

## 1. Introduction

To mitigate the negative impacts of greenhouse gas emissions on the global climate, policymakers are increasingly prioritizing sustainable energy technologies with reduced carbon footprints.<sup>1–4</sup> Nowadays, a diverse range of energy storage technologies, including chemical, electrochemical, mechanical, thermal, and electromagnetic systems, are being developed to meet specific application needs.<sup>5</sup> Among these renewable energy technologies, electrochemical energy storage devices, such as lithium-based batteries (lithium-ion and lithium–metal), redox flow batteries, and supercapacitors, are particularly promising due to their high energy conversion efficiency, scalability, and portability.<sup>3,6</sup>

The growing interest in utilizing conjugated conducting polymers in various electrochemical energy storage devices is driven by their pseudocapacitive properties, unique hybrid ionic/electronic conduction, rapid doping/de-doping behavior, relatively high specific capacity, high electrical conductivity, exceptional structural and thermal stability, low oxidation potential, and superior mechanical flexibility.<sup>7–10</sup> The presence of mixed ionic and electronic conduction (MIEC) in conjugated conducting polymers, despite its inherent complexity, has attracted substantial interest in the field of electrochemical energy storage devices.<sup>3,10,11</sup> In recent years, a diverse range of conjugated conducting polymers, including polypyrrole (PPy), polyaniline (PANI), polythiophene (PT), and poly(3,4-ethylene-dioxythiophene) (PEDOT), as well as their composites and derivatives, have been extensively employed in electrochemical energy storage devices.<sup>9,10,12–17</sup>

The pseudocapacitive characteristics of conjugated conducting polymers arise from reversible redox reactions coupled with doping/de-doping processes, which facilitates the movement of counterion dopants and ionic species between the polymer matrix and electrolyte.<sup>8,18,19</sup> The doping process in conjugated conducting polymer thin films can be electrochemically induced by applying an external electric field while the conjugated conducting polymer thin film is in contact with an electrolyte. In the doped state, conjugated polymer chains typically adopt an extended conformation to minimize repulsion between similar charges. Decreasing the magnitude of the applied electric field leads to the reduction of polycationic polymer chains and the expulsion of counterions from the conjugated polymer into the electrolyte, resulting in dedoping. In this de-doped state, the conjugated polymer chains often adopt a compact conformation.<sup>3</sup> The doping process in conducting polymers leads to the formation of charged species such as polarons and bipolarons along the conjugated backbone.<sup>8</sup> During de-doping process in p-type conducting polymers, bipolarons reduce to polarons, and polarons to neutral segments, while positively charged ions from the electrolyte enter the polymer thin film to maintain overall charge balance.<sup>10,20</sup>

To be effective as pseudocapacitive materials and achieve high specific capacitance, conjugated polymers must meet several criteria, including low molecular weight, an appropriate electrochemical potential window, high thermal and chemical

stability, good mechanical flexibility, excellent charge mobility, and high electrical conductivity.<sup>3,10</sup> The theoretical capacitance of a material represents the maximum charge it can hold relative to its mass.<sup>21</sup> Consequently, the maximum theoretical capacitance in conjugated conducting polymers corresponds to one charge per monomer unit, making low molecular weight polymers such as PANI, PPy, and P3HT generally preferred for achieving high specific capacitance.<sup>3</sup> However, conducting polymers with high molecular weights, such as PEDOT, also demonstrate excellent pseudocapacitance due to their superior ionic charge mobility, outstanding electrical conductivity, exceptional thermal and chemical stability, and resilience to expansion and contraction during electrochemical reactions.<sup>7,9,10,22–24</sup> Conjugated conducting polymers undergo expansion and contraction in response to electrochemical reactions, where redox processes induce elasticity. The inclusion or exclusion of anions from the polymer matrix causes strain, which varies with the amount of anionic redox species involved.<sup>9,25,26</sup> Conducting polymers with high molecular weights, such as PEDOT, effectively withstand the expansion and contraction that occur during electrochemical reactions.

The storage of electrochemical energy is governed by two principal mechanisms: the formation of electrochemical double layers on electrode surfaces through adsorption and faradaic charge storage in pseudocapacitive materials.<sup>8,27</sup> To enhance the storage capacity of electrochemical double layers, it is crucial to increase the electrode surface area available for adsorption. Various porous carbon materials have been investigated as high-surface-area electrodes for electrochemical double-layer capacitors (EDLCs).<sup>28</sup> The incorporation of a pseudocapacitive layer onto the carbon electrodes provides an additional reservoir for electrochemical charge. This faradaic charge storage is facilitated by reversible redox reactions occurring within the volume of the pseudocapacitive layer, leading to a substantial increase in total storage capacity.<sup>3,10</sup> The intercalation of ions within conjugated conducting polymer chains enables the use of the entire polymer volume for charge transfer, extending beyond the electrode/electrolyte interface and thereby enhancing pseudocapacitance.<sup>9,10,18</sup> High surface area is essential for efficient pseudocapacitive charge storage, as thinner layers over larger interfacial surface areas accelerate the charging and discharging rates for a given volume of conjugated polymer.

The size and nature of counterion dopants are critical in determining the electrochemical performance of conjugated conducting polymers.<sup>8,10,29</sup> Conjugated conducting polymers can be doped with a various range of counterions, from small monoatomic anions like chloride (Cl<sup>–</sup>) to large macro-molecular polyanions such as poly(styrene sulfonate) (PSS), tosylate, and 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ). Using small counterion dopants like chloride in conjugated conducting polymers facilitates redox exchange with the electrolyte and offers a broader electrochemical potential window, which is advantageous for electrochemical energy storage devices.<sup>8,10,30,31</sup> Enhancing the pseudocapacitive properties of conducting polymers can be achieved by increasing



their semi-crystalline characteristics and attaining longer polymer chains.<sup>7,8</sup> Ion diffusivity differs between crystalline domains and amorphous regions,<sup>3,19</sup> and conducting polymer thin films with an edge-on orientation exhibit higher charge storage capacities during charge/discharge cycles.<sup>10,31</sup> Additionally, smaller counter-ion dopants enhance ionic conductivity and accelerate redox reactions.

The large-scale adoption of electric vehicles (EVs), electronics, and drones has intensified the demand for high-energy, durable, and cost-effective lithium-ion batteries (LIBs).<sup>32</sup> Recent advancements in conjugated conducting polymers within Li-based batteries have significantly improved performance and stability.<sup>33,34</sup> Optimizing the anode surface area in LIBs is crucial: reducing anode surface area enhances energy density by minimizing lithium consumption for solid electrolyte interphase (SEI) formation, while increasing anode surface area boosts power density but lowers energy density due to higher lithium consumption.<sup>35</sup> The artificial SEI film, replicating the natural SEI layer, enhances Li-metal battery performance by regulating  $\text{Li}^+$  deposition and suppressing Li dendrite growth. Conducting polymers with excellent ionic conductivity, superior processability, and outstanding flexibility are ideal for use as artificial SEI.<sup>36</sup> To address the volumetric expansion of Si anodes in LIBs and prevent interfacial side reactions, stable SEI development is essential. Conducting polymer coatings on porous Si anodes enhance electrochemical performance by stabilizing the SEI layer, improving electronic conduction, rate capability, cycling stability, volumetric tolerance, electrolyte uptake, and electron transport.<sup>12,37,38</sup>

As a promising cathode for next-generation LIBs, Ni-rich NCM, particularly  $\text{NCM}_{811}$  ( $\text{LiNi}_{0.8}\text{Co}_{0.1}\text{Mn}_{0.1}\text{O}_2$ ), offers high energy density with a specific capacity of  $\sim 210 \text{ mA h g}^{-1}$ , good power densities, and reduced cost but faces challenges such as low thermal stability, poor cycle life, and rapid capacity degradation.<sup>32,39</sup> To address these issues, intensive research has focused on surface coating of NCM cathodes to enhance cycling performance using various materials, including metal oxides,<sup>40,41</sup> metal fluorides,<sup>42,43</sup> metal phosphates,<sup>44,45</sup> Li-ion conductive coatings (e.g.,  $\text{LiTi}_2\text{O}_4$ ),<sup>46,47</sup> and conducting polymers.<sup>48,49</sup> Compared to inorganic coatings, conducting polymers offer high electronic conductivity, environmental stability, and low cost, thus stabilizing cycling and improving rate capability. The lithium-sulfur (Li-S) battery is a promising Li-metal based technology, offering a high energy density of  $\sim 2600 \text{ W h kg}^{-1}$  and low-cost. However, the practical application of Li-S is hindered by challenges such as poor rate capability and limited rechargeability due to the insulating nature of  $\text{S}_8$  and  $\text{Li}_2\text{S}$ , rapid capacity degradation from the dissolution of lithium polysulfides ( $\text{Li}_2\text{S}_n$ ,  $3 \leq n \leq 6$ ), and significant volume expansion ( $\sim 80\%$ ) during charging.<sup>33,50-52</sup> To improve sulfur cathode utilization and cycling stability, various sulfur composites are used, including sulfur dispersed in conductive mediums like carbon-based materials,<sup>53,54</sup> and conducting polymers.<sup>55,56</sup> Coating the sulfur cathode in Li-S batteries with conducting polymers enhances capacity retention, improves overall capacity by restricting the shuttling of polysulfide intermediates, and mitigates electrode

degradation from sulfur's volume expansion due to the polymers' self-healing properties.<sup>55,57</sup>

Supercapacitors have attracted significant attention for their exceptional capacitance (100–1000× higher than conventional capacitors), high power density, rapid charge/discharge times (1–10 seconds), and long cycling life ( $> 30\,000$  hours), making them ideal for flexible electronics, electric vehicles, and renewable energy devices.<sup>2,7,58</sup> Enhanced capacitance is achieved by increasing electrode surface area, leading to the use of high-surface-area materials such as activated carbon (AC), carbon nanotubes (CNTs), graphene, graphene oxide (GO), carbon cloths (CCs), carbon fibers (CFs), and other nanoporous structures.<sup>2</sup> Pseudocapacitors, which typically offer higher energy density than EDLCs, bridge the gap between EDLCs and rechargeable batteries.<sup>2,20</sup> Conducting polymers and transition metal compounds ((TMCs)), such as metal oxides/sulfides/nitrides/carbides, are commonly used as electrode materials in pseudocapacitors.<sup>2</sup> The mechanical stability of conducting polymers, combined with their straightforward fabrication, redox properties, and mixed ionic and electrical conductivity, makes them ideal for use in flexible supercapacitors.<sup>2,3,20</sup> PANI stands out as a leading candidate among conducting polymers, noted for its lower molecular weight and impressive specific capacitance of  $1284 \text{ F g}^{-1}$ .<sup>59</sup> Conducting polymers generally show high specific capacitances, such as  $480 \text{ F g}^{-1}$  for PPy,<sup>60</sup> and  $210 \text{ F g}^{-1}$  for PEDOT.<sup>10,61</sup>

Redox flow batteries (RFBs) are ideal for large-scale, cost-effective electrochemical energy storage due to their simple and modular design, scalability, long-duration storage, rapid load response, high efficiency, safety, and long cycle life.<sup>62,63</sup> Common electrode materials in RFBs include low-cost porous carbons like carbon fiber paper (CFP), carbon cloth (CC), and graphite felt (GF).<sup>9,64,65</sup> Carbon cloth shows promise for RFB electrodes but suffers from low specific capacitance, limited surface area, and poor wettability due to its hydrophobic nature.<sup>9,10,66</sup> Similarly, graphite felt faces challenges like low electrochemical activity, limited surface area, poor wettability, and suboptimal chemical kinetics which results in low mass transfer.<sup>67</sup> To improve the electrochemical performance of carbon-based electrodes, various modification techniques have been explored, including wet chemical oxidation,<sup>68,69</sup> electrochemical oxidation,<sup>70,71</sup> thermal treatment,<sup>72,73</sup> plasma enhancement,<sup>74,75</sup> and utilization of conducting polymer coatings.<sup>9,10</sup> Applying thin layers of conducting polymers and their composites onto carbon electrodes has proven to be an effective method for improving the electrochemical performance of carbon-based electrodes in RFBs.<sup>9,10,16,76</sup> PEDOT, specifically, can be chemically tailored to regulate the diffusion of electroactive species within its structure, making it an excellent ion-selective material.<sup>16</sup>

This review focuses on the electrochemical characteristics of conjugated conducting polymers and their applications in electrochemical energy storage devices, such as Li-ion batteries, supercapacitors, and redox flow batteries. Various solution-based and vapor-based fabrication methods for conjugated conducting polymers are also discussed. Additionally, the impact of texture



and nanostructure on the electrical and ionic conductivity of conjugated polymers is examined, emphasizing their importance in the development of next-generation materials with diverse counterion dopants to meet future demands.

## 2. Fabrication methods for conducting polymers

Fabrication methods and corresponding process parameters can significantly influence various physical and chemical properties of conjugated conducting polymers, such as crystallinity, glass transition temperature, molecular weight, and morphology. Fabrication techniques for conducting polymers can be divided into two broad categories: solution-based and vapor-based, as illustrated in Fig. 1. Solution-based methods include: spin coating,<sup>77,78</sup> *in situ* chemical polymerization (ICP),<sup>79–81</sup> electropolymerization (electrochemical polymerization (ECP)).<sup>82–85</sup> Vapor-based techniques includes: vapor phase polymerization (VPP),<sup>86–93</sup> oxidative chemical vapor deposition (oCVD).<sup>2,9,94–99</sup>

In addition to these established methods, other polymerization techniques such as hydrothermal,<sup>100</sup> electrospinning,<sup>101</sup> self-assembly,<sup>102</sup> interfacial,<sup>103</sup> and plasma polymerization,<sup>104</sup> have also been used for fabricating conducting polymer thin films and composites.

### 2.1. Solution-based fabrication methods for conducting polymers

**2.1.1 Spin coating method.** Spin coating is a widely used solution-based method for fabricating polymeric thin films with thicknesses ranging from micrometers to nanometers. The spin coating process involves depositing a viscous polymer solution onto a rotating flat substrate, resulting in a uniform thin film. The fabrication process consists of four steps: (i) dispensing liquid precursors, (ii) substrate acceleration (spin-up), (iii) spin-off, and (iv) thermal annealing (Fig. 1a). The quality of the spin-coated thin film depends on several process parameters, including angular velocity, polymer molecular weight, liquid precursor viscosity, additive and solvent presence, additive/solvent volatility, and the surface

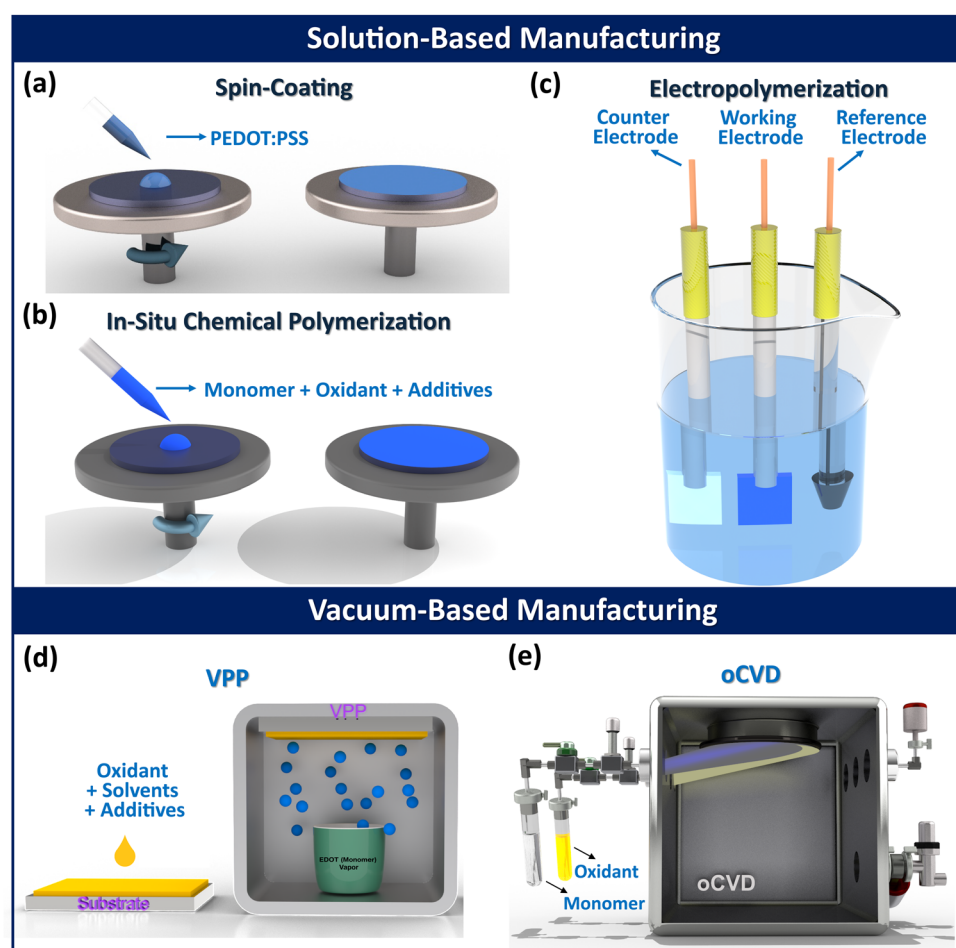


Fig. 1 Various methods commonly used for fabricating conjugated conducting polymers. Solution-based methods include (a) spin-coating,<sup>2</sup> (b) *in situ* chemical polymerization (ICP), and (c) electrochemical polymerization (ECP). Vapor-based methods include (d) vapor phase polymerization (VPP), and (e) oxidative chemical vapor deposition (oCVD).<sup>105</sup> (a) Reproduced with permission.<sup>2</sup> Copyright 2019, Wiley. (e) Reproduced with permission.<sup>105</sup> Copyright 2019, Science.



energy between the substrate, polymer film, and liquid precursor.<sup>106,107</sup>

The electrical conductivity of conducting polymer thin films fabricated by spin coating can be enhanced by adding organic polar solvents with high boiling points, such as dimethyl sulfoxide (DMSO), *N,N*-dimethyl formamide (DMF), glycerol, ethylene glycol (EG), *N,N*-dimethyl acetamide (DMAc), sorbitol, and dichloromethane (DMC).<sup>8,29,108–110</sup> The main advantages of the spin coating method are its quickness and low cost. However, its major drawbacks include: (i) inability to create ultrathin polymeric films (thickness  $<10$  nm), (ii) inability to provide conformal coatings, (iii) restriction to planar substrates, as the technique is ineffective for coating non-flat or complex surfaces, (iv) utilization of only a small fraction (2–5%) of the polymer solution, and (v) low quality and presence of contaminants in the fabricated thin films.<sup>2,106,111,112</sup>

**2.1.2. *In situ* chemical polymerization (ICP) method.** The ICP process begins by dissolving the monomer, oxidant, and additives into a solution, which is then deposited onto the substrate surface by spin coating, allowing polymerization and doping to occur simultaneously (Fig. 1b). The properties of the resultant polymeric thin film fabricated *via* the ICP method are highly influenced by factors such as molecular weight, type of oxidant, presence of organic solvents and additives, and the morphology of the deposited thin films. Oxidant characteristics (solubility, concentration, stability, anion and cation constituents, and oxidation strength) are crucial in the polymerization process in the ICP method because they determine the rate of monomer oxidation and influence the morphology and conductivity of the resultant conducting polymer thin film.<sup>113</sup>

Numerous oxidants are commonly used in the ICP method, including copper(II) chloride ( $\text{CuCl}_2$ ),<sup>114,115</sup> copper(II) perchlorate ( $\text{Cu}(\text{ClO}_4)_2$ ),<sup>116</sup> iron *p*-toluene sulfonic acid ( $\text{Fe}(\text{PTS})$ ),<sup>117,118</sup> iron chloride ( $\text{FeCl}_3$ ),<sup>117,118</sup> iron(III) *para*-tosylate ( $\text{Fe}(\text{Tos})_3$ ), and ammonium peroxydisulfate (APS) with *p*-toluene sulfonic acid (PTSA) as the oxidant and dopant, respectively.<sup>119</sup> Various organic solvents, such as dimethylformamide (DMF), *N*-methyl-2-pyrrolidone (NMP), acetonitrile–chloroform, and *n*-butanol, are used for fabricating different conducting polymers, including PEDOT, poly(ethylene terephthalate) (PET) fiber, and graphite/PMMA nanocomposites.<sup>120–123</sup> Additionally, different additives such as base inhibitors (*e.g.*, imidazole),<sup>124</sup> polymeric surfactants (*e.g.*, PEG–PPG–PEG), and high boiling point cosolvents (*e.g.*, NMP) are employed to modify the morphology and other physicochemical characteristics of the fabricated polymer thin films through the ICP method.<sup>113</sup>

The main benefits of the ICP technique over spin-coating are (i) a simpler and faster bottom-up approach and (ii) effective dispersion of nanoparticles with less aggregation. However, the ICP method has disadvantages compared to vapor phase methods, including (i) inability to provide conformal coatings due to surface tension effects, (ii) unsuitability for solvent-sensitive substrates, and (iii) potential for flocculant formation during polymerization.

**2.1.3. Electropolymerization method.** In the electropolymerization method, a thin film of a conducting polymer is

deposited on the surface of a conductive supporting electrode (the working electrode) by anodic oxidation of the monomer in a three-electrode cell setup, which includes a reference electrode, a counter electrode, and the working electrode (Fig. 1c). Common reference electrodes include Ag/AgCl and saturated calomel electrode (SCE), while platinum, stainless steel, and high-density graphite are used for the counter electrode. Various electropolymerization techniques can be employed, including potentiostatic (constant voltage), galvanostatic (constant current), and potentiodynamic (cyclic voltammetry). During electropolymerization, the monomer dissolves in a solvent containing an anionic doping salt (electrolyte solution). Applying a specific deposition potential ( $E_{\text{Polymerization}}$ ) oxidizes the monomer on the anode surface, forming a radical cation.<sup>82</sup> This radical cation reacts with other monomers in the solution to form oligomers, which, due to their lowered oxidation potential, more easily form further radical cations, thereby initiating polymerization and transitioning from monomer radicals to dimers, tetramers, octamers, and higher oligomers.<sup>125</sup> As chain length increases, solubility decreases, leading to film deposition on the anode surface. Simultaneously, anions or counterions are incorporated into the polymer matrix during film formation.

Selecting the appropriate solvent and electrolyte is critical for the electropolymerization method, as they must remain stable at the monomer's oxidation potential while providing ionic conductivity.<sup>82</sup> The solvent type (aqueous, non-aqueous, ionic liquid, protic, aprotic) and electrolyte properties (pH, composition) significantly influence the film properties and polymerization rate.<sup>82,126,127</sup> Organic solvents like DMSO, acetonitrile, dichloromethane, propylene carbonate, nitrobenzene, and methyl benzoate are commonly used in the electropolymerization method for fabricating conducting polymer thin films.<sup>128,129</sup> Aqueous solvents are preferred in the electropolymerization method due to advantages like easy handling, environmental safety, lower cost, and the availability of a wide range of possible counterions for incorporation.<sup>130</sup> Room temperature ionic liquids are also used as solvents in electrochemical polymerization method due to their non-volatility, wide electrochemical stability window, and environmental friendliness.<sup>82</sup>

The electropolymerization process is favored over other solution-based techniques for its excellent substrate adhesion, better control over film thickness and morphology, and prevention of extra chemical species accumulation (from oxidant or catalyst) on the polymeric film.<sup>131</sup> However, the electropolymerization method has a drawback compared to other solution-based methods, as it requires conductive substrates, limiting the choice of materials.

## 2.2. Vapor-based fabrication methods for conducting polymers

**2.2.1. Vapor phase polymerization (VPP) method.** The VPP method is a two-step fabrication for depositing conducting polymer thin films on various substrates.<sup>90,132,133</sup> In the first step, an oxidizing agent with low volatility is applied to the substrate surface *via* wet chemistry techniques, such as spin coating, and after heating and drying the oxidant thin film is placed upside down inside a vacuum chamber (Fig. 1d). In the



second step, the monomer vapor is exposed to the oxidant-coated substrate within the vacuum chamber, where polymerization occurs at the oxidant-monomer vapor interface.<sup>90,133,134</sup> For VPP, the chamber is usually kept at ambient pressure or a moderate vacuum of 5–30 Torr, with vacuum conditions becoming the norm and used frequently.<sup>92</sup> Notably, VPP can also be performed without drying the oxidant, allowing for the fabrication of conducting polymers on liquid droplet oxidants.<sup>135</sup> The VPP mechanism is similar to oxidative polymerization in solution, involving step growth polymerization between a monomer and an oxidizing agent.

Common oxidants used in VPP method include iron(III) chloride ( $\text{FeCl}_3$ ),<sup>91</sup> iron(III) tosylate [ $\text{Fe}(\text{OTs})_3$ ],<sup>136</sup> iron(III) trifluoromethanesulfonate [ $\text{Fe}(\text{OTf})_3$ ],<sup>137</sup> and copper(II) chloride ( $\text{CuCl}_2$ ).<sup>138</sup> These oxidants are typically dissolved in organic solvents like methanol, ethanol, butanol, or water to prepare spin-coatable precursors. The polymerization rate can be regulated using different solvents such as DMSO, NMP, DMF, and chelating ligands like EDTA.<sup>8,29,139</sup> The utilization of solvents and additives in the VPP method introduces potential coordination molecules into the oxidant solution, forming metal complexes and altering ligand coordination, thereby changing the oxidant's overall reactivity. Base inhibitors such as imidazole, polyethyleneimine (PEI), or pyridine, when dissolved with the oxidant, can enhance the film's conductivity by increasing the pH and inhibiting side reactions under acidic conditions, and reducing the polymerization rate.<sup>92,93,140</sup> The surfactant poly(ethylene glycol)-block-poly(propylene glycol)-block-poly(ethylene glycol) (PEG-PPG-PEG) has been reported to improve electrical conductivity of conjugated conducting polymer thin films by lowering the polymerization rate and preventing oxidant crystallization.<sup>141</sup>

The polymerization rate and characteristics of fabricated conducting polymers in VPP method can be controlled by adjusting parameters like exposure time of the substrate with the monomer vapor,<sup>92</sup> pressure of vacuum chamber,<sup>86</sup> substrate temperature,<sup>2,86,90,142</sup> relative humidity inside the polymerization chamber,<sup>87</sup> acidity level of oxidant,<sup>143</sup> and post-deposition acidic rinsing treatment.<sup>92</sup> The VPP method offers several key advantages for fabricating conducting polymers, including compatibility with various substrates, operation in a moderate vacuum environment for fabricating good quality thin films, fabrication of smooth thin films, and the ability to deposit on flexible or delicate substrates.<sup>90,92,133</sup> The drawbacks of the VPP method include the complexity of tuning interconnected process parameters, the initial formation of oxidant as a solid thin film in the absence of monomer preventing fabrication of ultrathin highly conformal coatings, and the risk of substrate surface damage from solution casting, making it incompatible with solution and temperature-sensitive substrates.<sup>90,92,133</sup>

**2.2.2. Oxidative chemical vapor deposition (oCVD) method.** The CVD method is a powerful manufacturing technique for large-scale production of inorganic semiconductors, conjugated conducting polymers, and functional polymers.<sup>2,144–146</sup> The oCVD method, first reported by Karen Gleason and her team in 2006,<sup>147</sup> was developed as a robust, simple, and solvent-free vapor-deposition technique for fabricating highly conformal conducting and semiconducting polymer thin films. To start

the oCVD procedure, the substrate is placed upside down on a temperature-controlled stage inside the vacuum chamber. The monomer and liquid oxidant, stored in separate heated glass jars, are connected to the reactor chamber *via* temperature-controlled feed lines (Fig. 1e).<sup>30</sup> Once vaporized, the monomer and oxidant vapors are fed into the vacuum chamber, where step-growth polymerization occurs on the temperature-controlled substrate.<sup>2,30</sup> An inert carrier gas, such as argon, is used for low volatile species or as a gas balancer for fundamental studies. The oxidizing agent in the oCVD method is heated and converted into vapor through sublimation for solid oxidant, or evaporation for liquid oxidants.

Common solid oxidants include iron(III) chloride ( $\text{FeCl}_3$ )<sup>30,95,98,148,149</sup> and copper(II) chloride ( $\text{CuCl}_2$ ),<sup>2,97</sup> while recently reported liquid oxidants include vanadium oxytrichloride ( $\text{VOCl}_3$ )<sup>9,30,105,150</sup> and antimony pentachloride ( $\text{SbCl}_5$ ).<sup>10,30,151,152</sup> Liquid oxidants are often more effective than solid oxidants due to their higher volatility, better flow rate control, and elimination of post-treatment acidic rinsing, as they leave negligible oxidant byproducts on the as-deposited thin films.<sup>2,30,105</sup> The characteristics of conducting polymers fabricated using the oCVD method can be controlled by tuning process parameters such as oxidant saturation ratio (OSR),<sup>105,152</sup> deposition temperature,<sup>10,153,154</sup> chamber pressure,<sup>2,30,146</sup> incorporation of water-assisted growth techniques,<sup>30,152,155</sup> and choice of oxidant.<sup>105</sup> The most important parameters for oxidants used in oCVD method are volatility, thermal stability, and the ability to drive the reaction forward.<sup>30,156</sup>

The oCVD method enables the fabrication of conducting polymers on any substrate type (planar/non-planar, organic/inorganic, porous/dense, conductor/insulator) and produces robust, patterned thin films *via* covalent grafting between the polymer film and substrate without requiring any linker molecule.<sup>2</sup> Given that majority of conjugated conducting polymers are insoluble in solvents, solvent-free and dry methods like vapor-based manufacturing (VPP, and oCVD) are highly desired.<sup>2,30,157</sup> The most significant advantages of oCVD over other techniques include its ability to form highly conformal coatings, independence from substrate type, compatibility with large-scale roll-to-roll (R2R) production, solvent-free single-step processing, and capability to create uniform thin films at low deposition temperatures.<sup>30,144,146,148</sup>

The Fabrication methods section provided an overview of various techniques used to fabricate conjugated conducting and semiconducting polymers. These fabrication methods often rely on specific oxidants to facilitate polymerization. The commonly explored oxidants employed in various fabrication processes are summarized in Table 1.

### 3. Specific characteristics of conjugated conducting polymers

#### 3.1. Charge transport mechanisms in conjugated conducting polymers

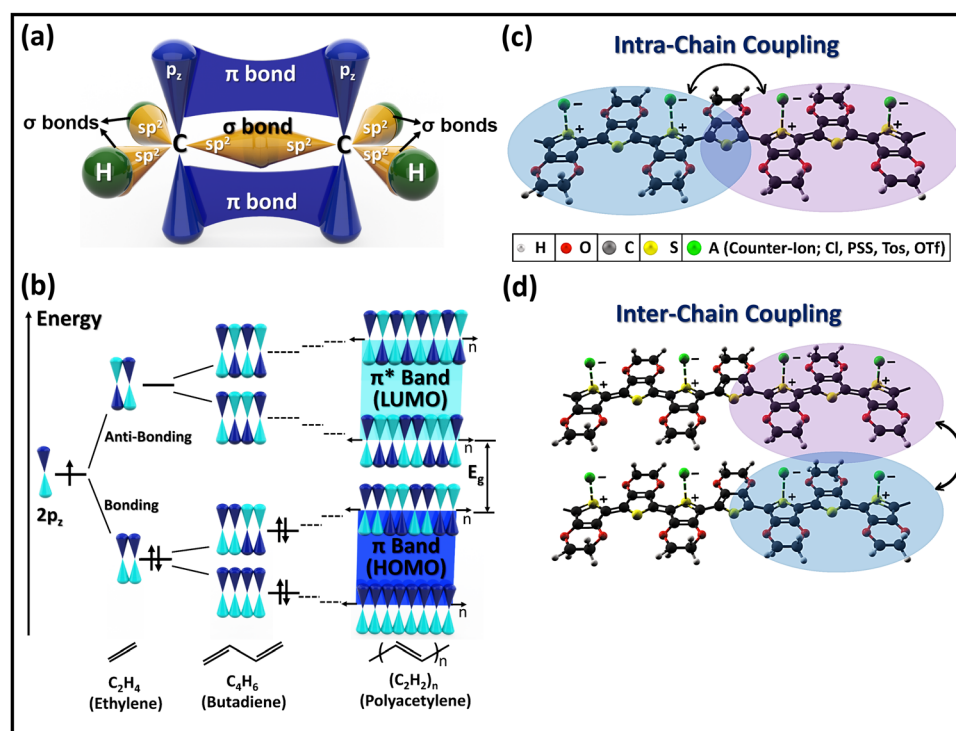
Organic polymers are typically insulators due to their large band gap, which is the significant energy difference between

**Table 1** Common oxidants used in various fabrication methods of conjugated polymers. The table provides details on oxidants commonly employed in the fabrication of conjugated conducting polymers, in both solution-based and vacuum-based processes

Fabrication method	Common explored oxidants
Spin coating	Polymer is typically dissolved in a solution, with $\text{FeCl}_3$ occasionally added as oxidant
Electropolymerization	Oxidation of a monomer is induced by voltage or current
<i>In situ</i> chemical polymerization (ICP)	$\text{FeCl}_3$ , $\text{CuCl}_2$ , $\text{Cu}(\text{ClO}_4)_2$ , $\text{Fe}(\text{pTS})$ , $\text{Fe}(\text{Tos})_3$ , ammonium peroxydisulfate (APS)
Vapor phase polymerization (VPP)	$\text{FeCl}_3$ , $\text{CuCl}_2$ , $\text{Fe}(\text{OTs})_3$ , $\text{Fe}(\text{OTf})_3$
Oxidative chemical vapor deposition (oCVD)	$\text{FeCl}_3$ , $\text{CuCl}_2$ , $\text{VOCl}_3$ , $\text{SbCl}_5$

the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) levels.<sup>8</sup> However, certain types of organic polymers, known as conjugated polymers, can exhibit semiconducting behavior due to the presence of alternating single ( $\sigma$ ) and double ( $\pi$ ) bonds along their polymer chain (Fig. 2a), creating an overlap of unbonded  $p_z$  orbitals that allows  $\pi$ -electrons to flow along the polymer backbone skeleton.<sup>3,8,29,158</sup> The doping process in conjugated polymers reduces the bandgap, thereby increasing their conductivity from an insulator to a conductive level.<sup>8,29</sup> The bonding ( $\pi$ ) and anti-bonding ( $\pi^*$ ) orbitals of carbons in the polymer chain eventually create the  $\pi$  (HOMO) and  $\pi^*$  (LUMO) bands (Fig. 2b).<sup>8</sup> The difference between the HOMO and LUMO energy levels, known as the bandgap, dictates the optical and optoelectronic characteristics of the polymer.<sup>8,152,159</sup> As the length of polymer chain increases, the bonding and antibonding orbitals get rearranged with different energy levels and practically reducing the bandgap which enhances conductivity.

There are two distinct pathways for the movement of charge carriers inside the conjugated polymer matrix: intrachain and interchain coupling (Fig. 2c and d).<sup>8</sup> Intrachain coupling allows charge carriers to move between adjacent localized polymer segments along the same polymer backbone, whereas interchain coupling facilitates charge transport through hopping between individual chains or within a folded chain. Charge carrier mobility is generally much faster along the conjugated backbone (intrachain transport) compared to interchain transport.<sup>8,29,30,105,160,161</sup> Interchain transport is a rate-limiting factor, and reducing the  $\pi$ - $\pi$  stacking distance is an effective method to enhance carrier mobility and overall electrical conductivity in various conjugated conducting polymers, including PEDOT,<sup>10,105,152,162–165</sup> P3HT,<sup>166,167</sup> and diketopyrrolopyrrole (DPP)-based donor-acceptor (D-A) polymers.<sup>168</sup> While intrachain coupling provides increased carrier mobility, a high packing order in chain orientation enhances interchain coupling, reducing charge localization caused by defects and amorphous regions.<sup>8</sup>



**Fig. 2** Charge transport in conjugated conducting polymers. (a) Schematic illustration of sigma and pi bonds in a conjugated polymer.<sup>8</sup> (b) Change in bandgap as a consequence of bonding and antibonding orbitals in conjugated polymers.<sup>8</sup> Schematic illustration of charge carrier transport through (c) intra-chain coupling,<sup>8</sup> and (d) inter-chain coupling.<sup>8</sup> (a)–(d) Reproduced with permission.<sup>8</sup> Copyright 2020, Elsevier.



The incorporation of dopants in the polymer results in the oxidation or reduction of the polymer backbone, forming local excitons or quasiparticles (polaron, bipolaron, and soliton) along the polymer chain.<sup>8,159</sup> These positively or negatively charged quasiparticles act as charge carriers along the conjugated backbone chains. Based on the polarity of the dominant charge carriers, conducting polymers can be classified as p-type (hole transport), n-type (electron transport), or ambipolar.<sup>8,29</sup>

The most widely studied and applicable p-type conjugated polymers include PEDOT, PANI, PPy, and PTh, as illustrated in Fig. 3a.<sup>29</sup> On the other hand, the most frequently reported n-type conjugated polymers are diketopyrrolopyrrole (DPP), naphthalene diimide (NDI), benzodifurandione-based oligo(*p*-phenylene vinylene) (BDOPV), isoindigo (IID), and benzothiadiazole (BT), shown in Fig. 3b.<sup>29</sup> Notably, n-type (reduced) conducting polymers exhibit less stability compared to p-type (oxidized) ones, making oxidized conducting polymers (p-type conducting polymers) more widely used in practical applications.<sup>3,8</sup>

### 3.2. Influence of nanostructure on electrical conductivity of conjugated conducting polymers

The electrical conductivity of conjugated conducting polymers can vary significantly depending on their semicrystalline characteristics such as crystallite size, semicrystalline orientation, and  $\pi$ - $\pi$  stacking distance.<sup>8,29,152,169–171</sup> With respect to the substrate surface, crystallites in conducting polymer thin films can exhibit a face-on orientations, edge-on orientations, end-on orientations, and a mixture of different orientations (isotropic).<sup>8</sup> The conjugated backbone lies parallel to the plane

of substrate in face-on orientation but the  $\pi$ - $\pi$  stacking is perpendicular; whereas for edge-on, both the conjugated backbone and  $\pi$ - $\pi$  stacking lie parallel to the plane of substrate (Fig. 4a).<sup>2,152,155</sup> Two of the most widely used conjugated conducting polymers, PEDOT and P3HT, exhibit a semicrystalline nature and assemble in an orthorhombic unit cell structure (Fig. 4b and c).<sup>8</sup> The conjugated polymer chains stack along the *b*-axis lattice parameter to form face-to-face layers, with  $\pi$ - $\pi$  stacking corresponding to half of the *b*-axis lattice parameter. In both highly edge-on and highly face-on orientations, the conjugated backbone chains lie in the plane of conjugated polymer thin films, enabling high in-plane electrical conductivity. However, a mixture of face-on and edge-on orientations reduces unperturbed percolation pathways, leading to reduced electrical conductivity.<sup>8,152</sup>

The possible charge carrier transport pathways in highly face-on and highly edge-on crystallite orientations in conjugated polymers are illustrated in Fig. 4d and e, respectively. The electrical conductivity in heavily doped conjugated conducting polymers with highly face-on (or highly edge-on orientation) and short  $\pi$ - $\pi$  stacking distance is quasi-1D.<sup>8,105,152</sup> In the quasi-1D conductivity, charge transport parallel to the substrate primarily occurs (i) along polymer chains that traverse multiple crystallites and bridge amorphous regions, and (ii) between chains in the crystalline regions.<sup>8,171</sup> Interchain transport is higher in the crystallite domains and slower in the less dense amorphous regions.<sup>105</sup> Reducing the  $\pi$ - $\pi$  stacking distance in conjugated conducting and semiconducting polymers increases both the charge transfer integral and carrier mobility and yields the electrical conductivity enhancement.<sup>8,105,152,155</sup>

Fig. 5 displays the timeline of gradual increment in the reported conductivity of a specific conducting polymer, PEDOT. The highest electrical conductivity achieved in PEDOT thin films to date is  $7520\text{ S cm}^{-1}$ , obtained using the water-assisted oCVD method (W-A oCVD) with  $\text{SbCl}_5$  as the oxidant.<sup>152</sup> Kim *et al.*<sup>172</sup> reported a conductivity of  $1418\text{ S cm}^{-1}$  in PEDOT thin films with EG as an additive. In 2014, Kim *et al.*<sup>173</sup> achieved a conductivity of  $4380\text{ S cm}^{-1}$  by using different concentrations of  $\text{H}_2\text{SO}_4$  as a post-treatment, which induced crystallinity and structural improvement. Worfolk *et al.*<sup>174</sup> used solution shearing during deposition that yielded PEDOT:PSS thin film with a conductivity of  $4600\text{ S cm}^{-1}$ . Gueye *et al.*<sup>175</sup> utilized the ICP method to deposit PEDOT thin films with  $\text{Fe}(\text{OTf})_3$  as the oxidant, and *N*-methyl pyrrolidone (NMP) as the polymerization rate controller, achieving a conductivity of  $5400\text{ S cm}^{-1}$ . Using the oCVD method with solid  $\text{FeCl}_3$  oxidant and HBr rinsing, PEDOT thin films exhibited a conductivity of  $6259\text{ S cm}^{-1}$ .<sup>153</sup> Moreover, using liquid  $\text{SbCl}_5$  as oxidant, the oCVD method yielded PEDOT thin films with a conductivity of  $5602\text{ S cm}^{-1}$ .<sup>152</sup> Post-deposition acidic rinsing steps, such as using  $\text{HCl}$ ,  $\text{HBr}$ , or  $\text{H}_2\text{SO}_4$ , have been shown to significantly enhance the electrical conductivity of conducting polymer thin films. Post-deposition acidic treatments improve conductivity by removing insulating impurities and inducing structural rearrangements that promote better charge transport.<sup>2,29,153,176</sup> For instance,  $\text{H}_2\text{SO}_4$  post-treatment has

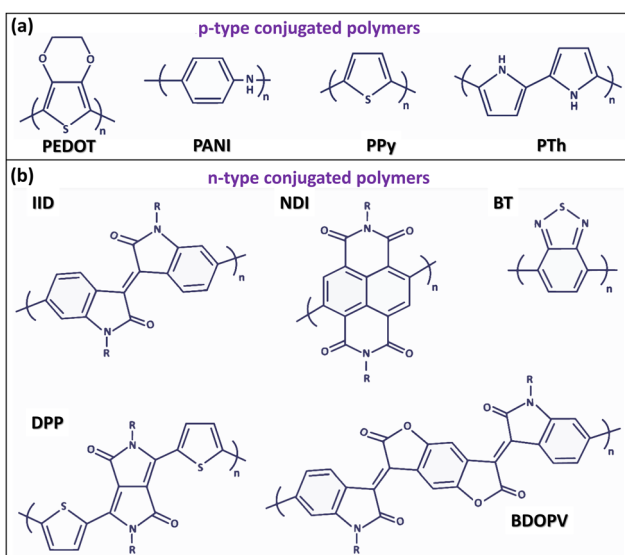


Fig. 3 Chemical structures of the most relevant p-type and n-type conjugated conducting polymers. (a) Chemical structures of p-type conjugated polymers, including poly(3,4-ethylenedioxythiophene) (PEDOT), polyaniline (PANI), and polythiophene (PTh).<sup>29</sup> (b) Chemical structures of significant n-type polymers, such as isoindigo (IID), naphthalene diimide (NDI), benzothiadiazole (BT), diketopyrrolopyrrole (DPP), and benzodifurandione-based oligo(*p*-phenylene vinylene) (BDOPV).<sup>29</sup> (a) and (b) Reproduced with permission.<sup>29</sup> Copyright 2022, MDPI.



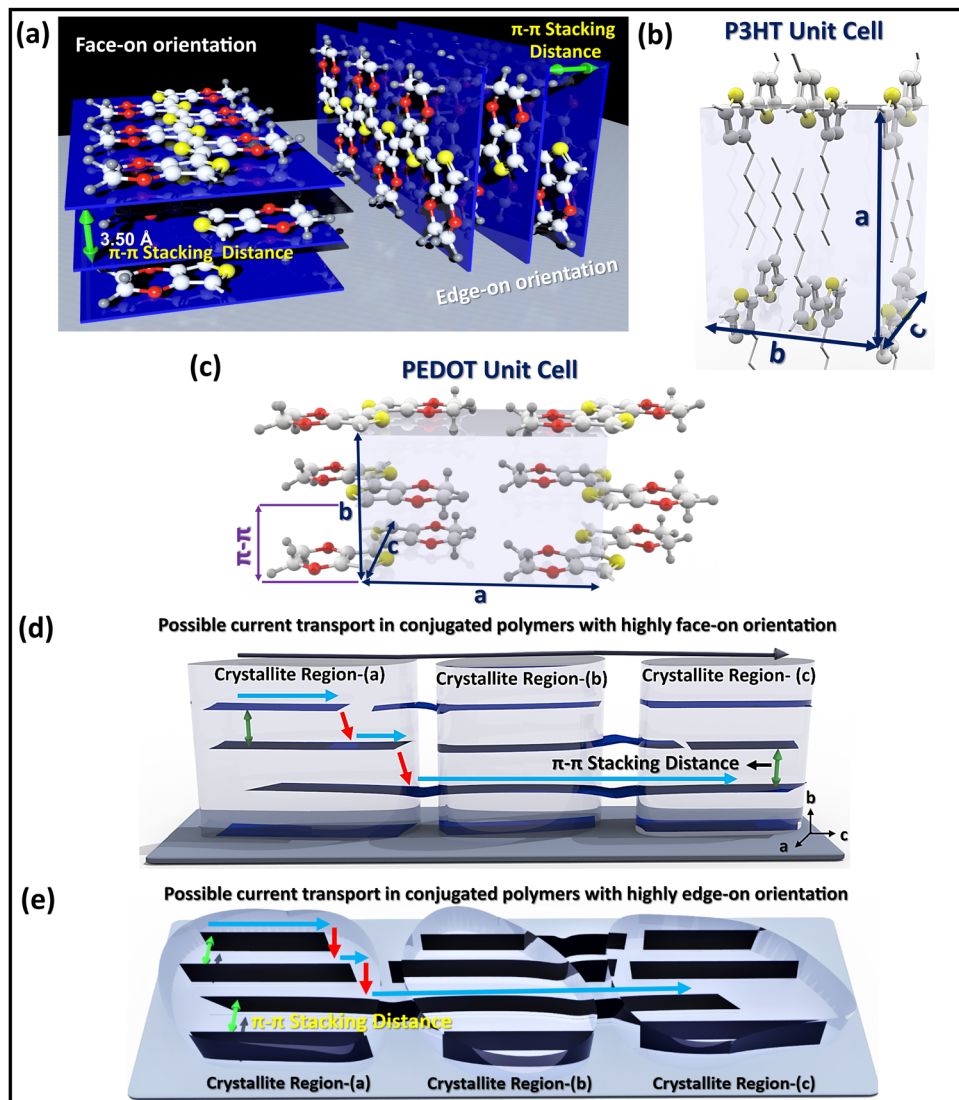


Fig. 4 Texture and nanostructure of conjugated conducting polymers. (a) Schematic illustration of face-on and edge-on orientation along with the  $\pi-\pi$  stacking distance, which can significantly affect the overall electrical conductivity.<sup>152</sup> Orthorhombic unit cell structure of (b) P3HT,<sup>8</sup> and (c) PEDOT.<sup>152</sup> Charge carrier transportation pathways between different ordered crystallite regions in conjugated conducting polymers with (d) highly face-on orientation,<sup>105</sup> and (e) highly edge-on orientation.<sup>8</sup> (a) and (c) Reproduced with permission.<sup>152</sup> Copyright 2020, Wiley. (b) and (e) Reproduced with permission.<sup>8</sup> Copyright 2020, Elsevier. (d) Reproduced with permission.<sup>105</sup> Copyright 2019, Science.

been reported to increase the crystallinity and overall electrical performance of PEDOT films.<sup>8,153,173</sup>

### 3.3. Molecular weight and glass transition temperature of conjugated conducting polymers

Conjugated conducting polymers exhibit both expansion and contraction in response to electrochemical reactions, where redox processes induce elasticity. The inclusion or exclusion of anions from the polymer matrix leads to straining.<sup>9,25,26</sup> The extent of strain depends on the amount of anionic redox species entering or departing from the polymer. For instance, applying a positive voltage along with a suitable electrolyte causes oxidation, leading to electron loss and anion entry into the polymer matrix, resulting in polymer expansion.<sup>9</sup> Conversely, reduction causes contraction. These electrochemical

deformations are useful in applications such as artificial muscles. Semi-crystalline polymers typically exhibit superior mechanical properties compared to amorphous polymers.<sup>177</sup>

The mechanical behavior of conjugated polymers depends on microscopic changes in molecular mobility, which are influenced by factors such as molecular weight, morphology, nanostructure, working temperature, and applied pressure.<sup>152</sup> Molecular weight is commonly characterized using the number-average molecular weight ( $M_n = \frac{\sum M_i \cdot N_i}{\sum N_i}$ ) or the weight-average molecular weight ( $M_w = \frac{\sum M_i^2 \cdot N_i}{\sum M_i \cdot N_i}$ ), where  $i$  is the degree of polymerization,  $M_i$  and  $N_i$  are the weight and number of molecules with degree of polymerization  $i$ .<sup>178,179</sup> Higher molecular weight in conjugated polymers enhances

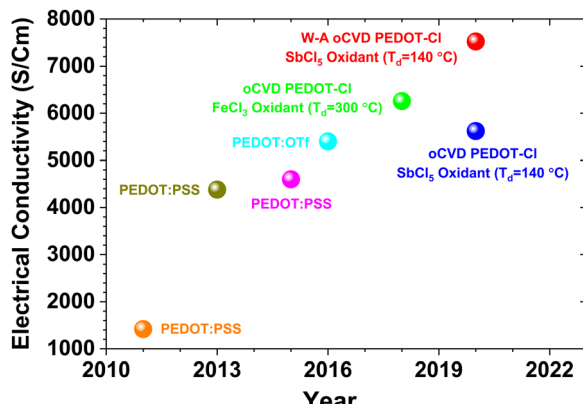


Fig. 5 Timeline of electrical conductivity for PEDOT thin films. The evolution of electrical conductivity enhancement in PEDOT thin films fabricated by various solution-based and vapor-based methods.<sup>30</sup> Reproduced with permission.<sup>30</sup> Copyright 2024, Springer.

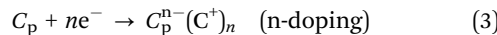
electrical conductivity, strength, and toughness due to increased density and better entanglement. Lower molecular weight polymers are brittle and cannot withstand high strain rates, whereas higher molecular weight polymers exhibit significant ductility and elongation.<sup>179</sup> The mechanical properties of conjugated conducting polymer thin films can be assessed using the strain-induced elastic buckling-based method.<sup>180</sup> The modulus of elasticity for conjugated polymer thin films ( $E_f$ ) is determined using the substrate's modulus of elasticity ( $E_s$ ), Poisson ratios of both the thin film ( $\vartheta_f$ ) and substrate ( $\vartheta_s$ ), buckling wavelength ( $\lambda_b$ ), and thin film thickness ( $d_f$ ) by using the following equation:<sup>180</sup>

$$E_f = 3E_s \left( \frac{1 - \vartheta_f^2}{1 - \vartheta_s^2} \right) \cdot \left( \frac{\lambda_b}{2\pi d_f} \right) \quad (1)$$

Reaching the glass transition temperature ( $T_g$ ) transforms the polymer from a glassy to a rubbery state, altering its mechanical properties.<sup>179</sup> The glass transition temperature represents a second-order phase transition and describes the thermally activated reorganization of polymer chains. Below  $T_g$  the polymer is glassy, and above  $T_g$  becomes elastomeric.  $T_g$  influences the morphological stability of the polymer's solid state, affecting thermal stability, dewetting, and delamination of thin films, and is critical for the mechanical properties needed in flexible optoelectronic applications.<sup>179</sup> The glass transition temperature is a critical processing parameter that dictates the kinetics of molecular organization in organic conjugated polymers during solidification,<sup>179</sup> or vacuum-based deposition methods.<sup>10,152,162</sup> In the oCVD method for fabricating PEDOT, when the deposition temperature reaches or exceeds the glass transition temperature of PEDOT (approximately 100 °C), the crystallites reorient from mixed face-on and edge-on orientation to predominantly face-on orientation.<sup>10,30,152,162</sup> The reorientation at around glass transition temperature aims to minimize interfacial energy between the substrate and the  $a-c$  plane of the PEDOT crystallite face, resulting in a highly face-on orientation, which is essential for enhanced electrical conductivity.<sup>8,30,162</sup>

### 3.4. Reversible doping/de-doping in conjugated conducting polymers

Conducting polymers can undergo p-doping with counter anions upon oxidation and n-doping with counter cations upon reduction. The simplified equations for these charging processes are as follows:<sup>20</sup>



where  $C_p$  represents the neutral conducting polymer,  $C_p^{n+}(A^-)_n$  and  $C_p^{n-}(C^+)_n$  indicate the oxidant- and reductant-doped conducting polymer with counterions,  $e^-$  denotes the electron,  $n$  states the number of electrons transferred,  $A^-$  represents counter anion, and  $C^+$  denotes counter cation.

The type of doping (p- or n-type) is typically determined from the required potential for the doping and the nature of conducting polymers, so the selection of appropriate conducting polymers based on the potential window is essential for optimal electrochemical device performance.<sup>181</sup> N-Type doping can sometimes result in undesirable properties such as high impedance, poor processability, chemical instability, and lower capacitance and conductivity due to the need for higher negative potentials than the electrolyte solvent breakdown potential.<sup>33,181,182</sup>

Doping in conducting polymers causes volumetric changes (swelling) that revert during de-doping (shrinking), which corresponds to the oxidation or reduction reaction.<sup>10,33,183</sup> Continuous electrochemical cycling causes large dimensional changes during charging and discharging, leading to material breakdown and degradation of the polymer structure, ultimately reducing electrochemical performance.<sup>184</sup> The doping process in conducting polymers results in the formation of charged particles such as polarons, bipolarons, and solitons along the polymer backbone.<sup>8</sup> During de-doping in p-type conducting polymers, bipolarons and polarons are reduced to polarons and neutral segments, respectively and positively charged ions from the electrolyte entering the film to balance the overall charge.<sup>8,10,19</sup> Inversely, positively charged ions of electrolyte are pushed out from the p-type conducting polymers during the next cycle of doping (charging) and the initial counterions again oxidize the polaron and neutral states to the bipolaron and polaron segments, respectively.<sup>10,183</sup>

Cyclic voltammetry (CV) is a fundamental technique for studying the doping/de-doping mechanism in conducting polymers. Applying a reversible potential at a fixed scan rate in CV causes conducting polymers to donate or accept electrons and ions from the electrolyte to move in and out of the polymer structure. The surface coverage ( $\Gamma$ ) is defined as the ratio of adsorbed molecules numbers on a specific surface to the number of molecules required to form a complete monolayer on that surface. The peak current ( $i_p$ ) of a reversible electrochemical system as a function of  $\Gamma$ , can be measured by the following equation:<sup>182</sup>

$$i_p = n^2 F^2 A \Gamma^3 \left[ \frac{\exp(\theta)}{RT(1 + \exp(\theta))} \right] \quad (4)$$



where  $n$  denotes the number of electrons,  $F$  is the Faraday constant ( $96\,485\text{ C mol}^{-1}$ ),  $A$  indicates the electrode surface area in  $\text{cm}^2$ ,  $\vartheta$  shows the surface coverage in  $\text{mol cm}^{-2}$ ,  $\vartheta$  specifies the scan rate in  $\text{mV s}^{-1}$ ,  $R$  is the universal gas constant ( $8.314\text{ J mol}^{-1}\text{ K}^{-1}$ ),  $T$  stands for temperature in  $\text{K}$ , and  $\theta = \frac{nF}{RT}(E - E^0)$  where  $(E - E^0)$  is the difference between cell potential and standard potential.

In general, when the redox reaction involves larger dopant ions which are sluggish to move, the reaction becomes diffusion-controlled and it was found that  $i_p$  is proportional to the square root of scan rate ( $\vartheta^{1/2}$ ).<sup>185</sup> For reversible, single-electron doping/de-doping reactions, CVs show symmetric redox peak. However, for redox reactions involving multi-electron transfers, CVs exhibit several peaks for oxidation and reduction. Moreover, the peak-to-peak distance is also observed to increase with the scan rate.<sup>186</sup> In cyclic voltammetry, the experimental timescale is controlled by the scan rate ( $\vartheta$ ), and the relationship between the observed current at any specific potential ( $i(V)$ ) can reveal whether the redox reaction is diffusion-controlled or surface-controlled. The general relationship between the observed current and scan rate for any material is described by the following equation:<sup>187</sup>

$$i(V) = k_1\vartheta^{1/2} + k_2\vartheta \quad (5)$$

where  $k_1$  and  $k_2$  are constants related to diffusion and capacitive currents, respectively.

The inner electrode surface stores charge *via* intercalation, while the outer surface, in contact with the electrolyte, stores charge *via* adsorption. Charge storage on the outer surface is instant and scan rate-independent, whereas storage on the inner surface is diffusion-controlled and scan rate-dependent.<sup>188</sup> The relationship between the inner surface (diffusion-controlled) and outer surface (surface-controlled) of an electrode to the total charges ( $Q_T$ ) measured using CV is expressed as follow:<sup>188,189</sup>

$$Q_T = Q_i + Q_o \quad (6)$$

where  $Q_i$  and  $Q_o$  represent the charges related to inner and outer surfaces, respectively.

Furthermore, the relationship between the total charges or capacity ( $Q$ ) as a function of  $\vartheta$  can be described through the following equation:<sup>58,189</sup>

$$Q(\vartheta) = Q_\infty + k\vartheta^{-1/2} \quad (7)$$

where the second term,  $k\vartheta^{-1/2}$  indicates the amount of charge stored due to semi-infinite diffusion,  $k$  is a constant, and  $Q_\infty$  describes the capacity related to high scan rates ( $\vartheta \rightarrow \infty$ ).

At  $\vartheta = 0$ , the value of  $Q$  represents the total capacity ( $Q_T$ ), encompassing contributions from capacities associated with both inner ( $Q_i$ ) and outer ( $Q_o$ ) surfaces, while at  $\vartheta \rightarrow \infty$ ,  $Q$  signifies the capacity specific to outer surfaces ( $Q_o$ ) exclusively.<sup>188</sup>

### 3.5. Pseudocapacitance characteristics of conjugated conducting polymers

In electrochemical double-layer capacitors (EDLCs), charge storage occurs at the electrode–electrolyte interface through

electrostatic (non-faradaic) interactions, as shown in Fig. 6a. The innermost layer, made up of solvent dipoles adsorbed onto the electrode surface, is known as the inner Helmholtz or Stern layer. Immediately next to this is the outer Helmholtz layer, which consists of an adsorbed layer of solvated ions. Beyond the outer Helmholtz layer lies the diffuse layer, where solvated ions and solvent molecules are randomly distributed into the bulk medium.<sup>190</sup>

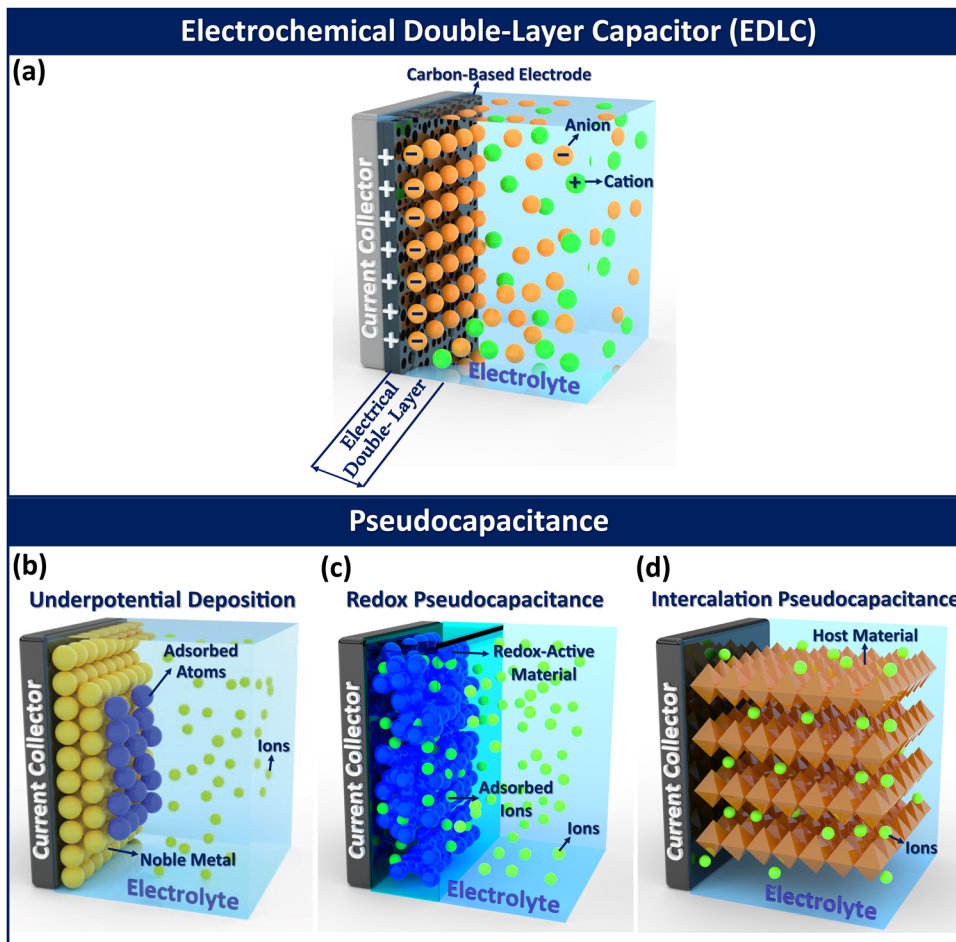
Activated carbon is the most widely recognized electrode material for EDLCs, offering a modest capacitance of up to  $200\text{ F cm}^{-2}$ , constrained by its specific surface area limitations. Typical pseudocapacitance also originates from the electrode surfaces but involves an associated faradaic charge-storage reaction, leading to significantly higher capacitance. While EDLCs depend exclusively on a physical charge storage mechanism,<sup>3</sup> pseudocapacitance describes a material's ability to replicate the electrochemical properties of capacitive electrodes (linear relationship between charge and potential) while utilizing a faradaic electron transfer mechanism for charge storage.<sup>58</sup> Utilizing redox reactions for energy storage, instead of electrostatic ion accumulation as seen in EDLCs, leads to battery-like behavior in pseudocapacitors during charging and discharging processes.<sup>20,182</sup>

Pseudocapacitors store energy through faradaic interactions, allowing them to achieve higher specific capacity and energy density compared to EDLCs.<sup>181</sup> The charge storage mechanisms in pseudocapacitors are categorized into three distinct faradaic processes: (i) underpotential deposition (also referred to as monolayer adsorption pseudocapacitance), (ii) surface redox pseudocapacitance, and (iii) intercalation pseudocapacitance (Fig. 6b–d). In underpotential deposition, charge storage is achieved *via* the adsorption of a monolayer of cations from a specific metal onto the surface of another metal at potentials more positive than their equilibrium potential (Fig. 6b). The underpotential deposition yields high capacitance but limited energy density due to a narrow potential window. Underpotential deposition can be exemplified by the deposition of lead (Pb) on a gold (Au) electrode surface. In this case, Pb is more readily deposited onto Au than onto itself due to the stronger interaction between Pb and Au compared to the Pb–Pb interaction in crystalline Pb metal.<sup>191</sup>

In surface redox pseudocapacitance, ions are electrochemically adsorbed on or near the electrode surface, with energy stored through a faradaic charge transfer mechanism (Fig. 6c). Conjugated conducting polymers demonstrate redox pseudocapacitance because they can undergo bulk intercalation, allowing ions to penetrate the entire polymer matrix, as illustrated in Fig. 6c. An example of redox pseudocapacitance can be observed in systems containing Ruthenium(IV) oxide ( $\text{RuO}_2$ ) nanoclusters.<sup>192</sup>

Intercalation pseudocapacitance occurs when the intercalation and deintercalation processes are not hindered by the slow solid-state diffusion of ions within the crystal structure of the electrode materials.<sup>58</sup> It is noteworthy that intercalation pseudocapacitance allows ions to enter the layers of redox-active materials without significant crystallographic phase changes,





**Fig. 6** Schematic illustration of various types of pseudocapacitance. Schematic illustration of (a) electrochemical double-layer capacitance (EDLC), (b) underpotential deposition, (c) redox pseudocapacitance, and (d) intercalation pseudocapacitance. Intercalation pseudocapacitance utilizes redox-active intercalation chemistry in electrode materials by incorporating surface redox pseudocapacitance and electrostatic pseudocapacitance. Conjugated conducting polymers exhibit redox pseudocapacitance due to their ability for bulk intercalation, which enables ions to penetrate the polymer structure volumetrically.

accompanied by a faradaic charge transfer process (Fig. 6d). Intercalation pseudocapacitance differs from battery systems because, unlike batteries where charge-discharge cycles often involve phase transformations of the host material, there are no phase changes in intercalation pseudocapacitance systems.<sup>193</sup> The intercalation pseudocapacitance is observed in systems where lithium ions intercalate into mesoporous and nanocrystalline films of orthorhombic niobium pentoxide ( $\text{Nb}_2\text{O}_5$ ) without any phase transformation.<sup>194</sup>

In conducting polymers, pseudocapacitive behavior stems from their reversible redox capabilities. During charging (or doping), the polymer chain oxidizes and releases electrons, while during discharging (or de-doping), the chain reduces and accepts electrons. Pseudocapacitance is measured by integrating the charging/discharging current ( $I$ ) over the entire potential window ( $E = V_f - V_i$ ) using the following equation:<sup>10,188</sup>

$$C = \frac{1}{E\vartheta} \int_{V_i}^{V_f} I \, dV \quad (8)$$

where  $V_f$  is the final potential of sweep,  $V_i$  denotes the initial potential of sweep, and  $\vartheta$  specifies the scan rate.

The gravimetric specific capacitance ( $C_{sp}$ ), which indicates the charge storage capacity and is directly related to the energy density of electrodes, can be calculated using the following equation:<sup>10,195</sup>

$$C_{sp} = \frac{\int_{V_i}^{V_f} I(V) \, dV}{2\vartheta \cdot m \cdot E} \quad (9)$$

where  $\int_{V_i}^{V_f} I(V) \, dV$  denotes the total area of the CV curve,  $\vartheta$  specifies the scan rate,  $m$  represents the mass of the electrode, and  $E$  specifies the difference between the initial and final potential of sweep ( $E = V_f - V_i$ ).

An additional common method to evaluate the behavior of electrochemical capacitors is the galvanostatic charge-discharge (GCD) test, where the specific capacitance ( $C$ ) can be calculated using the following equation:<sup>190</sup>

$$C = \frac{I\Delta t}{m\Delta V} \quad (10)$$

where  $I$  represents the discharge current,  $\Delta t$  indicates the discharge time,  $\Delta V$  describes the potential window and  $m$  is the mass of the active material in the electrode.

P-Type conjugated polymers are doped by oxidation, achievable through chemical reaction with an oxidant. A partially doped polymer chain for conjugated conducting polymer of poly-pyrrole (PPy, A = N-H), poly(thiophene) (PT, A = S), and poly(selenophene) (PSe, A = Se), showing benzoid and quinoid bonding (Fig. 7a).<sup>3</sup> Charge carriers include polarons (radical cations) and bipolarons (dication).<sup>8</sup> In doped p-type conjugated conducting polymers, polycationic chains maintain charge neutrality with negatively charged counterions. This charge neutrality links anion migration during doping/de-doping to electron mobility. Partial charge transfer with some molecular counterions forms charge transfer complexes between the polymer and dopant.<sup>3</sup> The doping process can be driven electrochemically by applying an external electric field to a conjugated polymer film in contact with an electrolyte.

In the doped state, polymer chains often extend to reduce repulsions between similar charges (Fig. 7b).<sup>3</sup> Lowering the magnitude of external electric field reduces the polycationic polymer chain, expelling counterions into the electrolyte and

resulting in dedoping. Reduced polymer chains typically adopt a compact conformation (Fig. 7c). The water solubility of PANI (polyaniline) makes it widely studied. Fully reduced PANI (leucoemeraldine) is yellow, while fully oxidized PANI (pernigraniline) is blue, with maximum ionic conductivity at 50% doping. Ionic conductivity is low at very low or very high doping levels, slowing electrochemical initiation unless pre-doped. This supports the hypothesis of ion transport between occupied and unoccupied states, with the rate proportional to the product of their concentrations.

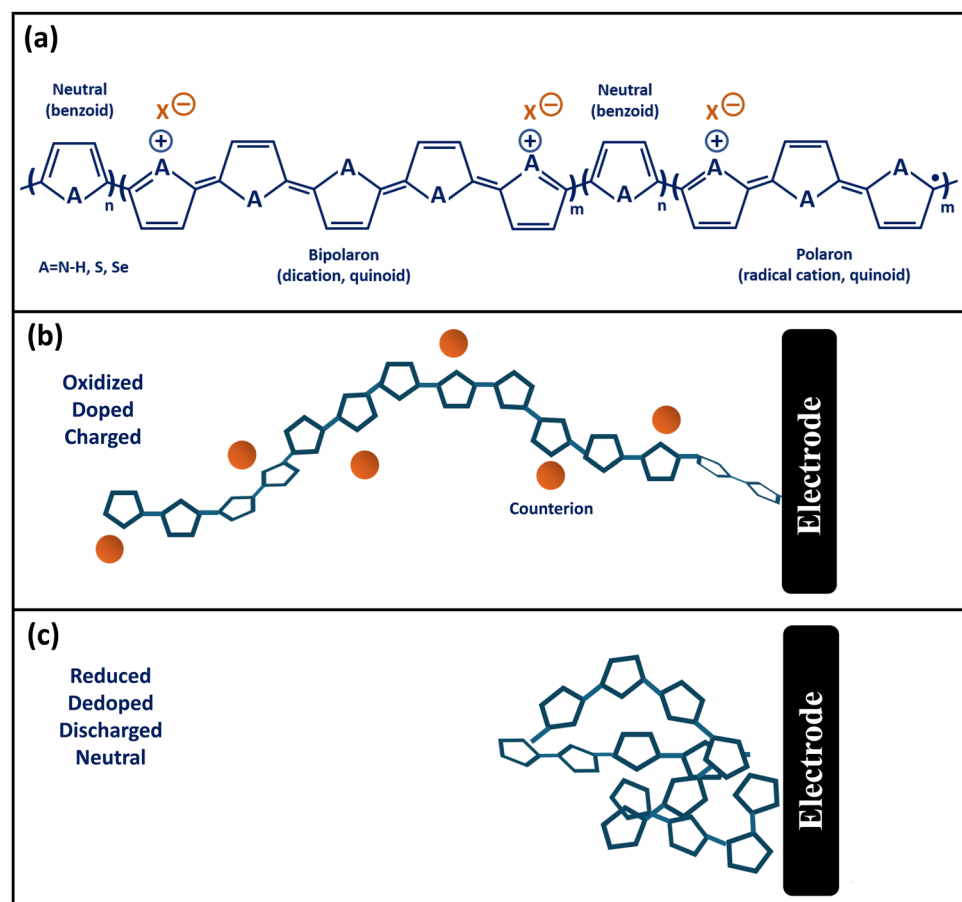
Key aspects contributing to the complex ionic transport in conjugated conducting polymers include:

(1) Ion concentration dynamics: driven by chemical potential gradients from electrical potential and ionic concentration differences, leading to ion diffusion and drift.

(2) Redox reactions: occur between ions and the polymer, altering the charge state of the conjugated backbone.

(3) Electronic states: new states like polaronic and bipolaronic states emerge in the density of state diagram.<sup>8,11,19</sup>

(4) Backbone reorganization: structural changes, such as from benzoid to quinoid configurations, occur upon doping (as shown in Fig. 7a).



**Fig. 7** Schematic illustration of conjugated conducting polymer structure during electrochemical reaction. (a) The chemical structure of a partially oxidized (doped) generic conjugated chain, having neutral segments in between localized bonding states corresponding to polaronic and bipolaronic charge carriers.<sup>3</sup> The neutral segments are in the benzoid state, while the oxidized segments display quinoid bonding. (b) Schematic illustration of conjugated polymer chain adjacent to an electrode in the (b) partially oxidized state,<sup>3</sup> and (c) fully reduced state.<sup>3</sup> (a)–(c) Reproduced with permission.<sup>3</sup> Copyright 2021, Walter de Gruyter.



Intrachain migration: localized charge carrier states, situated within specific molecular structure, can migrate along the backbone *via* intrachain hopping, while there is a different migration rates for polarons and bipolarons.<sup>8,196</sup>

(5) Interchain transfer: electron transfer between chains *via* hopping, which is a rate-limiting step.<sup>29,105,152</sup> Redox reactions between chains can also create interchain complexes in the solid conducting polymer matrix during charging, which relax during discharging.<sup>19</sup>

(6) Electrostatic forces: increase with more ion intercalation into the polymer film.

(7) Polymer conformation: the polymer chain extends to minimize electrostatic repulsion, altering its overall conformation (as illustrated in Fig. 7b)

(8) Morphology changes: conformation changes in the chains can affect the semicrystalline polymer's morphology. Also, electrochemical processes may differ in the ordered and disordered regions of the semicrystalline conjugated polymer.<sup>8-10,19,28</sup>

(9) Swelling: the electrolyte solution swells the polymer due to intercalation of redox-active components and solvent molecules.

### 3.6. Bulk intercalation and volumetric change in conjugated conducting polymers

The pseudocapacitive characteristics of conjugated conducting polymers differ from those of double-layer capacitors due to their capability for bulk intercalation, allowing ions to volumetrically penetrate the polymer structure, as exhibited in Fig. 6c (redox pseudocapacitance mechanism). Since the bulk of the conjugated conducting polymer participates in the redox reaction, rather than just the surface or interfacial layer between the electrode and electrolyte, pseudocapacitors typically provide higher specific capacitance and energy density than EDLCs.<sup>3,33</sup> The insertion and extraction of redox-active species generally lead to significant changes in the dimensions of the polymer. The total volumetric change in conducting polymers can be attributed to two distinct components: (i) changes in the bond length and conformation of the polymer chain, and (ii) osmotic expansion of the polymer phase, which is the structural change associated with the movement of solvent ions.<sup>184</sup> The concentration of ionic species within the polymer structure is determined by Donnan equilibrium,<sup>197</sup> and controlled by fulfilling two requirements: maintaining the electroneutrality of the polymer and preserving electrochemical equilibrium between the electrolyte and the polymer, as described by the following equations:<sup>184</sup>

$$\sum z_i c_i + \frac{q_p}{F} - c_A = 0 \quad (11)$$

where  $z_i$  represents the number of charges for different mobile ions,  $c_i$  indicates the concentration of mobile ions,  $q_p$  describes the volumetric density of charges in the polymer chain,  $F$  denotes Faraday constant, and  $c_A$  is the concentration of immobile anions.

Consequently, the maintenance of electrochemical equilibrium between the electrolyte and the polymer is governed by the following equation:<sup>184</sup>

$$\bar{\mu}_i^p = \bar{\mu}_i^I \quad (12)$$

where  $\bar{\mu}_i^p$  and  $\bar{\mu}_i^I$  denotes the electrochemical potentials of the polymer and electrolyte phases, respectively. The subscript  $i$  for both cases refer to the various ionic species present in the system.

The swelling and shrinkage of polymer structures during doping and de-doping processes make conducting polymers prone to rapid degradation with repetitive cycles of charging and discharging, leading to diminished capacity retention and cycling stability. To improve mechanical stability of conducting polymers during doping/de-doping process, researchers have developed composite conducting polymer nanostructures,<sup>22</sup> utilized advanced fabrication techniques like oCVD method to create rigid, durable polymeric thin films.<sup>9,10</sup>

### 3.7. Ionic conductivity in conjugated conducting polymers

A notable electrochemical property of conducting polymers is their ability to facilitate hybrid electrical and ionic conductivity by transferring both ions and electrons. The intercalation of ions within the conducting polymer matrix enables these materials to engage their full or partial volume during electrochemical redox reactions (doping/de-doping processes).<sup>8</sup> The ability of conducting polymers to facilitate the intercalation of ions from the electrolyte into their matrix effectively broadens the region of active electron transfer beyond the electrode surface or the electrode/electrolyte interface, thereby enhancing their pseudocapacitance characteristics.<sup>9,10,18</sup>

Ionic conduction in mixed ionic and electronic conductivity (MIEC) materials is typically described by the Nernst–Planck drift-diffusion model, which quantifies the ionic flux ( $\vec{N}_j$ ) as the number of moles of ions passing through a unit area per unit time ( $\text{mol cm}^{-2} \text{ s}^{-1}$ ).<sup>8</sup> The ionic flux is influenced by both the electrochemical gradient potential ( $\vec{\nabla} \mu_j$ ) and the convective velocity ( $\vec{v}$ ), as described by the following equation:<sup>8,198,199</sup>

$$\vec{N}_j = -\frac{D_j c_j}{RT} \vec{\nabla} \mu_j + c_j \vec{v} \quad (13)$$

where  $D_j$  represents the diffusion coefficient of species  $j$ ,  $c_j$  denotes the concentration of species  $j$ ,  $R$  signifies the ideal gas constant, and  $T$  denotes the temperature.

The electrochemical driving force induces both the concentration gradient of species  $j$  and the gradient in the applied electrostatic potential ( $\vec{\nabla} \phi$ ), as expressed by: (ref. 3 and 8)

$$\vec{\nabla} \mu_j = RT \vec{\nabla} \ln c_j + z_j F \vec{\nabla} \phi \quad (14)$$

where  $F$  represents Faraday's constant, and  $z_j$  denotes the charge of species  $j$ .

In the absence of convective flow ( $\vec{v} = 0$ ) and an applied electric field ( $\vec{\nabla} \phi$ ), the Nernst–Planck equation simplifies to Fick's law of diffusion. The diffusivity ( $D_j$ ) and mobility ( $\mu_j$ ) of



an ion are assumed to be interconnected *via* the Nernst-Einstein equation:<sup>8</sup>

$$\mu_j = \frac{F|z_j|D_j}{RT} \quad (15)$$

where  $F$  signifies the Faraday's constant,  $z_j$  is the charge of species  $j$ ,  $R$  denotes the gas constant, and  $T$  represents temperature.

The order and orientation of semicrystalline conjugated polymers are expected to affect ionic transport.<sup>10,31</sup> Ion diffusivity likely varies between crystalline domains and the surrounding amorphous regions.<sup>8,10,19</sup> The influence of tie-chains on ionic transport has received less attention, though noted in electrical conductivity. The orientation of crystallites relative to the substrate is anticipated to cause anisotropy in ionic diffusivity.<sup>3</sup> Variations in dopant levels, which alter electrostatic forces and polymer chain conformation, also impact ionic diffusivity.

The effects of counterion size,<sup>10,31</sup> and semicrystalline orientation<sup>10,25,200</sup> on ionic transport in these polymers are reported. The size and nature of the counterion dopant are crucial factors and significantly influencing the electrochemical performance of conjugated conducting polymers.<sup>8,10,29</sup>

Conducting polymers can be doped with various counterions, ranging from small monoatomic anions like chloride ( $\text{Cl}^-$ ) to large macromolecular polyanions such as poly(styrene sulfonate) (PSS).<sup>2,201</sup> The chemical structures of several negatively charged molecular counterions used as dopants for PEDOT conducting polymer, including PSS, tosylate, and hexafluorophosphate, are shown in Fig. 8a–c. Additionally, the molecular structure of the dopant 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ), commonly used as a counterion dopant with P3HT and other conjugated polymers, is presented in Fig. 8d. Using small counterion dopants, such as chloride ( $\text{Cl}^-$ ), in conjugated conducting polymer thin films facilitates redox exchange with the electrolyte and exhibits a wider electrochemical potential window compared to larger macromolecular dopants, providing a significant advantage for electrochemical energy storage devices.<sup>8,10,30,31</sup>

Minimizing  $\pi$ - $\pi$  stacking distance is crucial for enhancing charge carrier mobility and electron conductivity, which is achievable through a more ordered structure. Conversely, facile ion migration between the electrolyte and polymer matrix, leading to increased ionic conductivity, necessitates a more relaxed structure. Optimizing the structure of conducting

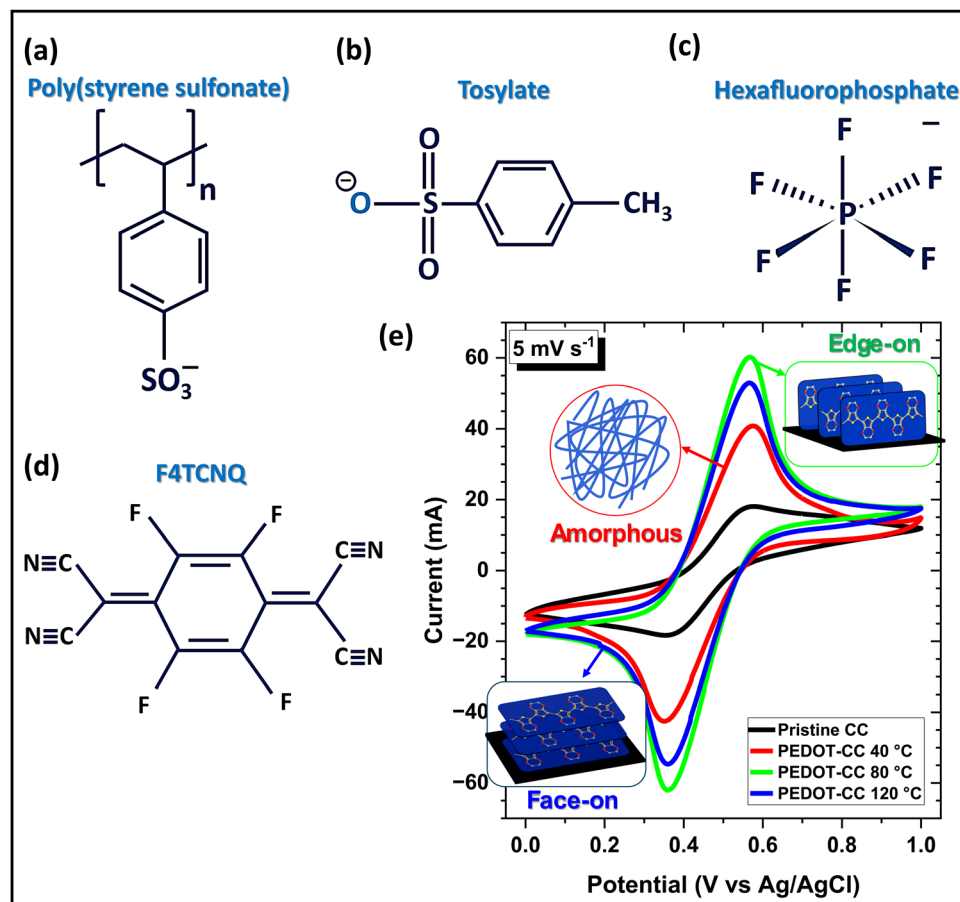


Fig. 8 Impact of counterion size and semicrystalline orientation on electrochemical properties of conjugated polymers. Chemical structure of counterion dopants used in conjugated polymer: (a) PSS, (b) tosylate, (c) hexafluorophosphate, and (d) F4TCNQ. (e) PEDOT thin film with edge-on semicrystalline orientation shows higher pseudocapacitance and specific capacitance compared to amorphous and face-on oriented counterparts.<sup>10</sup> (e) Reproduced with permission.<sup>10</sup> Copyright 2024, Wiley.



polymers to maximize both in-plane electrical conductivity and ionic conductivity presents a significant challenge due to the inherent trade-off between these parameters.<sup>3,7,29</sup> Notably, the edge-on orientation of crystallites in conjugated conducting polymers has been reported to enhance pseudocapacitance due to the inherent anisotropy in the direction of the conjugated backbone (Fig. 8c).<sup>10</sup> The edge-on orientation offers higher out-of-plane electrical conductivity, enabling faster ion exchange with the electrolyte and resulting in increased pseudocapacitance in conducting polymers.<sup>10,25,31</sup>

The ionic conductivity of a conducting polymer thin film can be measured by utilizing electrochemical impedance spectroscopy (EIS), where the conducting polymer film is used as the working electrode under various configurations.<sup>202–207</sup> The obtained impedance response from the input containing sinusoidal perturbation is comprised of double-layer formation, diffusion and migration of ions, transport of electrons, and transfer of charges.<sup>204</sup> In addition, electrochemical impedance models for conducting polymers describe various impedances as distributed parameters across the porous polymer thin film's thickness, rather than treating them as localized variables.<sup>208,209</sup> This approach considers the polymer film as a non-homogeneous structure, with distributed properties instead of a macroscopic boundary between the polymer and the electrolyte. The ionic resistance ( $R_{\text{ion}}$ ) and electronic resistance ( $R_e$ ) of the polymer phase and the interfacial faradaic impedance ( $Z_f$ ) are considered as distributed circuit elements in equivalent circuit model. Assuming rapid interfacial charge transfer, the real component of the impedance ( $Z_{\text{real}}$ ) can be defined by the following equation:<sup>204,205</sup>

$$Z_{\text{real}} = \frac{R_{\text{ion}} + R_e}{3} + R_s \quad (16)$$

where  $R_s$  designates the uncompensated resistance obtained from the real-axis intercept at the high-frequency region.

The value of  $Z_{\text{real}}$  can be obtained by extrapolating the low-frequency line to meet the real axis. Finally, the ionic conductivity ( $\sigma_{\text{ion}}$ ) of the conducting polymer film can be measured by the following equation:<sup>206,207</sup>

$$\sigma_{\text{ion}} = \frac{d}{R_{\text{ion}} \cdot A} \quad (17)$$

where  $d$  indicates the thickness of the polymer film and  $A$  specifies the film area.

## 4. Applications of conducting polymers in electrochemical energy storage devices

Energy storage systems must accommodate various demands depending on their application, whether providing high energy output over short durations (as seen in electric vehicles during acceleration) or delivering continuous energy over extended periods (as required for grid applications). Among the diverse available energy storage technologies, electrochemical energy

storage devices have quickly become prominent across multiple applications due to their superior attributes, including high coulombic efficiency, elevated energy and power densities, scalability, modularity, and rapid response capabilities. Transition metal oxides and conjugated conducting polymers have attracted significant attention in the field of electrochemical energy storage devices. Conjugated conducting polymers offer key advantages for commercial electrochemical energy storage, such as inherent flexibility, unique pseudocapacitive behavior, dual ionic/electronic conduction, fast doping/dedoping dynamics, bulk ionic intercalation, high specific capacity, and excellent structural and thermal stability. Moreover, the commercialization of electrochemical energy storage devices requires large-scale production, and the CVD process is a robust manufacturing method well-suited for large-scale production *via* roll-to-roll (R2R) processing. The R2R CVD technique is already widely employed in the large-scale fabrication of semiconductors,<sup>144</sup> and high-temperature superconductors.<sup>210–216</sup> Similarly, the commercialization of conjugated conducting polymers can be achieved through the R2R oCVD process, which is currently in the early stages of research and development.<sup>2,144,146,201,217</sup>

### 4.1. Lithium-based batteries

The large-scale adoption of electric vehicles (EVs), electronics, and drone applications has created an urgent demand for high-energy, durable, and cost-effective lithium-ion batteries (LIBs).<sup>32,218</sup> Lithium-based (Li-ion and Li-metal) batteries have become a cornerstone of modern electrochemical energy storage technology, offering superior energy density, higher efficiency, and extended longevity. Moreover, lithium-ion batteries (LiBs) are the most advanced secondary (rechargeable) batteries currently available for industrial-scale production. The outstanding attributes of Li-ion batteries include higher energy and power densities, greater thermal stability, low self-discharge rates, and extended cycle life.<sup>219</sup> A typical Li-ion battery comprises several key components, including a positive electrode (cathode), a negative electrode (anode), electrolytes, a separator, current collectors, electrolyte additives, and binders.

The active materials used as cathodes in Li-ion batteries typically consist of lithium transition metal layered oxides, such as lithium cobalt oxide (LCO), lithium manganese oxide (LMO), lithium iron phosphate (LFP), lithium nickel cobalt aluminum oxide (LNCA), and lithium nickel manganese cobalt oxide (LNMC). The atomic percentages of the constituent elements in the cathode can be adjusted to develop desirable characteristics. The most commonly used material for anodes in Li-ion batteries is graphite, which is typically coated on copper foil. However, graphite is susceptible to low usable specific capacity due to the limited intercalation of Li between adjacent graphene planes.<sup>220</sup> To develop anodes with higher energy and power densities, researchers are exploring materials such as silicon (Si), phosphorus (P), and lithium (Li), as well as their composites with graphite.<sup>220</sup>

The electrolytes in Li-ion batteries typically contain lithium salts such as lithium hexafluorophosphate (LiPF<sub>6</sub>), lithium

perchlorate ( $\text{LiClO}_4$ ), and lithium hexafluoroarsenate ( $\text{LiAsF}_6$ ), which are dissolved in organic solvents like ethyl methyl carbonate, dimethyl carbonate, diethyl carbonate, propylene carbonate, and ethylene carbonate.<sup>221</sup> To prevent unwanted side reactions and decomposition of the electrolytes, a variety of additives have been utilized, including vinylene carbonate, fluoroethylene carbonate, ammonium perfluoro-caprylate, and 4-propyl-[1,3,2]dioxathiolane-2,2-dioxide (PDTD).<sup>222,223</sup>

The separator, serving as a safety barrier between the anode and cathode, is electronically insulating but permeable to Li-ions and is typically made from polyethylene and polypropylene. Aluminum foils are commonly used as current collectors for the cathode, while copper foils are employed for the anode, both of which are ultimately connected to the battery terminals. Conductive additives like graphene are often incorporated into cathodes. The common binder supporting matrices include carboxymethyl cellulose (CMC), and polyvinylidene fluoride (PVDF) with *N*-methyl-2-pyrrolidone (NMP) as the solvent.<sup>224</sup>

The working principle of Li-ion batteries relies on the shuttling of lithium ions between the cathode (positive electrode) and anode (negative electrode) through the electrolyte.<sup>221</sup> During charging,  $\text{Li}^+$  ions de-intercalate from the cathode, diffuse through the electrolyte, pass through the nano-porous separator, and intercalate into the anode. Concurrently, electrons travel through the external circuit to maintain electrical neutrality. During discharge, lithium atoms in the anode release electrons, becoming  $\text{Li}^+$  ions that migrate back to the cathode. Simultaneously, electrons flow from the anode to the cathode *via* the external circuit, providing electrical energy. The movement of  $\text{Li}^+$  ions during battery operation generates significant heat due to Joule heating and the chemical energy involved in the charge/discharge processes. While heat generation is an intrinsic aspect of battery function, insufficient dissipation pathways in certain charge/discharge states can lead to excessive heating, posing a serious risk to battery safety.<sup>225</sup>

A current major focus in Li-ion batteries is enabling fast charging while maintaining rate capacity, which is a crucial requirement for the widespread adoption of electric vehicles (EVs).<sup>226,227</sup> The target benchmark for EV batteries is to achieve a fast charging rate that allows for 80% battery capacity to be replenished in 10 minutes or less. The current rate (C-rate) is defined as the charging or discharging current density required to achieve or utilize the theoretical capacity of a battery within  $1 \text{ C}^{-1} \text{ hour}^{-1}$ .<sup>228</sup> The C-rate metric measures the rate at which a battery is charged or discharged, normalized against its maximum capacity. Despite the challenges, achieving fast charging (high C-rate charging) remains a key requirement for advancing Li-ion battery technology. However, several issues associated with fast charging, such as structural instability, poor charge transport kinetics, and accelerated degradation, need to be addressed.<sup>229</sup>

**4.1.1. Solid electrolyte interphase (SEI) formation.** Lithium (Li) is considered as one of the most promising materials for anodes due to its high theoretical capacity ( $3861 \text{ mA h g}^{-1}$ ) and low electrochemical potential ( $-3.04 \text{ V vs. standard hydrogen}$

electrode (SHE)).<sup>220</sup> However, its high chemical reactivity leads to the formation of a solid electrolyte interphase (SEI) layer, which adversely affects device performance. The formation of the SEI layer during the initial cycles results in the irreversible consumption of Li-ions and the decomposition of electrolytes, leading to approximately 10% degradation of the initial capacity.<sup>35</sup> Additionally, Li metal particles tend to form dendrites upon deposition, which can result in short circuit failure, thermal runaway, and explosion hazards.<sup>230</sup> One of the major safety concerns related to Li metal batteries is the formation of dendrites, which occur due to disordered Li nucleation on the unstable SEI, which is formed between the highly reactive Li metal and the electrolyte.<sup>231-233</sup> In general, battery systems based on  $\text{Li}^+$ ,  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Mg}^{2+}$ , and  $\text{Ca}^{2+}$  ions including solid-state batteries (SSBs), suffer from issues like dendrite growth and electrode-electrolyte interface instabilities, indicating the crucial need for engineering of the SEI layer for their optimization.<sup>234-237</sup>

The SEI layer is a passivation layer that forms on the anode at the electrode-electrolyte interface when the redox potential of the electrode falls outside the electrochemical stability window of the electrolyte.<sup>238</sup> The SEI layer results from the reaction between a thermodynamically unstable electrode and the electrolyte during the initial cycles, comprising decomposition products of the electrolyte. Due to its complex microstructure and chemical composition, the ionic conduction within the SEI involves multiple pathways: (i) interstitial and vacancy mechanisms in the inorganic regions, (ii) inter- and intrachain hopping in the polymeric regions, (iii) conduction through grain boundaries, and (iv) surface diffusion in highly porous SEIs. Ion diffusion in the inorganic component of the SEI typically serves as the rate-limiting step.

Optimizing the anode surface area is crucial for enhancing both energy and power density in Li-ion batteries with regard to the stable SEI formation. Minimizing the anode surface area is advantageous for energy density, as it reduces lithium consumption for SEI formation; however, it is detrimental to power density due to solid-state diffusion limitations. Conversely, increasing the anode surface area improves power density but consumes more Li for SEI formation, thereby reducing energy density.<sup>35</sup> Introducing an artificial SEI layer is a sophisticated approach to enhancing overall battery performance by stabilizing the Li-metal anode. This surface engineering method, which utilizes fluorides, phosphates, and polymers, has been shown to improve electrode stability, increase electronic and ionic conductivity, and suppress the dissolution of transition-metal elements from the cathode.<sup>239</sup>

The artificial SEI film acts as a protective layer on the surface of the Li metal electrode, replicating the functionalities of the SEI layer. By regulating  $\text{Li}^+$  deposition and mechanically suppressing Li dendrite growth, the ionic conductive polymeric thin film effectively enhances the electrochemical performance of Li-metal batteries. Polymer materials with excellent ionic conductivity, superior processability, and high flexibility are considered ideal for artificial SEI films.<sup>239</sup> Conducting polymer thin films, such as polypyrrole (PPy), can function as an



artificial SEI layer to achieve a stable, dendrite-free Li metal anode by allowing Li ions to interact with the pyrrole ring through cation- $\pi$  interactions, resulting in the dispersion of Li salt within the PPy thin film.<sup>36</sup> PEDOT-based copolymers, PEDOT *co*-poly(ethylene glycol) (PEDOT-*co*-PEG), when deposited on the surface of a lithium anode, have been reported to act as protective layers, forming a stable and less resistive SEI layer while suppressing the growth of Li dendrites (Fig. 9a).<sup>240</sup> The utilization of a layered conductive polyaniline (LCP) coating on Si nanoparticles, *via* a trimethoxysilyl propyl aniline (TMSPA) bridge, forms an LCP-integrated hybrid SEI layer with high conductivity and mechanical stability, enhancing cycling stability and maintaining high coulombic efficiency for Si anodes (Fig. 9b).<sup>241</sup>

**4.1.2. Utilization of conducting polymer coatings on anodes in Li-ion batteries.** Although graphite has traditionally been used as the anode in Li-ion batteries, it has several limitations, including low intrinsic capacity, slow Li-ion intercalation kinetics from the graphite bulk phase, and capacity loss during high-current charging.<sup>196</sup> As Li-ion batteries approach their theoretical limits with graphite-based anodes, significant attention has recently shifted towards batteries with lithium metal anodes (such as lithium-air (Li-O<sub>2</sub>) batteries,<sup>242</sup> and lithium-sulfur (Li-S) batteries<sup>55</sup>), as well as the use of silicon in lithium-silicon (Li-Si) batteries.<sup>243</sup> To develop Li-ion batteries with higher energy density, silicon (Si) has garnered significant attention as a prospective anode material due to its

high theoretical gravimetric specific capacity of  $\sim 4200$  mA h g<sup>-1</sup>, compared to  $\sim 370$  mA h g<sup>-1</sup> for graphite.<sup>243</sup> Additionally, Si is the second-most abundant element in the Earth's crust, is environmentally friendly, and has a relatively low discharge voltage.<sup>244</sup> However, Si anodes face challenges such as large volumetric changes during lithiation/delithiation processes, which result in fractures, loss of electrical contact, and unstable SEI growth, ultimately leading to poor capacity retention.<sup>245</sup>

Addressing the significant volumetric expansion of Si anodes requires the development of a stable SEI to prevent interfacial side reactions. Applying conducting polymer coatings to porous Si anodes significantly improves electrochemical performance by forming a conformal surface coating that stabilizes the SEI layer, enhances electronic conduction, improves rate capability and cycling stability, increases volumetric tolerance, enhances electrolyte uptake, and facilitates electron transport.<sup>12,37,38,246,247</sup>

Yu *et al.*<sup>12</sup> implemented an n-doped, porous biphenyl-polyoxadiazole (bPOD) conducting polymer layer on a Si anode, observing significant improvement in its rate capacity due to the hybrid ionic-electronic conductivity of the bPOD porous layer and better control of SEI formation by inhibiting direct exposure between the Si and electrolyte (Fig. 10a). Furthermore, the Si-bPOD electrode exhibited higher coulombic efficiency and specific capacity compared to its pristine counterparts, with excellent capacity retention of 85% and 80% after 500 and 800 cycles, respectively (Fig. 10b).<sup>12</sup> The construction of

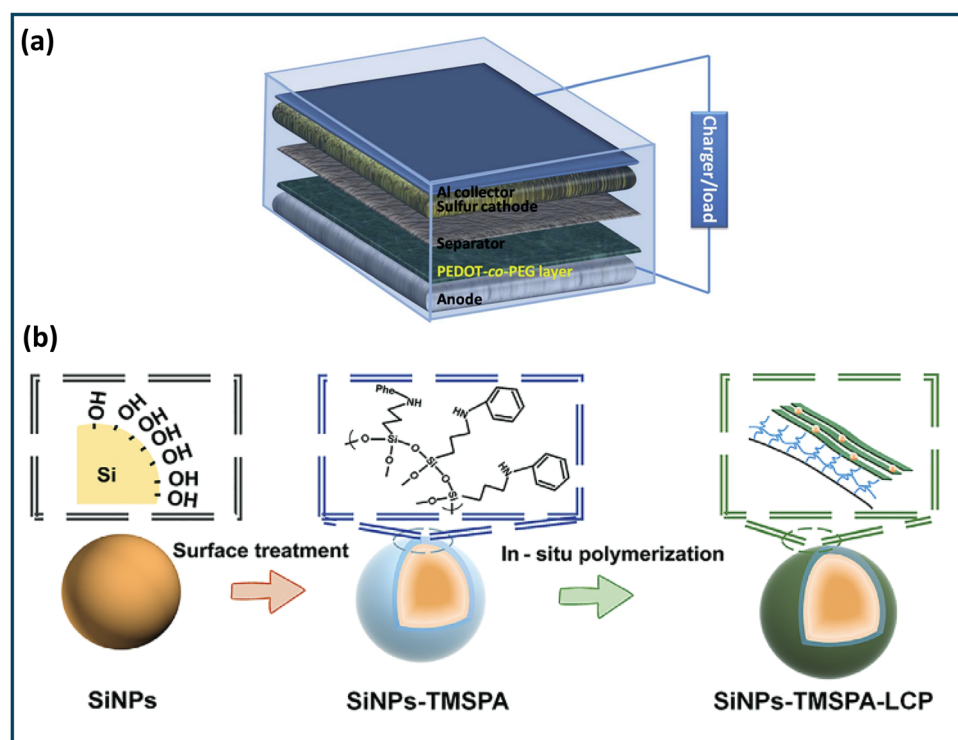
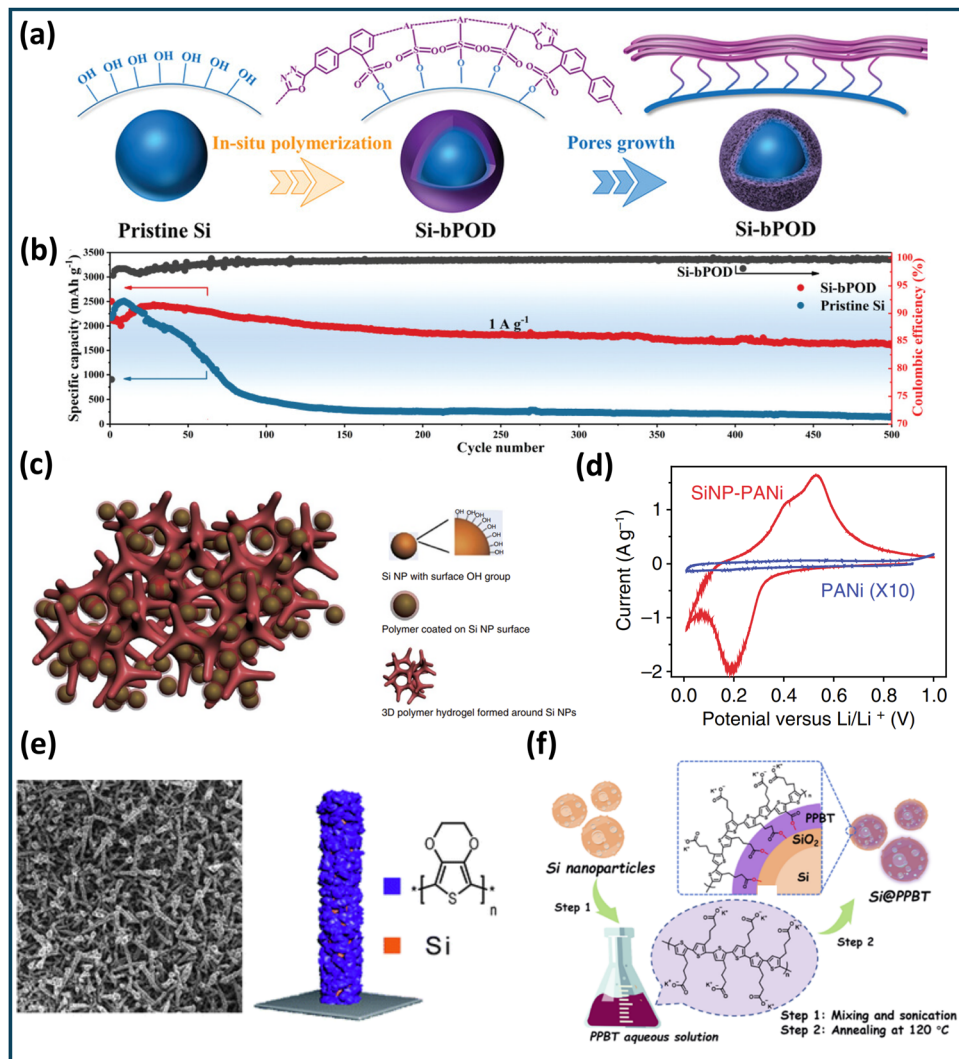


Fig. 9 Artificial solid-electrolyte interface (SEI) with conducting polymers for Li-based batteries. (a) Schematic representation of a Li-S battery featuring a lithium sheet anode protected by a PEDOT-*co*-PEG layer.<sup>240</sup> (b) Schematic representation of Si nanoparticles (SiNPs) modification through *in situ* polymerization to form a layered conductive polyaniline (LCP) coating.<sup>241</sup> (a) Reproduced with permission.<sup>240</sup> Copyright 2014, Royal Society of Chemistry. (b) Reproduced with permission.<sup>241</sup> Copyright 2022, Wiley.





**Fig. 10** Utilization of conducting polymers in anodes for Li-ion batteries. (a) Schematic illustration of fabricating a porous bPOD layer on a Si electrode.<sup>12</sup> (b) Long-term cycling performance comparison between pristine Si and Si-bPOD electrodes.<sup>12</sup> (c) Schematic illustration of a PANI hydrogel 3D network encapsulating Si nanoparticles.<sup>246</sup> (d) CV diagram for SiNP-PANI hydrogel and PANI hydrogel within the potential range of 0.01–1 V vs. Li/Li<sup>+</sup>.<sup>246</sup> (e) Morphology and schematic illustration of PEDOT-coated Si nanowires (SiNWs).<sup>247</sup> (f) Schematic illustration of procedure for fabricating PPBT conducting polymer on Si anode.<sup>38</sup> (a) and (b) Reproduced with permission.<sup>12</sup> Copyright 2023, Wiley. (c) and (d) Reproduced with permission.<sup>246</sup> Copyright 2013, Nature. (e) Reproduced with permission.<sup>247</sup> Copyright 2012, Royal Society of Chemistry. (f) Reproduced with permission.<sup>38</sup> Copyright 2012, American Chemistry Society.

conducting polymer hydrogels made of PANI over Si nanoparticles (SiNPs) exhibited an enhancement in current density, with approximately two orders of magnitude higher performance than the dried PANI hydrogels (Fig. 10c and d).<sup>246</sup> The utilization of a PEDOT coating on Si nanowires (SiNWs) improves cycling stability (the capacity retention after 100 cycles increased from 30% to 80%) and maintains structural integrity, preventing the cracks commonly found in non-coated SiNWs after cycling (Fig. 10e).<sup>247</sup> The utilization of a poly[3-(potassium-4-butanoate)thiophene] (PPBT) conducting polymer layer on a Si anode significantly enhanced rate capability and cycling stability by increasing tolerance of volumetric changes, improving electrolyte uptake, facilitating electron transport, and stabilizing the SEI layer (Fig. 10f).<sup>38</sup>

A few other promising anode materials, such as Germanium (Ge), manganese-based oxides ( $Mn_xO_y$ ), and aligned carbon nanotubes (ACNTs), have also been combined with conducting polymers to mitigate their inherent limitations and enhance electrochemical performance.<sup>248–250</sup> Ge has significantly higher conductivity and Li-ion diffusivity (about 400 times higher) than Si at room temperature, but it has a lower specific capacity ( $1600 \text{ mA h g}^{-1}$ ) and also undergoes large volume changes and rapid capacity fading.<sup>251</sup> The deposition of PEDOT:PSS conducting polymer on Ge anodes has demonstrated enhanced rate capability and cycling stability, indicating improved structural integrity and reduced pulverization and exfoliation of the Ge shell.<sup>248</sup> The deposition of PEDOT on  $Mn_2O_3$  nanowires has been shown to reduce surface resistance, prevent the

pulverization of surface electron channels, and enhance cycling stability, achieving a reversible capacity of  $1450 \text{ mA h g}^{-1}$  after 200 cycles.<sup>249</sup>

**4.1.3. Utilization of conducting polymer coatings on cathodes in Li-ion batteries.** As a promising cathode for next-generation lithium-ion batteries, Ni-rich NCM, specifically NCM<sub>811</sub> ( $\text{LiNi}_{0.8}\text{Co}_{0.1}\text{Mn}_{0.1}\text{O}_2$ ), exhibits high energy density with a high specific capacity of  $\sim 210 \text{ mA h g}^{-1}$ , good power density, and reduced cost but faces challenges such as low thermal stability, poor cycle life, and rapid capacity degradation that must be addressed.<sup>32,39</sup> Due to the critical role of the surface/interface in electrochemical degradation, surface coating of NCM cathodes has been intensively investigated to enhance cycling performance using various coatings, including metal oxides (e.g.,  $\text{TiO}_2$ ,  $\text{ZrO}_2$ , and  $\text{Al}_2\text{O}_3$ ),<sup>40,41,252</sup> metal fluorides (e.g.,  $\text{AlF}_3$ ,  $\text{CaF}_2$ , and  $\text{LiF}$ ),<sup>42,43</sup> metal phosphates (e.g.,  $\text{Li}_3\text{PO}_4$ ,  $\text{AlPO}_4$ ,  $\text{FePO}_4$ ),<sup>44,45</sup> Li-ion conductive coating ( $\text{LiTi}_2\text{O}_4$ ,  $\text{Li}_2\text{ZrO}_3$  and  $\text{Li}_4\text{Mn}_5\text{O}_{12}$ ),<sup>46,47</sup> and conducting polymers (e.g., PEDOT, PANI, PPy).<sup>48,49</sup> Compared to inorganic coatings, conducting polymers, offering high electronic conductivity, environmental stability, and low cost, can stabilize cycling and improve rate capability.

Conducting polymers have been effectively utilized as coating layers on various cathode surfaces in lithium-ion batteries, acting as barriers between the cathode and electrolyte to prevent undesirable side reactions, provide thermal and structural stability, enhance long-term cycling stability, and improve high-rate performance.<sup>15,39,253–255</sup> The application of a conformal PEDOT coating on the cathode surface has been shown to decrease cell impedance, reduce interfacial charge transfer resistance between the electrode and electrolyte, and improve electron transport to the current collectors.<sup>2,256</sup> The favorable chemical and coordination bonds (between oxygen, sulfur (S) in the PEDOT structure, and manganese (Mn) in the cathode structure) stabilize the cathode surface, inhibiting Mn dissolution and preventing SEI growth, which are primary aging mechanisms.<sup>256,257</sup>

Uniformly coating NCM<sub>811</sub> with conducting polymers is challenging due to poor bonding between the materials. The oCVD method is a versatile manufacturing technique for fabricating highly conformal, ultrathin coatings of conducting polymers on various cathode materials in Li-ion batteries, such as NMC and LMO, resulting in enhanced stability and electrochemical performance.<sup>2,8,15,255,256</sup> These improvements are attributed to oCVD's ability to conformally coat individual particles beyond the surface layer, significantly enhancing crystal and interfacial stability for all particles by providing a stable cathode–electrolyte interface, suppressing microcrack formations, and preventing irreversible reactions.

Gan *et al.*<sup>253</sup> demonstrated that a polyvinylpyrrolidone (PVP)-induced PANI coating on NCM<sub>811</sub> cathode surfaces enhanced stability and rate performance, achieving the highest capacities at various scan rates by preventing side reactions and facilitating fast diffusion channels for Li-ions (Fig. 11a). The surface of NCM<sub>811</sub> was modified with a PANI–PEG hybrid layer to simultaneously utilize PANI's high electronic conductivity

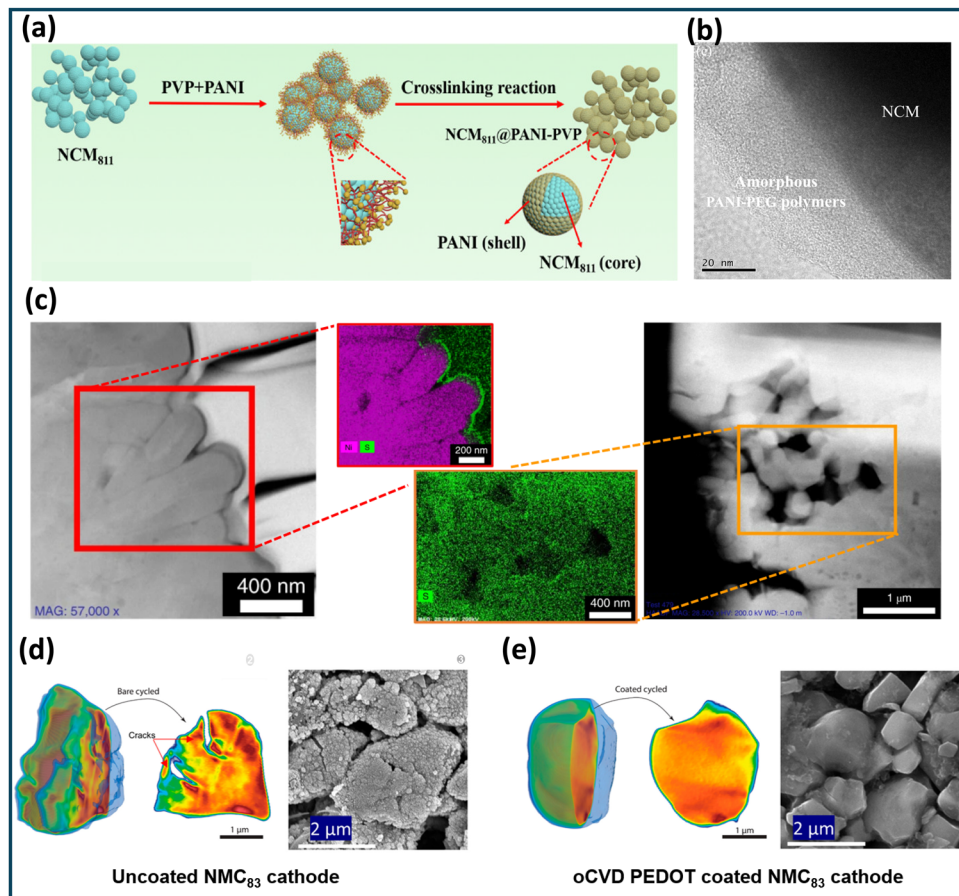
and PEG's exceptional ionic conductivity, creating a facile diffusion pathway for Li ions and resulting in improved capacity and cycling performance, particularly at high C-rates (Fig. 11b).<sup>254</sup> Applying a highly conformal oCVD PEDOT layer on various NCM cathodes enhanced thermal and structural stability, improving capacity retention by suppressing undesired structural evolution during charge–discharge cycles (layered to spinel structure), as shown in Fig. 11c.<sup>15</sup>

The thermal stability of NMC can be assessed by comparing oxygen release profiles for highly charged pristine and PEDOT-coated NMC, to identify side reactions on the coated cathode surface.<sup>258</sup> Previous studies have shown that oxygen species (*i.e.*,  $\text{O}_2^-$ ,  $\text{O}^-$ ,  $\text{O}_2^{2-}$  and  $\text{O}_2$ ) are released during the phase transition of charged NMC upon heating, which can cause thermal runaway by reacting with the flammable electrolyte.<sup>258</sup> The deposition of a conformal oCVD PEDOT thin film on the single-crystal Ni-rich NMC cathode (NMC<sub>83</sub>:  $\text{LiNi}_{0.83}\text{Mn}_{0.1}\text{Co}_{0.07}\text{O}_2$ ) significantly improved capacity retention, achieving 85.3% after 100 cycles at a 0.5C rate, compared to 59.6% for the pristine electrode.<sup>255</sup> The post-cycled microstructure of the pristine NMC<sub>83</sub> cathode exhibited substantial crack formation due to heterogeneous phase transitions from layered to spinel and rock-salt structures, leading to internal strain and capacity loss; whereas PEDOT-coated NMC<sub>83</sub> retained its initial structure (Fig. 11d and e).<sup>32</sup>

Another cathode material for next-generation Li-ion batteries, the Li-rich cathode ( $\text{Li}_{1.2}\text{Ni}_{0.2}\text{Mn}_{0.6}\text{O}_2$ ), demonstrates a higher reversible capacity ( $\sim 250 \text{ mA h g}^{-1}$ ) compared to traditional cathodes ( $\text{LiCoO}_2$ ,  $\text{LiFePO}_4$ , and  $\text{LiMn}_2\text{O}_4$ ) with capacities around  $170 \text{ mA h g}^{-1}$ .<sup>259,260</sup> However, side reactions at high voltages lead to undesired phase transitions, and the insulating nature of  $\text{Li}_2\text{MnO}_3$  formed during cycling results in poor rate capability and low cycling performance.<sup>259</sup> To address these issues, depositing PEDOT:PSS with optimized thickness on the Li-rich cathode improves discharge capacity and rate capability due to the conductive nature of PEDOT and its role as a barrier between the cathode and electrolyte, preventing undesired reactions.<sup>259</sup> Su *et al.*<sup>261</sup> deposited various polymeric materials, including PEDOT, on the  $\text{LiCoO}_2$  (LCO) cathode surface and observed significant enhancements in  $\text{Li}^+$  kinetics, resulting in increased rate capability at C-rates  $> 5\text{C}$  and enhanced cycle life at high voltage of 4.5 V (Fig. 12a and b).

The lithium–sulfur (Li–S) battery is considered one of the most promising Li–metal-based battery technologies, offering a high gravimetric energy density of  $\sim 2600 \text{ W h kg}^{-1}$  and serving as a low-cost energy storage method. However, the practical utilization of Li–S batteries faces several challenges: (i) poor rate capability and limited electrode rechargeability due to the insulating nature of  $\text{S}_8$  and  $\text{Li}_2\text{S}$ , (ii) rapid capacity degradation resulting from the loss or dissolution of active material, as soluble lithium polysulfides ( $\text{Li}_2\text{S}_n$ , where  $3 \leq n \leq 6$ ) migrate from the cathode to the electrolyte and anode, causing electrolyte degradation and a non-uniform anode microstructure, and (iii) significant volume increase ( $\sim 80\%$ ) during charging, as  $\text{S}_8$  converts to low-density  $\text{Li}_2\text{S}$ .<sup>33,50–52</sup> To enhance active material utilization and cycling stability of the sulfur cathode, various





**Fig. 11** Utilization of conducting polymer coatings on various Ni-rich NCM cathode surfaces. (a) Schematic illustration showing the mechanism of developing the PANI layer on the surface of NCM<sub>811</sub> cathode using PVP surfactant.<sup>253</sup> (b) TEM images of NCM<sub>811</sub> coated with PANI-PEG copolymer.<sup>254</sup> (c) STEM-HAADF images with corresponding EDS mappings of the secondary particles surface (left) and inner primary particles surface (right) of oCVD PEDOT-coated NCM.<sup>15</sup> Ptychographic X-ray computed tomography and corresponding SEM images showing particles after cycling between 2.8 and 4.6 V for 200 cycles of (d) pristine NMC<sub>83</sub> cathode and (e) oCVD PEDOT coated NMC<sub>83</sub> cathode.<sup>32</sup> (a) Reproduced with permission.<sup>253</sup> Copyright 2019, American Chemistry Society. (b) Reproduced with permission.<sup>254</sup> Copyright 2018, American Chemistry Society. (c) Reproduced with permission.<sup>15</sup> Copyright 2019, Nature. (d) and (e) Reproduced with permission.<sup>32</sup> Copyright 2022, American Chemical Society.

advanced sulfur composites have been employed, including dispersing sulfur in conductive mediums such as carbon-based materials,<sup>53,54</sup> metal oxides/carbides, and conducting polymers like PANI, PPy, and PEDOT.<sup>55,56,262,263</sup> Coating the sulfur cathode in Li-S batteries with conjugated conducting polymers provides substantial benefits, such as enhanced capacity retention and improved overall capacity by restricting the shuttling of polysulfide intermediates. Additionally, conducting polymers help mitigate electrode degradation caused by the volume expansion of sulfur-based cathode materials due to their self-healing properties.<sup>55,57</sup>

Zhang *et al.*<sup>55</sup> demonstrated a practical Li-S application using pouch cells by depositing a highly conformal oCVD PEDOT layer on sulfur particles, achieving notable capacity retention of 85% after 300 cycles at 0.5C due to the restriction of polysulfide shuttling in the cathode (Fig. 12c and d). Yang *et al.*<sup>262</sup> deposited a layer of PEDOT:PSS across the surface of a mesoporous carbon/sulfur composite (CMK-3/S) cathode and observed that the polymer layer effectively confined the

shuttling of polysulfide intermediates and prevented dissolution during cycling, resulting in enhanced capacity retention and coulombic efficiency (Fig. 12e). To encapsulate sulfur and its polysulfide compounds within the cathode, PANI nanotubes are synthesized with sulfur to form a 3D, cross-linked sulfur-PANI composite, leading to improved cycling stability and rate capability.<sup>263</sup> Xie *et al.*<sup>56</sup> deposited an approximately 60-nm PPy layer on sulfur cathodes *via* *in situ* polymerization, forming S/PPy composites with a core-shell structure, which prevented sulfur particle aggregation and resulted in faster ion transfer and enhanced C-rate performance (Fig. 12f).

**4.1.4. Utilization of conducting polymers as separator and binder materials in Li-ion batteries.** Achieving a uniform Li<sup>+</sup> flux towards the electrode reduces Li dendrite growth. Therefore, optimizing pore density and structure in separators is crucial for the performance of lithium metal batteries, necessitating the development of new materials.<sup>264,265</sup> Conducting polymers in overoxidized or de-doped states, have been used as separator materials in lithium-ion batteries

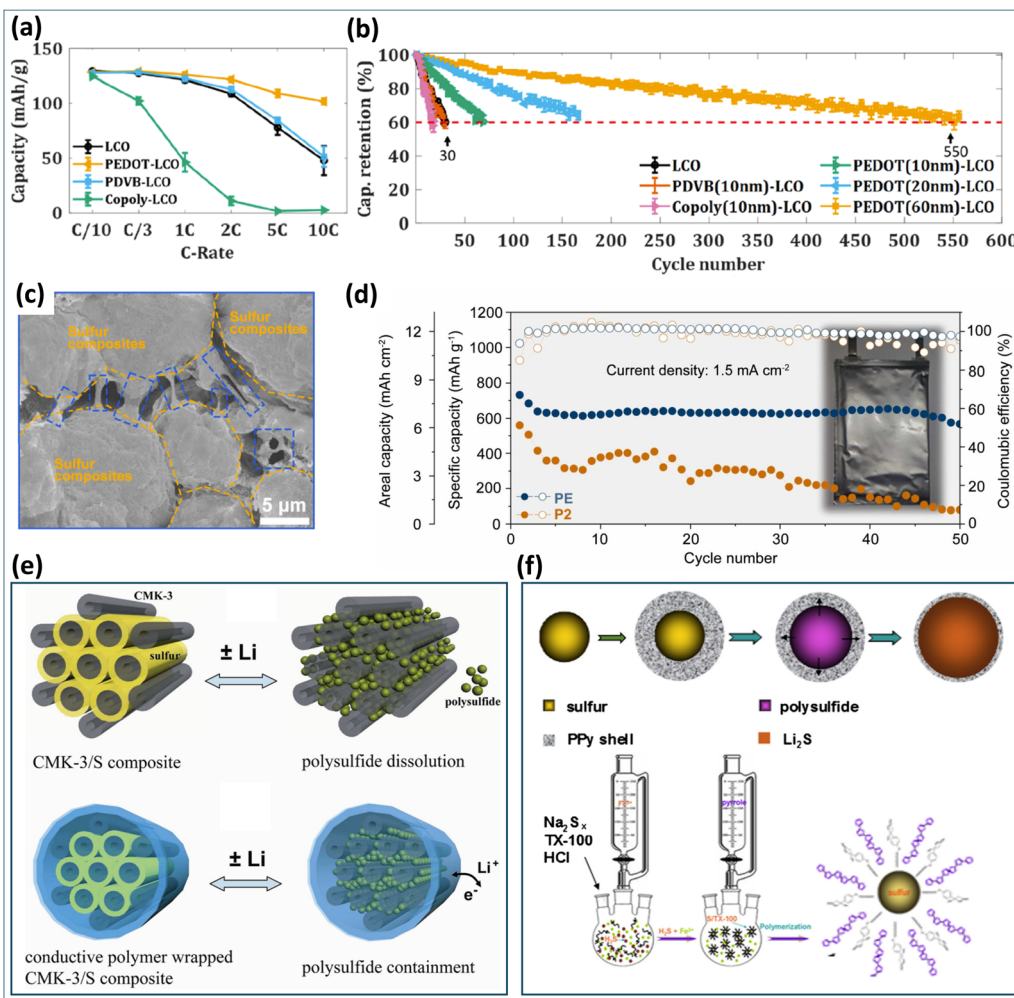


Fig. 12 Utilization of conducting polymer coatings on various sulfur cathodes in Li-S batteries. (a) Comparison of cell capacities for various polymer-coated LCO cathodes.<sup>261</sup> (b) Capacity retention of different LCO cathodes at C/2 within 3.0–4.5 V.<sup>261</sup> (c) SEM images of OCVD PEDOT-coated sulfur cathode.<sup>55</sup> (d) Cycling performance of Li-S pouch cell with the PEDOT-coated sulfur (PE), and pristine sulfur (P2).<sup>55</sup> (e) Schematic illustration showing PEDOT:PSS coating on CMK-3/S composite electrode, blocking polysulfide dissolution.<sup>262</sup> (f) Illustration of the synthesis process for fabricating a PPy layer on sulfur cathodes to form S/PPy composites via *in situ* polymerization.<sup>56</sup> (a) and (b) Reproduced with permission.<sup>261</sup> Copyright 2021, American Chemical Society. (c) and (d) Reproduced with permission.<sup>55</sup> Copyright 2023, Elsevier. (e) Reproduced with permission.<sup>262</sup> Copyright 2011, American Chemical Society. (f) Reproduced with permission.<sup>56</sup> Copyright 2016, Elsevier.

because of their numerous advantageous properties, such as thermal stability, mechanical flexibility, excellent electrolyte wettability, high ionic conductivity, and reversible doping/de-doping capabilities.<sup>264,266</sup> The commonly used separator material, commercial porous polyolefin film, can be enhanced by incorporating poly(3-butyl thiophene) (P3BT) conducting polymer into its micropores, serving as a self-actuated safety mechanism for overcharge protection.<sup>266</sup> The de-doped polymer utilized in the separator permits electrolyte flow while blocking electron transport. However, during overcharging, the increased cathode potential oxidizes the polymer, making it conductive and forming a bridge between the anode and cathode to create a bypass current and prevent overcharging. Wang *et al.*<sup>264</sup> utilized overoxidized porous PPy paper as a mesoporous separator for Li-metal batteries, and achieved homogeneous current distribution due to uniform pore size and high electrolyte wettability.

Additionally, conducting polymers and their composites can serve as binder materials for Li-ion battery electrodes because they provide flexible electronic contacts, maintain adhesive strength during volumetric changes, and act as both binder and conductive additive, enabling high active material loading.<sup>267–269</sup> Zhong *et al.*<sup>268</sup> used a water-soluble conductive composite binder with carboxymethyl chitosan (CCTS) as the binder and PEDOT:PSS as the conductive agent for lithium iron phosphate (LFP) cathodes, and achieved homogeneous, continuous conducting bridges and improved rate performance. Das *et al.*<sup>267</sup> demonstrated a carbon black-free LFP electrode in which PEDOT:PSS served as both binder and conducting additive, resulting in enhanced rate capacity and cycling stability. Combining carboxymethyl cellulose (CMC) with PEDOT:PSS as a water-soluble composite binder for a Si anode results in superior cycling and rate capacity,

as well as faster kinetics, compared to the acetylene black/CMC binder.<sup>269</sup>

#### 4.2. Utilization of conducting polymer coatings in supercapacitors

Capacitance refers to a material's ability to store electric charge, and devices that store energy in the form of charges are commonly known as capacitors. Supercapacitors (also referred to as ultracapacitors or electrochemical capacitors) are energy storage devices that offer significantly higher capacitance compared to traditional dielectric capacitors. Supercapacitor devices are engineered to fill the gap between conventional capacitors and rechargeable secondary batteries and balancing

specific power with energy density, as depicted in the Ragone plot in Fig. 13a. Supercapacitors have gained considerable attention for their exceptional capacitance (100–1000× higher than a conventional capacitors), high power density, rapid charge/discharge times (1–10 second), and extended cycling life (>30 000 h), making them ideal for applications such as flexible electronics, electric vehicles, and renewable energy.<sup>7,20,58,270</sup> Supercapacitors have found applications in several crucial areas, including electric vehicles, power grids, and aerospace technology.<sup>2,271</sup>

Supercapacitors typically consist of a current collector, an electrode active material (which may be submerged in a liquid electrolyte or exist as a solid-state in solid-state capacitors), and

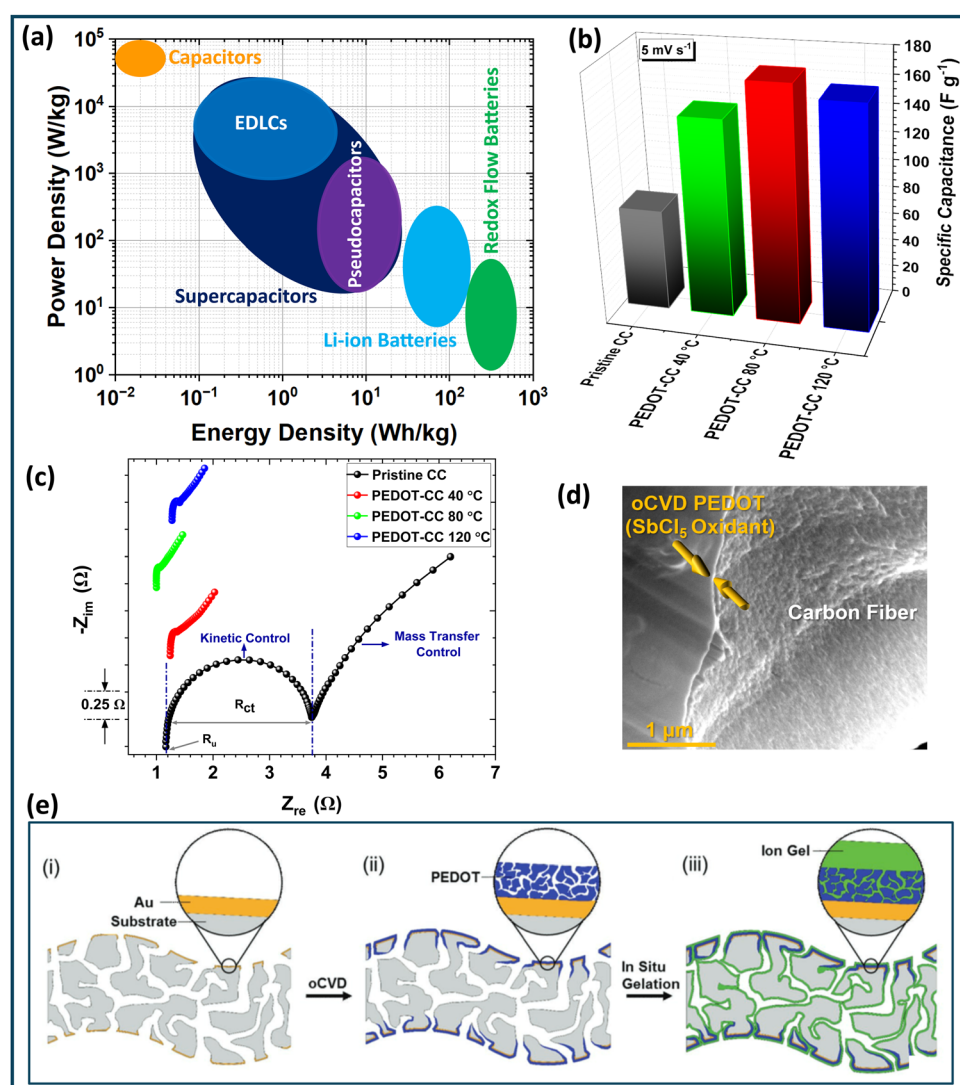


Fig. 13 Utilization of PEDOT conducting polymer coatings in supercapacitors. (a) Ragone plot illustrating the relative positions of EDLC and pseudocapacitors, showing how supercapacitors bridge the gap between batteries and traditional capacitors in terms of energy and power density. (b) Specific capacitance values for carbon cloth (CC) and oCVD PEDOT-coated carbon cloth (PEDOT-CC) electrodes. The oCVD PEDOT thin films fabricated at different deposition temperatures.<sup>10</sup> (c) Nyquist plots for pristine carbon cloth and PEDOT-coated carbon cloths fabricated at different deposition temperatures.<sup>10</sup> (d) Magnified SEM image showing an individual carbon fiber with a highly conformal oCVD PEDOT coating.<sup>10</sup> (e) Schematic illustrations of the intermediate and final configurations of a vapor-printed solid-state supercapacitor, where gray for porous substrate, gold for current collector, blue for PEDOT, and green for ion gel electrolyte.<sup>272</sup> (b)–(d) Reproduced with permission.<sup>10</sup> Copyright 2024, Wiley. (e) Reproduced with permission.<sup>272</sup> Copyright 2017, Wiley.



a porous dielectric separator between the two electrodes.<sup>2</sup> Two key performance metrics for supercapacitors are energy density (or specific energy) and power density (or specific power), which are evaluated in relation to parameters such as mass, volume, or area. The energy density ( $E_{D,X}$ ) and power density ( $P_{D,X}$ ) in supercapacitors are defined as follows:<sup>181</sup>

$$E_{D,X} = \frac{1}{2} C_{S,X} \Delta V^2 \quad (18)$$

$$P_{D,X} = \frac{E_{D,X}}{\Delta t} \quad (19)$$

where parameter X defines the methods (e.g. X = m for mass or X = v for volume),  $C_{S,X}$  denotes the specific capacity,  $\Delta V$  represents the potential window,  $\Delta t$  describes the discharge time.

Depending on the energy storage mechanism, supercapacitors can be classified into three categories: (i) electrochemical double layer capacitors (EDLCs), which store charge electrostatically by forming a non-faradaic double layer at the electrode surface, (ii) pseudocapacitors, which store charge electrochemically through faradaic electron transfer *via* fast redox reactions, and (iii) hybrid supercapacitors, which combine both EDLC and pseudocapacitor charge storage mechanisms, and utilizing hybrid or asymmetric electrodes.<sup>273</sup>

In EDLCs, energy storage is facilitated by a non-faradaic process, where charge accumulates at the electrode-electrolyte interface, forming an electrochemical double layer that is responsible for the capacitance. In supercapacitors, capacitance increases with larger electrode surface areas, leading to the exploration of high-surface-area materials like carbon nanotubes (CNTs) and other nanoporous structures as potential electrodes.<sup>2</sup> Carbon-based materials with high surface areas, such as activated carbon, CNTs, and graphene are widely used as electrode materials in EDLCs. The specific capacitance of EDLCs is determined by the exposed surface area and charge storage capacity of the carbon-based electrode materials, but it is inherently limited by the non-faradaic charge storage mechanism. In contrast, pseudocapacitors offer 10 to 100 times higher specific capacitance due to the faradaic (redox-reaction) charge storage mechanism that occurs within the bulk of the active materials.<sup>3</sup> EDLCs offer higher power density but lower energy density compared to rechargeable batteries. Pseudocapacitors generally exhibit significantly higher specific capacitances compared to carbon materials utilizing the EDLC mechanism, as the entire bulk of the pseudocapacitive material, rather than just the surface layer, participates in redox reactions.<sup>10,20</sup> Pseudocapacitors, on the other hand, typically achieve significantly greater energy density than EDLCs, positioning them in an intermediate specific energy range between EDLCs and rechargeable batteries.<sup>2</sup>

Recently, various flexible supercapacitor designs have been explored, including wire-shaped supercapacitors,<sup>274</sup> fiber-shaped coaxial supercapacitors,<sup>275</sup> carbon based textiles,<sup>10</sup> and highly stretchable fiber-shaped supercapacitors.<sup>276</sup> High surface area carbon-based materials like activated carbon (AC), carbon nanotubes (CNTs), graphene, graphene oxide (GO),

carbon cloths (CCs), and carbon fibers (CFs) are commonly employed as active electrode materials in non-faradaic EDLCs. In contrast, conjugated conducting polymers and transition metal compounds (TMCs) such as metal oxides (e.g., RuO<sub>2</sub>, MnO<sub>2</sub>, V<sub>2</sub>O<sub>5</sub>, and Fe<sub>2</sub>O<sub>3</sub>), metal sulfides, metal nitrides, and metal carbides are commonly and effectively employed as electrode materials for faradaic pseudocapacitors.<sup>2</sup> The mechanical stability of conjugated conducting polymers is effectively utilized in the fabrication of flexible supercapacitors due to their simple synthesis, redox properties, and electrical conductivity.<sup>2,20</sup>

Conducting polymers, in particular, have shown great potential as supercapacitor electrode materials due to their unique combination of hybrid ion-electron transfer capabilities, fast doping/de-doping mechanisms, excellent conductivity, mechanical flexibility, and high environmental stability.<sup>2,10</sup> Conducting polymers are promising pseudocapacitor materials for flexible supercapacitors, advancing energy storage devices for future flexible electronics due to their high redox capacitance and inherent elasticity.

The most commonly utilized conducting polymers in supercapacitor applications include PANI, PEDOT, PPy, and PT, which are typically fabricated using solution-based or vapor-based techniques. PANI stands out as a leading candidate among conjugated conducting polymers for supercapacitors due to its excellent pseudocapacitive properties, as demonstrated by PANI nanowire arrays achieving a high specific capacitance of up to 950 F g<sup>-1</sup>.<sup>277</sup> Typically, conjugated conducting polymers possess multiple redox states, allowing for the precise control of electrical conductivity across the entire spectrum from insulator to metal, making them suitable for use in supercapacitors operating *via* a pseudocapacitance mechanism.<sup>2,8,278</sup> Conducting polymers generally exhibit high specific capacitances, such as 1284 F g<sup>-1</sup> for PANI,<sup>59</sup> 480 F g<sup>-1</sup> for PPy,<sup>60</sup> and 210 F g<sup>-1</sup> for PEDOT.<sup>10,61</sup>

PEDOT is a widely used conducting polymers in supercapacitor devices due to its desirable properties such as possessing a wide potential electrochemical window over which the capacitance is high, excellent thermal and chemical stability, high charge mobility and electrical conductivity that induces fast electrochemical kinetics, and excellent cycling stability.<sup>2,20,279</sup> However, PEDOT has a disadvantage of relatively high molecular weight which leads to the approximately low specific capacitance, especially compared to PANI which exhibits the highest theoretical specific capacitance among conjugated conducting polymers.<sup>2</sup>

To address the cycle stability limitations of polyaniline, it is commonly composited with carbonaceous materials such as graphene, activated carbon, or graphite. Carbonaceous materials such as CNTs, graphene, and graphite hold significant potential across various fields due to their exceptional structural, mechanical, and electrical properties. A significant drawback of polyaniline–carbon based composites is their tendency to aggregate during synthesis, which reduces active sites and impairs ion mobility, ultimately compromising supercapacitor performance. To address this issue, researchers have shifted their focus to 3D porous carbon structures.



Supercapacitor devices composed entirely of conducting polymers can be configured in three main types:<sup>2,20,280</sup>

(1) Type I (symmetric): utilizes the same p-type conducting polymer for both electrodes.

(2) Type II (asymmetric): employs two different p-type conducting polymers with distinct electroactivity ranges for the electrodes.

(3) Type III (symmetric): uses the same conducting polymer for both electrodes, but the p-doped form serves as the positive electrode and the n-doped serving as the negative electrode.

Additionally, hybrid (or asymmetric) devices can be constructed with a conducting polymer as the positive electrode and a carbon or lithium-based material as the negative electrode. Among these configurations, the Type III conducting polymer-based supercapacitors offer the most promising theoretical performance due to their ability to maintain both electrodes in highly conductive states, allowing operation over a broader potential range and achieving a significantly higher maximum cell potential ( $V_{max,c}$ ). In Type I and II conducting polymer-based supercapacitors, the positive electrode remains in a fully doped (conductive) state when fully charged, while the negative electrode stays in a de-doped (neutral) state. During discharge, both electrodes shift to partially doped states, which limits the achievable maximum cell voltage. In contrast, Type III conducting polymer-based supercapacitors have both electrodes in highly conductive doped states (p-doped and n-doped) when fully charged, and these can transition to partially p-doped or n-doped states upon discharge, depending on the potential difference between the electrodes. This configuration enables the device to operate within a broader potential window, leading to a significantly higher  $V_{max,c}$ .<sup>7,20</sup> Conjugated conducting polymers can be positively charged (p-doped) or negatively charged (n-doped) through redox reaction and ion insertion from the electrolyte into their matrix to stabilize the charge.<sup>2</sup> However, the practical application of Type III is limited by the challenges associated with fabricating stable n-type conducting polymers.<sup>2,20,33</sup>

Carbon cloth possesses desirable properties for use as flexible electrodes in various energy storage devices but suffer from limited wettability and low specific capacitance.<sup>10</sup> The oCVD PEDOT-coated carbon cloth, fabricated at various deposition temperatures, displayed highly conformal coatings and significantly enhanced specific capacitance, with the highest value of  $170.94 \text{ F g}^{-1}$  achieved in a sample fabricated at  $80^\circ\text{C}$ , representing a 2.3-fold enhancement over the pristine carbon cloth (Fig. 13b-d).<sup>10</sup> PEDOT-coated carbon cloth samples exhibit lower charge transfer resistance compared to the pristine carbon cloth (Fig. 13c), and the optimal performance of oCVD PEDOT-coated carbon cloths at  $80^\circ\text{C}$  is due to a higher ionic dopant concentration and edge-on orientation, enhancing out-of-plane electrical conductivity.<sup>10</sup> Liu *et al.*<sup>272</sup> fabricated the highly conformal oCVD PEDOT thin film on both side of a flexible porous substrate to integrate all components of a supercapacitor into a flexible, porous substrate with a sandwich-like structure (Fig. 13e). This single-step process precisely controlled vapor penetration, enabling the substrate's

interior to function as both a separator and solid-state electrolyte reservoir, resulting in a supercapacitor with high volumetric specific capacitance ( $11.3 \text{ F cm}^{-3}$ ), power density ( $0.42 \text{ W cm}^{-3}$ ), and energy density ( $2.98 \text{ mW h cm}^{-3}$ ).<sup>272</sup>

Conducting polymers can be fabricated into diverse morphologies such as thin films, hydrogels, rods, particles, and conformal nanostructures, each exhibiting unique optical, electrical, mechanical, and electrochemical properties. Numerous studies have indicated a significant correlation between the morphology of conducting polymers and their electrochemical performance. Dubal *et al.*<sup>281</sup> synthesized PPy nanostructures with different morphology (nanobelts, nanobricks, and nanosheets) on steel substrates *via* electrochemical polymerization, finding that PPy nanosheets achieved a higher specific capacitance of  $586 \text{ F g}^{-1}$  at the scan rate of  $2 \text{ mV s}^{-1}$ , compared to nanobricks ( $357 \text{ F g}^{-1}$ ) and nanobelts with the value of  $296 \text{ F g}^{-1}$  (Fig. 14a).

$\text{MnO}_2$  is a promising pseudocapacitive material for high-performance supercapacitors due to its high theoretical specific capacitance, low cost, environmental friendliness, and natural abundance, but its performance is often hindered by poor electronic and ionic conductivities, limiting power density and cycling stability.<sup>282</sup> The incorporation of PEDOT on  $\text{MnO}_2$  has been reported to improve cycling stability and increase capacitance.<sup>282</sup> The graphene/ $\text{MnO}_2$ /PEDOT:PSS (GMP) ternary composite, demonstrating exceptional cycling stability with over 95% capacitance retention after 3000 cycles and high capacitance, attributed to the significant pseudocapacitance of PEDOT:PSS (Fig. 14b).<sup>282</sup>

The integration of PANI nanowires on a flexible, free-standing graphene film with a 3D interconnected porous structure (3D-RGO film) enhances the specific capacitance to  $385 \text{ F g}^{-1}$ , with a rate retention of 94% (Fig. 14c).<sup>271</sup> The fabrication of a three-dimensional (3D) PANI-graphene nanoribbon (GNR)-carbon nanotube (CNT) composite (PANI-GNR-CNT composite) *via* *in situ* polymerization enhances PANI's cycle life, resulting in 89% capacity retention after 1000 cycles.<sup>284</sup> Smolin *et al.*<sup>283</sup> integrated a nanometer-thin film of oCVD PANI onto carbide-derived carbon (CDC) with a bimodal (micro/mesoporous) pore size distribution, enabling PANI to penetrate pores as small as 1.7 nm. The resulting oCVD PANI-integrated CDC supercapacitors achieved a gravimetric capacitance of  $136 \text{ F g}^{-1}$ , more than double that of uncoated CDC electrodes ( $60 \text{ F g}^{-1}$ ) at a scan rate of  $10 \text{ mV s}^{-1}$ , and demonstrated excellent cyclability, with only a 10% decrease in capacitance after 10 000 cycles (Fig. 14d).<sup>283</sup>

#### 4.3. Utilization of conducting polymer coatings in redox flow battery batteries

Redox flow batteries (RFBs) are highly suitable for large-scale, cost-effective electrochemical energy storage due to their simple and modular design, ease of scalability, long-duration energy storage capabilities, rapid response to load demand, high efficiency, enhanced safety compared to conventional sealed battery systems, and long cycle life.<sup>9,62,63</sup> Unlike conventional solid-state batteries, redox flow batteries store redox-



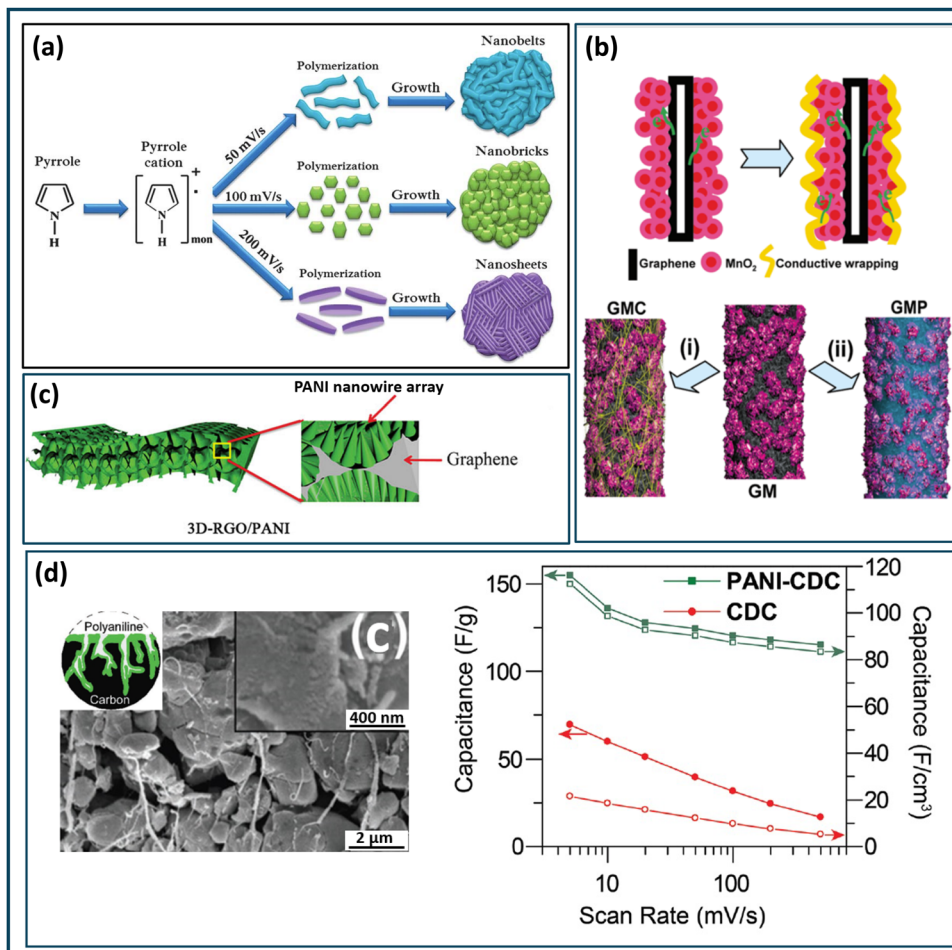


Fig. 14 Utilization of conducting polymer coatings in supercapacitors. (a) Schematic illustration of different PPy nanostructures made of nanobelts, nanobricks, and nanosheets, which are fabricated by the electropolymerization, which yields different specific capacitance.<sup>281</sup> (b) Schematic illustration of graphene/MnO<sub>2</sub>/CNT (GMC) and graphene/MnO<sub>2</sub>/conducting polymer (GMP) systems, showing GM nanostructures wrapped with CNTs or conducting polymers (black: graphene; rose: MnO<sub>2</sub>; yellow: CNTs; blue: conducting polymer).<sup>282</sup> (c) Schematic illustration of graphene composites formed through the polymerization of PANI nanowires.<sup>271</sup> (d) Cross-sectional SEM images of PANI-coated Mo<sub>2</sub>C-CDC electrodes (left) and capacitance comparison of pristine and PANI-coated Mo<sub>2</sub>C-CDC electrodes across various scan rates (right).<sup>283</sup> (a) Reproduced with permission.<sup>281</sup> Copyright 2012, Royal Society of Chemistry. (b) Reproduced with permission.<sup>282</sup> Copyright 2011, American Chemical Society. (c) Reproduced with permission.<sup>271</sup> Copyright 2013, Wiley. (d) Reproduced with permission.<sup>283</sup> Copyright 2017, Wiley.

active materials in external reservoirs, circulating them to electrodes, such as carbon-based electrodes, for energy storage and discharge instead of confining them within solid electrolytes. The overall power density of a redox flow battery system is determined by the size of its electrode stack, while the total energy density depends on the amount of electrolyte stored in the reservoirs.<sup>285,286</sup> To enhance the energy density of redox flow batteries (RFBs), two primary strategies have been employed: increasing the solubility of the molecular reactants and maximizing the redox potential difference between the catholyte and anolyte.

A redox flow battery system comprises three key components: (i) separate storage reservoirs for two different electrolytes (anolyte (which flows through the anode), and catholyte (which flows through the cathode)); (ii) stacks of electrodes (anodes and cathodes) separated by a porous, ion-selective membrane; and (iii) a flow system assembly (Fig. 15a). During operation,

electrolytes containing dissolved electroactive species are pumped into their respective electrode compartments, where they undergo oxidation or reduction by releasing or accepting electrons at the electrode surface. When the RFB system discharges, the anolyte is oxidized at the anode, releasing electrons that travel through the current collectors to the external load and accumulate at the cathode, where the catholyte accepts the electrons and is reduced.<sup>65</sup> Additionally, charged species or supporting ions pass through the ion exchange membrane to complete the reaction and maintain electroneutrality.<sup>62</sup> During charging, the reactions reverse: the catholyte is oxidized and releases electrons, while the anolyte accepts electrons and is reduced. In conventional redox flow batteries, no phase changes occur in the reactants, whereas hybrid redox flow battery systems involve phase changes in at least one of the electrode reactions.<sup>287</sup>

Inorganic redox couples, such as Fe<sup>3+/2+</sup>/Cr<sup>2+/3+</sup>, Cr<sup>2+/3+</sup>, Ti<sup>3+/4+</sup>, Zn-Br, Sn<sup>2+/4+</sup>, and V<sup>2+/3+/4+/5+</sup>, have been used for

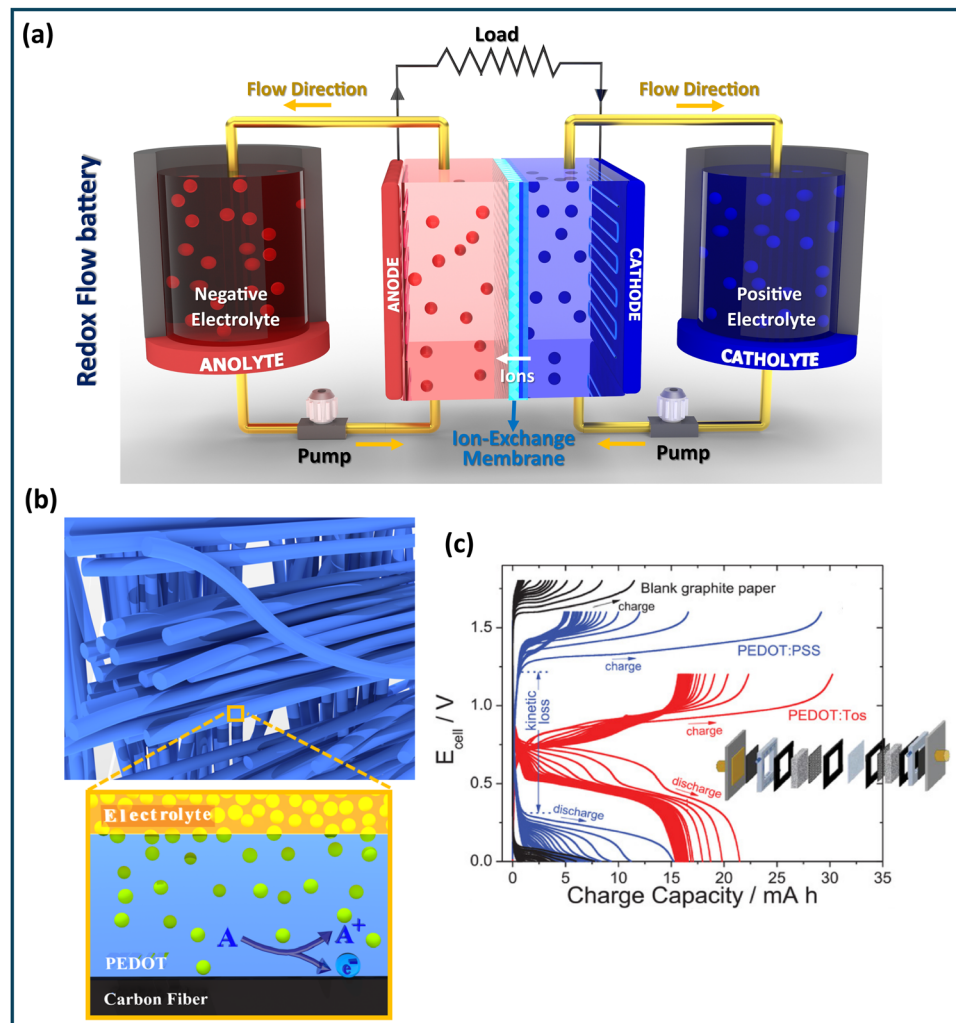


Fig. 15 Utilization of conducting polymers in redox flow batteries. (a) Schematic illustration of redox flow battery system. (b) Schematic illustration exhibits the highly conformal coating of oCVD PEDOT around each carbon fiber, highlighting the penetration of ionic species from the electrolyte through the PEDOT layer and ultimately reaching to the underlying carbon fiber substrate.<sup>10</sup> (c) Comparison of cell voltage versus capacity for blank graphite paper, PEDOT:Tos, and PEDOT:PSS.<sup>16</sup> (b) Reproduced with permission.<sup>10</sup> Copyright 2024, Wiley. (c) Reproduced with permission.<sup>16</sup> Copyright 2020, Wiley.

energy storage in RFBs. Despite the availability of various redox molecules, rechargeable has shifted toward organic molecules due to the abundance of their constituent elements and the flexibility of organic synthesis for chemical design.<sup>288</sup> Additionally, organic redox couples like viologen, anthraquinone, and 2,2,6,6-tetramethylpiperidine 1-oxyl (commonly known as TEMPO), have been investigated in RFBs. However, only a few have been commercialized, including all-vanadium redox flow batteries (VRFB),<sup>63</sup> Fe-based and Mn-based RFBs,<sup>289,290</sup> and Zn-based hybrid flow battery systems.<sup>287,291</sup> As an alternative to the high-cost all-vanadium RFB, RFBs using the same electroactive species with various oxidation states, such as all-chromium, all-iron, all-lead, and all-copper have been proposed.<sup>292</sup> Notably, all-iron RFBs have gained significant attention due to their low toxicity, material abundance, and low cost.<sup>292,293</sup> The substantial initial expense limits the widespread adoption of RFBs, with recent reports indicating that the cell stack alone accounts for about 31% of the

total cost.<sup>293</sup> Given that electrodes and membranes form the cell stack, optimizing these components is essential for cost reduction. Electrodes in RFBs must ensure strong electrochemical interactions with various redox couples and maintain stability in both acidic and alkaline environments.

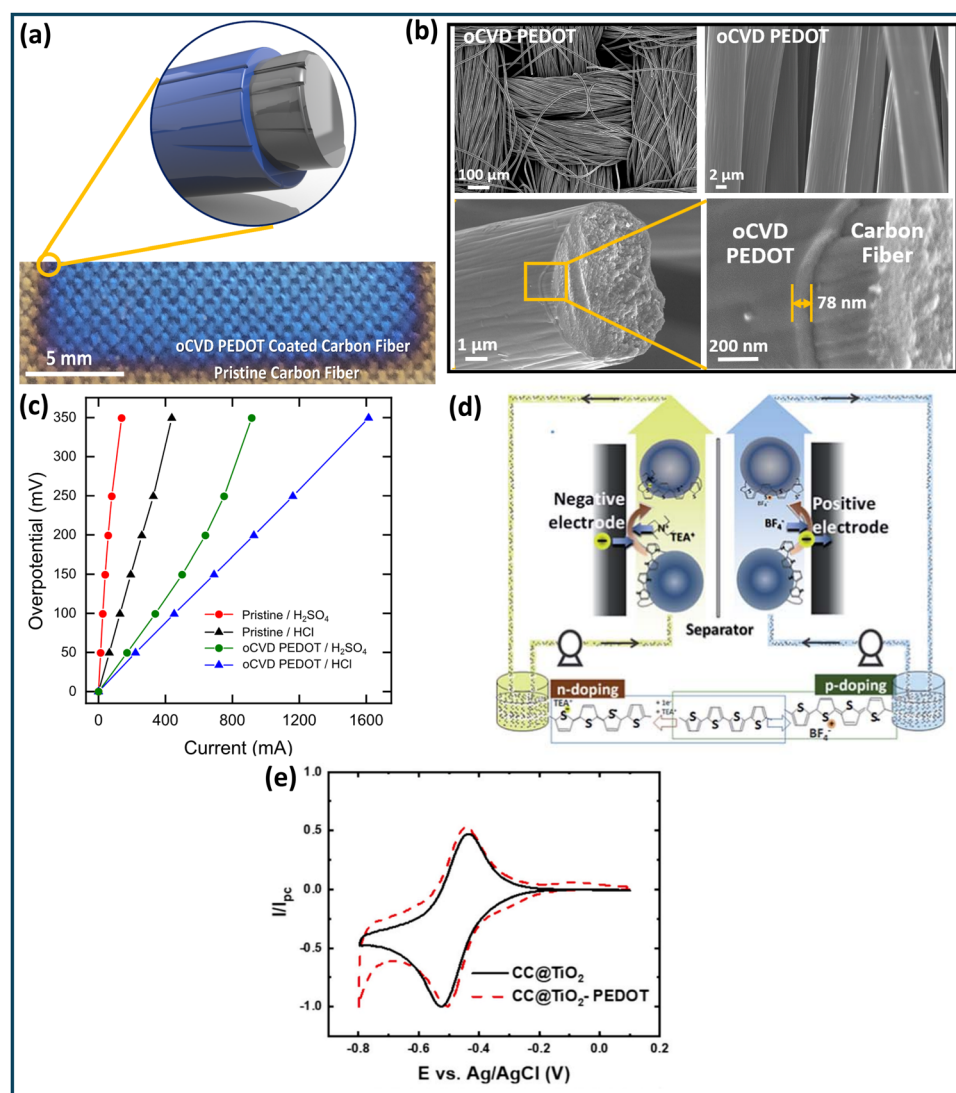
Common electrode materials used in RFBs include low-cost porous carbon options such as carbon fiber paper (CFP), carbon cloth (CC), graphite felt (GF), and their composites with other materials.<sup>9,64,65</sup> Carbon cloth holds significant promise for RFB electrodes, but it suffers from low specific capacitance, limited specific surface area, and poor wettability due to its hydrophobic nature.<sup>9,10,66</sup> Similarly, graphite felt faces challenges such as low electrochemical activity, limited specific surface area, poor wettability, and suboptimal chemical kinetics, leading to low mass transfer.<sup>67,294,295</sup> To address these issues and enhance the performance of carbon-based electrodes, various modification techniques have been explored,



including wet chemical oxidation,<sup>68,69</sup> electrochemical oxidation,<sup>70,71</sup> thermal treatment,<sup>72,73</sup> plasma enhancement,<sup>74,75</sup> and utilization of conducting polymer coatings.<sup>9,10</sup>

Depositing thin layers of conducting polymers and their composites on carbon electrodes has emerged as a promising technique to enhance the electrochemical performance of carbon-based electrodes.<sup>9,10,16,76,296,297</sup> Organic conjugated conducting polymers are gaining significant attention for RFBs due to their cost-effectiveness, abundance, non-toxicity, minimal environmental impact, and ease of chemical design.<sup>9,10,298,299</sup> PEDOT, in particular, can be chemically engineered to control the diffusion of electroactive species through its volume, making it suitable as an ion-selective material.<sup>16</sup> As the charge density or

doping level of PEDOT thin films increases, electroneutrality is maintained by either expelling cations or inserting anions, depending on the mobility of the counterion dopants.<sup>19</sup> For large, immobile counterions such as PSS, cations are expelled from the conjugated polymer chain, and the access of anions from the electrolyte is impeded due to the Donnan exclusion effect.<sup>197</sup> Conversely, if the dopant is small and exchangeable (such as chloride or tosylate), it facilitates anion insertion from the electrolyte into the conducting polymer matrix to balance the charge transfer.<sup>9,10,19</sup> The Donnan exclusion effect (also known as Gibbs–Donnan effect), describes the distribution of charged ions near a permeable membrane and has been applied to explain the low permeability of vanadium in anion exchange membranes (AEMs) for redox flow batteries.<sup>300</sup>



**Fig. 16** Utilization of conducting polymers in redox flow battery electrodes. (a) Photograph of oCVD PEDOT-coated carbon cloth electrodes integrated into the redox flow battery.<sup>9</sup> (b) SEM images showcasing the highly conformal coating of oCVD PEDOT on carbon cloth electrodes.<sup>9</sup> (c) Cell polarization data at 50% state-of-charge (0.25 M  $\text{Fe}^{2+}$ /0.25 M  $\text{Fe}^{3+}$ ), comparing the performance of pristine and oCVD PEDOT-coated carbon cloth electrodes in different electrolytes with a flow rate of 15  $\text{mL min}^{-1}$ .<sup>9</sup> (d) Schematic illustration of a redox flow battery utilizing polythiophene microparticles as both the cathodic and anodic redox-active species.<sup>17</sup> (e) CV graphs of carbon cloth electrodes coated with  $\text{TiO}_2$  and  $\text{TiO}_2$ -PEDOT.<sup>302</sup> (a)–(c) Reproduced with permission.<sup>9</sup> Copyright 2020, Wiley. (d) Reproduced with permission.<sup>17</sup> Copyright 2014, Royal Society of Chemical. (e) Reproduced with permission.<sup>302</sup> Copyright 2024, Elsevier.

PEDOT conducting polymers exhibit notable inherent molecular-level porosity, setting them apart from conventional heterogeneous electrocatalysts.<sup>8,25,301</sup> During the electrochemical reaction, the de-doping of PEDOT thin films involves expelling counterion dopants and incorporating anions from the electrolyte into the polymer matrix.<sup>10,19</sup> The fabrication of oCVD PEDOT thin films with small, easily exchangeable primary counterions, such as chloride, enhances electrochemical reactions by facilitating the transport of ionic species from the electrolyte through the PEDOT pseudocapacitive layer (Fig. 15b).<sup>10</sup> Vagin *et al.*<sup>16</sup> demonstrated congruent ionic charge transport from the electrolyte (4,5-dibenzoquinone-1,3-benzenedisulfonate (tiron)) along with mobile Tos counterion dopants in PEDOT:Tos, whereas such behavior was suppressed in PEDOT:PSS due to the immobile polyanion PSS counterion dopant. This congruent ionic transport in PEDOT:Tos significantly improved the charge and discharge capacity of the flow cell compared to PEDOT:PSS (Fig. 15c).<sup>16</sup> Generally, incorporating smaller counterions within the PEDOT polymer chain enhances doping/de-doping capability, promotes ion exchange with the electrolyte, and results in superior overall electrochemical performance.<sup>2,10</sup>

The application of a highly conformal oCVD PEDOT layer over porous carbon cloth electrodes in redox flow batteries reduces ohmic, kinetic, and mass transport resistances compared to pristine carbon cloth electrodes, leading to increased current density (Fig. 16a–c).<sup>9</sup> Compared to unmodified carbon cloth electrodes, oCVD PEDOT-coated carbon cloth electrodes significantly increase the maximum current density at an applied overpotential of 350 mV, achieving 6.7-fold and 3.7-fold enhancements in iron sulfate and iron chloride, respectively (Fig. 16c).<sup>9</sup> Oh *et al.*<sup>17</sup> developed a symmetric, metal-free, all-organic redox flow battery using polythiophene microparticles as the redox couple in a non-aqueous electrolyte. This innovative design achieved a high cell potential of 2.5 V, stable cycling performance, and an energy efficiency of 60.9% (Fig. 16d). The performance of carbon cloth electrodes was enhanced by coating them with a TiO<sub>2</sub> layer followed by the application of a PEDOT thin film (CC@TiO<sub>2</sub>-PEDOT) and exhibited high reversibility for the V(II)/V(III) redox reaction (Fig. 16e).<sup>302</sup>

## 5. Conclusion and outlook

The synergy between scientific understanding and technological objectives has led to significant advancements in enhancing the electronic, optical, electrochemical, and thermal properties of conjugated conducting polymers. Moreover, their chemical reactivity, biocompatibility, magnetic properties, and mechanical flexibility make them highly attractive for a wide range of applications. In order to have a real-world impact, conjugated conducting polymers must be stable, cost-effective, scalable to large areas, and present minimal environmental, health, and safety (EHS) risks. To meet these demands, advanced manufacturing techniques are required that can economically fabricate thin films of conjugated conducting polymers over large areas while achieving the desired nanostructure within

microscale layers. A crucial aspect of harnessing the full capabilities of semi-crystalline conjugated conducting polymers lies in controlling and leveraging their nanoscale order and orientation.

Engineering the texture and nanostructure of conjugated conducting polymers is crucial for advancing their utilization in applications like electrochemical energy storage devices, flexible optoelectronics, and wearable devices. Key factors such as semi-crystalline orientation,  $\pi$ – $\pi$  stacking, crystallite size, and inter-chain and inter-crystallite couplings must be precisely controlled during synthesis to optimize charge carrier mobility. Improving the pseudocapacitive properties of conducting polymers can be achieved by enhancing their semi-crystalline structure and extending polymer chain length. Ion diffusivity varies between crystalline and amorphous regions, and edge-on oriented conducting polymer thin films showing greater charge storage capacity during cycling. The size and type of counterion dopants are crucial for the electrochemical performance of conjugated conducting polymers. Small counterion dopants like chloride enhance redox exchange and expand the electrochemical potential window, benefiting electrochemical energy storage devices. Directly fabricating ultra-thin, highly conformal coatings of conducting polymers *via* vapor phase deposition onto flexible, porous substrates has proven promising for creating flexible supercapacitors with high specific capacitance, as it preserves the substrate's high surface area, providing more active sites for redox reactions.

N-Type conjugated conducting polymers are crucial for many devices, including organic energy storage, electrochemical transistors, and spintronic devices. However, their development has lagged behind p-type counterparts. For air-stable n-type polymers, the LUMO level must be below  $-4.0$  eV, which also improves n-type doping efficiency. Porous materials, with their reduced resistance to mass transport, are ideal for applications requiring molecular diffusion, such as adsorption, separation, sensing, and catalysis. In electrochemical energy storage, hierarchical pore structures improve access to redox-active sites and help accommodate mechanical stresses from doping and de-doping. Porous conjugated conducting polymers often have a high surface area ( $>1000$  m<sup>2</sup> g<sup>-1</sup>) without relying on heavy metals, using only earth-abundant elements like hydrogen, carbon, oxygen, nitrogen, and sulfur, making them suitable for large-area electrochemical energy storage devices.

Conducting polymers have become promising materials in Li-ion batteries, addressing limitations of traditional electrodes with their mixed ionic and electrical conductivity, flexibility, rapid doping/de-doping, stability, and reversible redox capabilities. The tunable nanostructure of conjugated conducting polymers allows for the design of materials with enhanced energy density, cycle stability, and rate performance. Additionally, incorporating conducting polymers improves electrode–electrolyte interfaces, mitigating issues like SEI formation and dendrite growth. As research progresses, optimizing conjugated polymer nanostructures, doping levels, and hybridization with other materials will be key to advancing Li-ion battery technology, leading to more efficient, durable, and sustainable energy storage solutions.



Research and development of conjugated conducting polymer-based flexible supercapacitors have progressed rapidly, with substantial efforts aimed at positioning these supercapacitors as a new platform for flexible energy storage devices. Interestingly, the direct growth of ultrathin, highly conformal coatings of conjugated conducting polymers *via* vapor phase deposition onto flexible, porous substrates has emerged as a promising approach for achieving flexible supercapacitors with high specific capacitance. This method offers a significant advantage by preserving the high surface area of the porous, flexible substrate, thereby facilitating more active sites for redox reactions. To fully realize the potential of conjugated conducting polymers in flexible supercapacitors, further advancements are crucial, particularly in optimizing specific power and energy density by exploiting the broader potential window provided by eco-friendly ionic liquid electrolyte systems. Achieving large-scale commercial applications will require not only innovative strategies but also a deeper and more refined understanding of the effects of *in situ* doping and the nanostructure of conjugated polymer architectures.

Conducting polymers hold significant promise for enhancing the performance of redox flow batteries. Traditional carbon-based electrode materials, such as carbon cloth and graphite felt, often face limitations like low specific capacitance, limited surface area, and poor wettability, which hinder their efficiency. However, the application of ultrathin, highly conformal coatings of conducting polymers on carbon-based electrodes has emerged as an effective strategy to overcome these challenges. Conjugated conducting polymers can be chemically engineered to improve the diffusion of electroactive species, enhance ion selectivity, and ultimately boost the overall electrochemical performance of redox flow battery electrodes, paving the way for more efficient and durable energy storage solutions.

## Data availability

No software or code have been included as part of this review article.

## Conflicts of interest

Authors declare no competing interests.

## Acknowledgements

M. H. G. acknowledges the Faculty Startup Support from the San Diego State University (SDSU).

## References

- 1 N. Wunderling, M. Willeit, J. F. Donges and R. Winkelmann, *Nat. Commun.*, 2020, **11**, 5177.
- 2 M. Heydari Gharahcheshmeh and K. K. Gleason, *Adv. Mater. Interfaces*, 2019, **6**, 1801564.
- 3 K. K. Gleason and M. H. Gharahcheshmeh, *Conjugated Polymers at Nanoscale: Engineering Orientation, Nanostructure, and Properties*, Walter de Gruyter GmbH & Co KG, 2021.
- 4 A. Varyani, M. Salehi and M. Heydari Gharahcheshmeh, *Energies*, 2024, **17**, 2865.
- 5 S. Ould Amrouche, D. Rekioua, T. Rekioua and S. Bacha, *Int. J. Hydrogen Energy*, 2016, **41**, 20914–20927.
- 6 I. Tantis, S. Talande, V. Tzitzios, G. Basina, V. Shrivastav, A. Bakandritsos and R. Zboril, *Adv. Funct. Mater.*, 2023, **33**, 2209360.
- 7 A. M. Bryan, L. M. Santino, Y. Lu, S. Acharya and J. M. D'Arcy, *Chem. Mater.*, 2016, **28**, 5989–5998.
- 8 M. Heydari Gharahcheshmeh and K. Gleason, *Mater. Today Adv.*, 2020, **8**, 100086.
- 9 M. Heydari Gharahcheshmeh, C. T. C. Wan, Y. Ashraf Gandomi, K. V. Greco, A. Forner-Cuenca, Y. M. Chiang, F. R. Brushett and K. K. Gleason, *Adv. Mater. Interfaces*, 2020, **7**, 2000855.
- 10 M. Heydari Gharahcheshmeh and K. Chowdhury, *Adv. Mater. Interfaces*, 2024, **11**, 2400118.
- 11 J. G. Ibanez, M. E. Rincón, S. Gutierrez-Granados, M. H. Chahma, O. A. Jaramillo-Quintero and B. A. Frontana-Uribe, *Chem. Rev.*, 2018, **118**, 4731–4816.
- 12 Y. Yu, C. Yang, Y. Jiang, J. Zhu, Y. Zhao, S. Liang, K. Wang, Y. Zhou, Y. Liu and J. Zhang, *Small*, 2023, **19**, 2303779.
- 13 B. Liu, K. Zhu, K. Ye, J. Yan, G. Wang and D. Cao, *J. Power Sources*, 2022, **552**, 232226.
- 14 L. M. Zhu, A. W. Lei, Y. L. Cao, X. P. Ai and H. X. Yang, *Chem. Commun.*, 2013, **49**, 567–569.
- 15 G.-L. Xu, Q. Liu, K. K. S. Lau, Y. Liu, X. Liu, H. Gao, X. Zhou, M. Zhuang, Y. Ren and J. Li, *Nat. Energy*, 2019, **4**, 484–494.
- 16 M. Vagin, C. Che, V. Gueskine, M. Berggren and X. Crispin, *Adv. Funct. Mater.*, 2020, **30**, 2007009.
- 17 S. H. Oh, C. W. Lee, D. H. Chun, J. D. Jeon, J. Shim, K. H. Shin and J. H. Yang, *J. Mater. Chem. A*, 2014, **2**, 19994–19998.
- 18 S.-M. Kim, C.-H. Kim, Y. Kim, N. Kim, W.-J. Lee, E.-H. Lee, D. Kim, S. Park, K. Lee, J. Rivnay and M.-H. Yoon, *Nat. Commun.*, 2018, **9**, 3858.
- 19 J. Heinze, B. A. Frontana-Uribe and S. Ludwigs, *Chem. Rev.*, 2010, **110**, 4724–4771.
- 20 G. A. Snook, P. Kao and A. S. Best, *J. Power Sources*, 2011, **196**, 1–12.
- 21 T. B. Schon, B. T. McAllister, P.-F. Li and D. S. Seferos, *Chem. Soc. Rev.*, 2016, **45**, 6345–6404.
- 22 C. Bauer, M. Kirchner and A. Krueger, *Energy Adv.*, 2024, **3**, 1422–1430.
- 23 Y. Ma, K. Xu, X. Liu, S. Yao, X. Li, Y. Si and X. Li, *Chem. Eng. J.*, 2024, 154348.
- 24 M. G. Tadesse, N. Simon and J. F. Lübben, *Adv. Energy Sustainability Res.*, 2024, 2400006.
- 25 K. K. Gleason, F. Brushett, C. Wan, A. Forner-Cuenca, M. H. Gharahcheshmeh and Y. A. Gandomi, *U.S. Pat.*, 11677078, 2023.



26 I. E. Jacobs, G. d'Avino, V. Lemaur, Y. Lin, Y. Huang, C. Chen, T. F. Harrelson, W. Wood, L. J. Spalek and T. Mustafa, *J. Am. Chem. Soc.*, 2022, **144**, 3005–3019.

27 F. Schauer, *J. Appl. Phys.*, 2020, **128**, 150902.

28 S. P. Arnold, J. K. Harris, B. Neelamraju, M. Rudolph and E. L. Ratcliff, *Synth. Met.*, 2019, **253**, 26–33.

29 M. Heydari Gharahcheshmeh and K. K. Gleason, *Energies*, 2022, **15**, 3661.

30 M. Heydari Gharahcheshmeh, *MRS Commun.*, 2024, DOI: [10.1557/s43579-024-00593-w](https://doi.org/10.1557/s43579-024-00593-w).

31 P. Moni, J. Lau, A. C. Mohr, T. C. Lin, S. H. Tolbert, B. Dunn and K. K. Gleason, *ACS Appl. Energy Mater.*, 2018, **1**, 7093–7105.

32 Q. Liu, Y.-T. Liu, C. Zhao, Q.-S. Weng, J. Deng, I. Hwang, Y. Jiang, C. Sun, T. Li and W. Xu, *ACS Nano*, 2022, **16**, 14527–14538.

33 M. E. Abdelhamid, A. P. O'Mullane and G. A. Snook, *RSC Adv.*, 2015, **5**, 11611–11626.

34 J. Rehmen, T. Pathirana, L. Garcia-Quintana, R. Kerr, P. C. Howlett, K. Zuber, C. Pozo-Gonzalo and D. R. Evans, *ACS Appl. Nano Mater.*, 2020, **3**, 3820–3828.

35 S. J. An, J. Li, C. Daniel, D. Mohanty, S. Nagpure and D. L. Wood, *Carbon*, 2016, **105**, 52–76.

36 Z. Wen, Y. Peng, J. Cong, H. Hua, Y. Lin, J. Xiong, J. Zeng and J. Zhao, *Nano Res.*, 2019, **12**, 2535–2542.

37 S. Li, J. Huang, J. Wang and K. Han, *Mater. Lett.*, 2021, **293**, 129712.

38 R. Na, K. Minnici, G. Zhang, N. Lu, M. A. González, G. Wang and E. Reichmanis, *ACS Appl. Mater. Interfaces*, 2019, **11**, 40034–40042.

39 T. Li, X.-Z. Yuan, L. Zhang, D. Song, K. Shi and C. Bock, *Electrochem. Energy Rev.*, 2020, **3**, 43–80.

40 B. Zhang, J. Shen, Q. Wang, C. Hu, B. Luo, Y. Liu, Z. Xiao and X. Ou, *Energy Environ. Mater.*, 2023, **6**, e12270.

41 Y. Li, X. Liu, D. Ren, H. Hsu, G.-L. Xu, J. Hou, L. Wang, X. Feng, L. Lu and W. Xu, *Nano Energy*, 2020, **71**, 104643.

42 Q. Xie, Z. Hu, C. Zhao, S. Zhang and K. Liu, *RSC Adv.*, 2015, **5**, 50859–50864.

43 Z. Du, W. Peng, Z. Wang, H. Guo, Q. Hu and X. Li, *Ionics*, 2018, **24**, 3717–3724.

44 Y.-H. Du, H. Sheng, X.-H. Meng, X.-D. Zhang, Y.-G. Zou, J.-Y. Liang, M. Fan, F. Wang, J. Tang and F.-F. Cao, *Nano Energy*, 2022, **94**, 106901.

45 X. Ding, Y.-X. Li, S. Wang, J.-M. Dong, A. Yasmin, Q. Hu, Z.-Y. Wen and C.-H. Chen, *Nano Energy*, 2019, **61**, 411–419.

46 L. Li, M. Xu, Q. Yao, Z. Chen, L. Song, Z. Zhang, C. Gao, P. Wang, Z. Yu and Y. Lai, *ACS Appl. Mater. Interfaces*, 2016, **8**, 30879–30889.

47 T. He, Y. Lu, Y. Su, L. Bao, J. Tan, L. Chen, Q. Zhang, W. Li, S. Chen and F. Wu, *ChemSusChem*, 2018, **11**, 1639–1648.

48 Q. Gan, N. Qin, Y. Zhu, Z. Huang, F. Zhang, S. Gu, J. Xie, K. Zhang, L. Lu and Z. Lu, *ACS Appl. Mater. Interfaces*, 2019, **11**, 12594–12604.

49 G.-L. Xu, Q. Liu, K. K. Lau, Y. Liu, X. Liu, H. Gao, X. Zhou, M. Zhuang, Y. Ren and J. Li, *Nat. Energy*, 2019, **4**, 484–494.

50 P. G. Bruce, S. A. Freunberger, L. J. Hardwick and J.-M. Tarascon, *Nat. Mater.*, 2012, **11**, 19–29.

51 H. Xiang, N. Deng, H. Zhao, X. Wang, L. Wei, M. Wang, B. Cheng and W. Kang, *J. Energy Chem.*, 2021, **58**, 523–556.

52 R. Fang, S. Zhao, Z. Sun, D.-W. Wang, H.-M. Cheng and F. Li, *Adv. Mater.*, 2017, **29**, 1606823.

53 X. Ji, K. T. Lee and L. F. Nazar, *Nat. Mater.*, 2009, **8**, 500–506.

54 G. Zhou, D.-W. Wang, F. Li, P.-X. Hou, L. Yin, C. Liu, G. Q. M. Lu, I. R. Gentle and H.-M. Cheng, *Energy Environ. Sci.*, 2012, **5**, 8901–8906.

55 Y. Zhang, H. W. Song, K. R. Crompton, X. Yang, K. Zhao and S. Lee, *Nano Energy*, 2023, **115**, 108756.

56 Y. Xie, H. Zhao, H. Cheng, C. Hu, W. Fang, J. Fang, J. Xu and Z. Chen, *Appl. Energy*, 2016, **175**, 522–528.

57 M. D. Hager, P. Greil, C. Leyens, S. Van Der Zwaag and U. S. Schubert, *Adv. Mater.*, 2010, **22**, 5424–5430.

58 V. Augustyn, P. Simon and B. Dunn, *Energy Environ. Sci.*, 2014, **7**, 1597–1614.

59 C. Peng, D. Hu and G. Z. Chen, *Chem. Commun.*, 2011, **47**, 4105–4107.

60 J. Zang, S.-J. Bao, C. M. Li, H. Bian, X. Cui, Q. Bao, C. Q. Sun, J. Guo and K. Lian, *J. Phys. Chem. C*, 2008, **112**, 14843–14847.

61 H. Randriamahazaka, C. Plesse, D. Teyssie and C. Chevrot, *Electrochim. Acta*, 2005, **50**, 1515–1522.

62 E. Sánchez-Díez, E. Ventosa, M. Guarnieri, A. Trovò, C. Flox, R. Marcilla, F. Soavi, P. Mazur, E. Aranzabe and R. Ferret, *J. Power Sources*, 2021, **481**, 228804.

63 M. Gencten and Y. Sahin, *Int. J. Energy Res.*, 2020, **44**, 7903–7923.

64 A. Forner-Cuena, E. E. Penn, A. M. Oliveira and F. R. Brushett, *J. Electrochem. Soc.*, 2019, **166**, A2230–A2241.

65 J. Li, M. Al-Yasiri, H. Pham and J. Park, in *Advanced Materials for Battery Separators*, ed. S. Thomas, D. Rouxel, N. Kalarikkal, B. Kottathodi and H. J. Maria, Elsevier, 2024, pp. 327–347, DOI: [10.1016/B978-0-12-817507-1.00003-X](https://doi.org/10.1016/B978-0-12-817507-1.00003-X).

66 M. Maleki, G. A. El-Nagar, D. Bernsmeier, J. Schneider and C. Roth, *Sci. Rep.*, 2020, **10**, 11153.

67 I. Azpitarte, U. Eletxigerra, A. Barros, E. Aranzabe and R. Cid, *Batteries*, 2023, **9**, 39.

68 L. Yue, W. Li, F. Sun, L. Zhao and L. Xing, *Carbon*, 2010, **48**, 3079–3090.

69 G. Wang, H. Wang, X. Lu, Y. Ling, M. Yu, T. Zhai, Y. Tong and Y. Li, *Adv. Mater.*, 2014, **26**, 2676–2682.

70 X.-G. Li, K.-L. Huang, S.-Q. Liu, N. Tan and L.-Q. Chen, *Trans. Nonferrous Met. Soc. China*, 2007, **17**, 195–199.

71 R. Wang and Y. Li, *Energy Storage Mater.*, 2020, **31**, 230–251.

72 P. C. Ghimire, R. Schweiss, G. G. Scherer, T. M. Lim, N. Wai, A. Bhattacharai and Q. Yan, *Carbon*, 2019, **155**, 176–185.

73 Y.-J. Gu, W. Wen and J.-M. Wu, *J. Mater. Chem. A*, 2018, **6**, 21078–21086.

74 B. Ouyang, Y. Zhang, Y. Wang, Z. Zhang, H. J. Fan and R. S. Rawat, *J. Mater. Chem. A*, 2016, **4**, 17801–17808.



75 D. Dixon, D. J. Babu, J. Langner, M. Bruns, L. Pfaffmann, A. Bhaskar, J. J. Schneider, F. Scheiba and H. Ehrenberg, *J. Power Sources*, 2016, **332**, 240–248.

76 M. G. Sumdani, M. R. Islam, A. N. A. Yahaya and S. I. Safie, *Polym. Eng. Sci.*, 2022, **62**, 269–303.

77 K. Choudhary, A. X. Chen, G. M. Pitch, R. Runser, A. Urbina, T. J. Dunn, M. Kodur, A. T. Kleinschmidt, B. G. Wang, J. A. Bunch, D. P. Fenning, A. L. Ayzner and D. J. Lipomi, *ACS Appl. Mater. Interfaces*, 2021, **13**, 51436–51446.

78 H. A. M. Mustafa and D. A. Jameel, *J. Appl. Sci. Technol. Trends*, 2021, **2**, 91–95.

79 K. R. Thines, E. C. Abdullah, N. M. Mubarak and M. Ruthiraan, *Biomass Bioenergy*, 2017, **98**, 95–111.

80 Y. Yan, J. Ju, S. Dong, Y. Wang, L. Huang, L. Cui, F. Jiang, Q. Wang, Y. Zhang and G. Cui, *Adv. Sci.*, 2021, **8**, 2003887.

81 Z. Yang, L. Ding, P. Wu, Y. Liu, F. Nie and F. Huang, *Chem. Eng. J.*, 2015, **268**, 60–66.

82 S. Cosnier and A. Karyakin, *Electropolymerization: concepts, materials and applications*, John Wiley & Sons, 2011.

83 G. Sabouraud, S. Sadki and N. Brodie, *Chem. Soc. Rev.*, 2000, **29**, 283–293.

84 C. Zhao, Z. Chen, W. Wang, P. Xiong, B. Li, M. Li, J. Yang and Y. Xu, *Angew. Chem., Int. Ed.*, 2020, **59**, 11992–11998.

85 C. Friebe, M. D. Hager, A. Winter and U. S. Schubert, *Adv. Mater.*, 2012, **24**, 332–345.

86 M. Fabretto, J.-P. Autere, D. Hoglinger, S. Field and P. Murphy, *Thin Solid Films*, 2011, **519**, 2544–2549.

87 M. Fabretto, K. Zuber, C. Hall and P. Murphy, *Macromol. Rapid Commun.*, 2008, **29**, 1403–1409.

88 A. T. Lawal and G. G. Wallace, *Talanta*, 2014, **119**, 133–143.

89 A. Mohammadi, M. A. Hasan, B. Liedberg, I. Lundström and W. R. Salaneck, *Synth. Met.*, 1986, **14**, 189–197.

90 D. Bhattacharyya, R. M. Howden, D. C. Borrelli and K. K. Gleason, *J. Polym. Sci., Part B: Polym. Phys.*, 2012, **50**, 1329–1351.

91 J. Kim, E. Kim, Y. Won, H. Lee and K. Suh, *Synth. Met.*, 2003, **139**, 485–489.

92 P. A. Levermore, L. Chen, X. Wang, R. Das and D. D. C. Bradley, *Adv. Mater.*, 2007, **19**, 2379–2385.

93 B. Winther-Jensen and K. West, *Macromolecules*, 2004, **37**, 4538–4543.

94 T. Bashir, M. Ali, S.-W. Cho, N.-K. Persson and M. Skrifvars, *Polym. Adv. Technol.*, 2013, **24**, 210–219.

95 T. Bashir, M. Skrifvars and N.-K. Persson, *Polym. Adv. Technol.*, 2012, **23**, 611–617.

96 K. K. Gleason, *CVD Polym.*, 2015, 1–11, DOI: [10.1002/9783527690275.ch1](https://doi.org/10.1002/9783527690275.ch1).

97 S. G. Im, D. Kusters, W. Choi, S. H. Baxamusa, M. C. M. de Sanden and K. K. Gleason, *ACS Nano*, 2008, **2**, 1959–1967.

98 S. Lee, D. C. Paine and K. K. Gleason, *Adv. Funct. Mater.*, 2014, **24**, 7187–7196.

99 F. Muralter, A. M. Coclite and K. K. S. Lau, *Adv. Electron. Mater.*, 2021, **7**, 2000871.

100 M. Rajesh, C. J. Raj, R. Manikandan, B. C. Kim, S. Y. Park and K. H. Yu, *Mater. Today Energy*, 2017, **6**, 96–104.

101 A. G. MacDiarmid, W. E. Jones Jr, I. D. Norris, J. Gao, A. T. Johnson Jr, N. J. Pinto, J. Hone, B. Han, F. K. Ko and H. Okuzaki, *Synth. Met.*, 2001, **119**, 27–30.

102 Q. Tang, J. Wu, X. Sun, Q. Li and J. Lin, *Langmuir*, 2009, **25**, 5253–5257.

103 Y. Song, J.-B. Fan and S. Wang, *Mater. Chem. Front.*, 2017, **1**, 1028–1040.

104 C.-S. Park, D. H. Kim, B. J. Shin, D. Y. Kim, H.-K. Lee and H.-S. Tae, *Materials*, 2016, **9**, 812.

105 M. Heydari Gharahcheshmeh, M. M. Tavakoli, E. F. Gleason, M. T. Robinson, J. Kong and K. K. Gleason, *Sci. Adv.*, 2019, **5**, eaay0414.

106 R. Das and A. Chanda, *Nano-size polymers: Preparation, properties, applications*, 2016, pp. 283–306.

107 K. Norrman, A. Ghanbari-Siahkali and N. B. Larsen, *Annu. Rep. Prog. Chem., Sect. C: Phys. Chem.*, 2005, **101**, 174–201.

108 H. Yan and H. Okuzaki, *Synth. Met.*, 2009, **159**, 2225–2228.

109 G.-F. Wang, X.-M. Tao, J. H. Xin and B. Fei, *Nanoscale Res. Lett.*, 2009, **4**, 613–617.

110 C. Gong, H. B. Yang, Q. L. Song, Z. S. Lu and C. M. Li, *Sol. Energy Mater. Sol. Cells*, 2012, **100**, 115–119.

111 M. D. Tyona, *Adv. Mater. Res.*, 2013, **2**, 195.

112 A. Boudrioua, M. Chakaroun and A. Fischer, in *Organic Lasers*, ed. A. Boudrioua, M. Chakaroun and A. Fischer, Elsevier, 2017, pp. 49–93, DOI: [10.1016/B978-1-78548-158-1.50002-X](https://doi.org/10.1016/B978-1-78548-158-1.50002-X).

113 Y. Jiang, T. Liu and Y. Zhou, *Adv. Funct. Mater.*, 2020, **30**, 2006213.

114 V. Goel, R. Tanwar and U. Mandal, *J. Chem. Technol. Biotechnol.*, 2021, **96**, 502–513.

115 M. S. Cho, S. Y. Kim, J. D. Nam and Y. Lee, *Synth. Met.*, 2008, **158**, 865–869.

116 A. Seijas-Da Silva, J. Noguera-Gomez, P. J. Rodríguez-Canto, J. F. Sanchez-Royo, A. Cantarero, C. M. Gómez and R. Abargues, *J. Phys. Chem. C*, 2020, **124**, 22884–22892.

117 K. H. Hong, K. W. Oh and T. J. Kang, *J. Appl. Polym. Sci.*, 2005, **97**, 1326–1332.

118 X. Wang, S. Meng, W. Ma, J. Pionteck, M. Gnanaseelan, Z. Zhou, B. Sun, Z. Qin and M. Zhu, *React. Funct. Polym.*, 2017, **112**, 74–80.

119 M. Ferenets and A. Harlin, *Thin Solid Films*, 2007, **515**, 5324–5328.

120 L. A. A. Pettersson, F. Carlsson, O. Inganäs and H. Arwin, *Thin Solid Films*, 1998, **313–314**, 356–361.

121 I. S. Tsagkalias, T. K. Manios and D. S. Achilias, *Polymers*, 2017, **9**, 432.

122 J. Wang, Z. Shi, Y. Ge, Y. Wang, J. Fan and J. Yin, *Mater. Chem. Phys.*, 2012, **136**, 43–50.

123 M. K. Erdogan, M. Karakisla and M. Sacak, *J. Text. Inst.*, 2019, **110**, 282–288.

124 M. N. Gueye, A. Carella, J. Faure-Vincent, R. Demadrille and J.-P. Simonato, *Prog. Mater. Sci.*, 2020, **108**, 100616.

125 C. Friebe, M. D. Hager, A. Winter and U. S. Schubert, *Adv. Mater.*, 2012, **24**, 332–345.

126 Y. Seki, M. Takahashi and M. Takashiri, *RSC Adv.*, 2019, **9**, 15957–15965.



127 A. Yazdanpanah, A. Ramedani, A. Abrishamkar, P. B. Milan, Z. S. Moghadan, N. P. S. Chauhan, F. Sefat and M. Mozafari, in *Fundamentals and Emerging Applications of Polyaniline*, ed. M. Mozafari and N. P. S. Chauhan, Elsevier, 2019, pp. 105–119, DOI: [10.1016/B978-0-12-817915-4.00006-3](https://doi.org/10.1016/B978-0-12-817915-4.00006-3).

128 J. Yano, Y. Ota and A. Kitani, *Mater. Lett.*, 2004, **58**, 1934–1937.

129 E. Schab-Balcerzak, *Electropolymerization*, BoD–Books on Demand, 2011.

130 M. M. Gvozdenović, B. Jugović, J. S. Stevanović and B. Grgur, *Hem. Ind.*, 2014, **68**, 673–684.

131 S. Sadki, P. Schottland, N. Brodie and G. Sabouraud, *Chem. Soc. Rev.*, 2000, **29**, 283–293.

132 E. Stussi, R. Stella and D. De Rossi, *Sens. Actuators, B*, 1997, **43**, 180–185.

133 A. T. Lawal and G. G. Wallace, *Talanta*, 2014, **119**, 133–143.

134 B. Winther-Jensen, J. Chen, K. West and G. Wallace, *Macromolecules*, 2004, **37**, 5930–5935.

135 D. Evans, M. Fabretto, M. Mueller, K. Zuber, R. Short and P. Murphy, *J. Mater. Chem.*, 2012, **22**, 14889–14895.

136 T. Y. Kim, C. M. Park, J. E. Kim and K. S. Suh, *Synth. Met.*, 2005, **149**, 169–174.

137 N. Massonnet, A. Carella, A. De Geyer, J. Faure-Vincent and J.-P. Simonato, *Chem. Sci.*, 2015, **6**, 412–417.

138 M. S. Cho, S. Y. Kim, J. D. Nam and Y. Lee, *Synth. Met.*, 2008, **158**, 865–869.

139 P. Hojati-Talemi, C. Bächler, M. Fabretto, P. Murphy and D. Evans, *ACS Appl. Mater. Interfaces*, 2013, **5**, 11654–11660.

140 B. Li, D. Ni, J. Fan and J. Zhou, *Chem. Eng. J.*, 2022, **445**, 136818.

141 M. Fabretto, M. Müller, K. Zuber and P. Murphy, *Macromol. Rapid Commun.*, 2009, **30**, 1846–1851.

142 D. Wu, J. Zhang, W. Dong, H. Chen, X. Huang, B. Sun and L. Chen, *Synth. Met.*, 2013, **176**, 86–91.

143 B. Winther-Jensen, D. W. Breiby and K. West, *Synth. Met.*, 2005, **152**, 1–4.

144 L. Sun, G. Yuan, L. Gao, J. Yang, M. Chhowalla, M. H. Gharahcheshmeh, K. K. Gleason, Y. S. Choi, B. H. Hong and Z. Liu, *Nat. Rev. Methods Primers*, 2021, **1**, 5.

145 M. T. Robinson, J. Tung, M. Heydari Gharahcheshmeh and K. K. Gleason, *Adv. Funct. Mater.*, 2021, **31**, 2101310.

146 K. K. Gleason, *CVD polymers: fabrication of organic surfaces and devices*, John Wiley & Sons, 2015.

147 J. P. Lock, S. G. Im and K. K. Gleason, *Macromolecules*, 2006, **39**, 5326–5329.

148 M. Heydari Gharahcheshmeh and K. K. Gleason, *Adv. Mater. Interfaces*, 2019, **6**, 1970016.

149 M. K. Charyton, J. Crépellière, K. Kotwica, M. Gora, W. Le, G. Frache, P. Grysian, J. Guillot, K. Baba and N. D. Boscher, *J. Mater. Chem. C*, 2024, **12**, 11368–11377.

150 M. M. Tavakoli, M. H. Gharahcheshmeh, N. Moody, M. G. Bawendi, K. K. Gleason and J. Kong, *Adv. Mater. Interfaces*, 2020, **7**, 2000498.

151 S. Kaviani, M. Mohammadi Ghaleni, E. Tavakoli and S. Nejati, *ACS Appl. Polym. Mater.*, 2019, **1**, 552–560.

152 M. Heydari Gharahcheshmeh, M. T. Robinson, E. F. Gleason and K. K. Gleason, *Adv. Funct. Mater.*, 2021, **31**, 2008712.

153 X. Wang, X. Zhang, L. Sun, D. Lee, S. Lee, M. Wang, J. Zhao, Y. Shao-Horn, M. Dincă and T. Palacios, *Sci. Adv.*, 2018, **4**, eaat5780.

154 G. Drewelow, H. W. Song, Z.-T. Jiang and S. Lee, *Appl. Surf. Sci.*, 2020, **501**, 144105.

155 M. Heydari Gharahcheshmeh, M. T. Robinson, E. F. Gleason and K. K. Gleason, *Adv. Funct. Mater.*, 2021, **31**, 2170097.

156 W. E. Tenhaeff and K. K. Gleason, *Adv. Funct. Mater.*, 2008, **18**, 979–992.

157 K. Zuber, H. Shere, J. Rehmen, V. Wheaton, M. Fabretto, P. J. Murphy and D. R. Evans, *ACS Omega*, 2018, **3**, 12679–12687.

158 A. H. Malik, F. Habib, M. J. Qazi, M. A. Ganayee, Z. Ahmad and M. A. Yatoo, *J. Polym. Res.*, 2023, **30**, 115.

159 J. L. Bredas and G. B. Street, *Acc. Chem. Res.*, 1985, **18**, 309–315.

160 R. J. Hoofman, M. P. De Haas, L. D. Siebbeles and J. M. Warman, *Nature*, 1998, **392**, 54–56.

161 H. Sirringhaus, P. Brown, R. Friend, M. M. Nielsen, K. Bechgaard, B. Langeveld-Voss, A. Spiering, R. A. Janssen, E. Meijer and P. Herwig, *Nature*, 1999, **401**, 685–688.

162 K. Chowdhury, S. K. Behura, M. Rahimi and M. Heydari Gharahcheshmeh, *ACS Appl. Energy Mater.*, 2024, **7**, 1068–1079.

163 S. Kee, N. Kim, B. S. Kim, S. Park, Y. H. Jang, S. H. Lee, J. Kim, J. Kim, S. Kwon and K. Lee, *Adv. Mater.*, 2016, **28**, 8625–8631.

164 C. M. Palumbiny, F. Liu, T. P. Russell, A. Hexemer, C. Wang and P. Müller-Buschbaum, *Adv. Mater.*, 2015, **27**, 3391–3397.

165 N. Kim, B. H. Lee, D. Choi, G. Kim, H. Kim, J.-R. Kim, J. Lee, Y. H. Kahng and K. Lee, *Phys. Rev. Lett.*, 2012, **109**, 106405.

166 S. Y. Son, Y. Kim, J. Lee, G.-Y. Lee, W.-T. Park, Y.-Y. Noh, C. E. Park and T. Park, *J. Am. Chem. Soc.*, 2016, **138**, 8096–8103.

167 M. S. Chen, J. R. Niskala, D. A. Unruh, C. K. Chu, O. P. Lee and J. M. Fréchet, *Chem. Mater.*, 2013, **25**, 4088–4096.

168 E. Collini and G. D. Scholes, *Science*, 2009, **323**, 369–373.

169 S. P. Arnold, J. K. Harris, B. Neelamraju, M. Rudolph and E. L. Ratcliff, *Synth. Met.*, 2019, **253**, 26–33.

170 R. Noriega, J. Rivnay, K. Vandewal, F. P. V. Koch, N. Stingelin, P. Smith, M. F. Toney and A. Salleo, *Nat. Mater.*, 2013, **12**, 1038–1044.

171 M. H. Gharahcheshmeh and K. K. Gleason, *Conjugated Polymers for Next-Generation Applications*, Elsevier, 2022, ch. 9, vol. 2, pp. 283–311.

172 Y. H. Kim, C. Sachse, M. L. Machala, C. May, L. Müller-Meskamp and K. Leo, *Adv. Funct. Mater.*, 2011, **21**, 1076–1081.

173 N. Kim, S. Kee, S. H. Lee, B. H. Lee, Y. H. Kahng, Y.-R. Jo, B.-J. Kim and K. Lee, *Adv. Mater.*, 2014, **26**, 2268–2272.



174 B. J. Worfolk, S. C. Andrews, S. Park, J. Reinspach, N. Liu, M. F. Toney, S. C. Mannsfeld and Z. Bao, *Proc. Natl. Acad. Sci. U. S. A.*, 2015, **112**, 14138–14143.

175 M. N. Gueye, A. Carella, N. Massonnet, E. Yvenou, S. Brenet, J. Faure-Vincent, S. Pouget, F. Rieutord, H. Okuno, A. Benayad, R. Demadrille and J.-P. Simonato, *Chem. Mater.*, 2016, **28**, 3462–3468.

176 R. M. Howden, E. D. McVay and K. K. Gleason, *J. Mater. Chem. A*, 2013, **1**, 1334–1340.

177 Y. Lin, E. Bilotti, C. W. Bastiaansen and T. Peijs, *Polym. Eng. Sci.*, 2020, **60**, 2351–2376.

178 S. E. Root, S. Savagatrup, A. D. Printz, D. Rodriguez and D. J. Lipomi, *Chem. Rev.*, 2017, **117**, 6467–6499.

179 C. Müller, *Chem. Mater.*, 2015, **27**, 2740–2754.

180 C. M. Stafford, C. Harrison, K. L. Beers, A. Karim, E. J. Amis, M. R. VanLandingham, H.-C. Kim, W. Volksen, R. D. Miller and E. E. Simonyi, *Nat. Mater.*, 2004, **3**, 545–550.

181 Z. Wang, M. Zhu, Z. Pei, Q. Xue, H. Li, Y. Huang and C. Zhi, *Mater. Sci. Eng., R*, 2020, **139**, 100520.

182 T.-H. Le, Y. Kim and H. Yoon, *Polymers*, 2017, **9**, 150.

183 G. Rebetez, O. Bardagot, J. Affolter, J. Réhault and N. Banerji, *Adv. Funct. Mater.*, 2022, **32**, 2105821.

184 L. Bay, T. Jacobsen, S. Skaarup and K. West, *J. Phys. Chem. B*, 2001, **105**, 8492–8497.

185 N. Elgrishi, K. J. Rountree, B. D. McCarthy, E. S. Rountree, T. T. Eisenhart and J. L. Dempsey, *J. Chem. Edu.*, 2018, **95**, 197–206.

186 J.-J. Han, N. Zhang, D.-L. Liu, H. Ma, T. Han and D.-D. Sun, *Ionics*, 2020, **26**, 1029–1038.

187 T. C. Liu, W. G. Pell, B. E. Conway and S. L. Roberson, *J. Electrochem. Soc.*, 1998, **145**, 1882.

188 T. S. Mathis, N. Kurra, X. Wang, D. Pinto, P. Simon and Y. Gogotsi, *Adv. Energy Mater.*, 2019, **9**, 1902007.

189 S. Ardizzone, G. Fregonara and S. Trasatti, *Electrochim. Acta*, 1990, **35**, 263–267.

190 K. C. S. Lakshmi and B. Vedhanarayanan, *Batteries*, 2023, **9**, 202.

191 B.-Y. Chang, E. Ahn and S.-M. Park, *J. Phys. Chem. C*, 2008, **112**, 16902–16909.

192 D. Rochefort and A.-L. Pont, *Electrochim. Commun.*, 2006, **8**, 1539–1543.

193 M. R. Lukatskaya, B. Dunn and Y. Gogotsi, *Nat. Commun.*, 2016, **7**, 12647.

194 V. Augustyn, J. Come, M. A. Lowe, J. W. Kim, P.-L. Taberna, S. H. Tolbert, H. D. Abruña, P. Simon and B. Dunn, *Nat. Mater.*, 2013, **12**, 518–522.

195 S. Vijayakumar, S.-H. Lee and K.-S. Ryu, *Electrochim. Acta*, 2015, **182**, 979–986.

196 M. G. Voss, J. R. Challa, D. T. Scholes, P. Y. Yee, E. C. Wu, X. Liu, S. J. Park, O. León Ruiz, S. Subramaniyan and M. Chen, *Adv. Mater.*, 2021, **33**, 2000228.

197 F. G. Donnan, *Z. Electrochem.*, 1911, **17**, 572–581.

198 X. Jin, R. E. White and K. Huang, *J. Electrochem. Soc.*, 2016, **163**, A2702.

199 T. Sokalski, P. Lingenfelter and A. Lewenstam, *J. Phys. Chem. B*, 2003, **107**, 2443–2452.

200 P. Moni, A. Al-Obeidi and K. K. Gleason, *Beilstein J. Nanotechnol.*, 2017, **8**, 723–735.

201 K. K. Gleason, *Org. Mater.*, 2022, **4**, 261–267.

202 N. A. Kukhta, A. Marks and C. K. Luscombe, *Chem. Rev.*, 2022, **122**, 4325–4355.

203 B. X. Dong, C. Nowak, J. W. Onorato, J. Strzalka, F. A. Escobedo, C. K. Luscombe, P. F. Nealey and S. N. Patel, *Chem. Mater.*, 2019, **31**, 1418–1429.

204 B. Zayat, P. Das, B. C. Thompson and S. R. Narayan, *J. Phys. Chem. C*, 2021, **125**, 7533–7541.

205 P. Das, B. Zayat, Q. Wei, C. Z. Salamat, I.-B. Magdău, R. Elizalde-Segovia, D. Rawlings, D. Lee, G. Pace, A. Irshad, L. Ye, A. Schmitt, R. A. Segelman, T. F. Miller III, S. H. Tolbert, B. S. Dunn, S. R. Narayan and B. C. Thompson, *Chem. Mater.*, 2020, **32**, 9176–9189.

206 Y. Dobashi, A. Fannir, M. Farajollahi, A. Mahmoudzadeh, A. Usgaocar, D. Yao, G. T. M. Nguyen, C. Plesse, F. Vidal and J. D. W. Madden, *Electrochim. Acta*, 2017, **247**, 149–162.

207 K. Leš and C.-S. Jordan, *RSC Adv.*, 2020, **10**, 41296–41304.

208 G. Garcia-Belmonte, J. Bisquert, E. C. Pereira and F. Fabregat-Santiago, *J. Electroanal. Chem.*, 2001, **508**, 48–58.

209 J. Bisquert, G. G. Belmonte, F. F. Santiago, N. S. Ferriols, M. Yamashita and E. C. Pereira, *Electrochim. Commun.*, 2000, **2**, 601–605.

210 M. H. Gharahcheshmeh, E. Galstyan, A. Xu, J. Kukunuru, R. Katta, Y. Zhang, G. Majkic, X. Li and V. Selvamanickam, *Supercond. Sci. Technol.*, 2016, **30**, 015016.

211 V. Selvamanickam, R. Mallick, X. Tao, Y. Yao, M. H. Gharahcheshmeh, A. Xu, Y. Zhang, E. Galstyan and G. Majkic, *Supercond. Sci. Technol.*, 2016, **29**, 085016.

212 A. Xu, Y. Zhang, M. H. Gharahcheshmeh, L. Delgado, N. Khatri, Y. Liu, E. Galstyan and V. Selvamanickam, *IEEE Trans. Appl. Supercond.*, 2017, **27**, 1–5.

213 M. H. Gharahcheshmeh, E. Galstyan, J. Kukunuru, R. Katta, G. Majkic, X.-F. Li and V. Selvamanickam, *IEEE Trans. Appl. Supercond.*, 2017, **27**, 1–5.

214 V. Selvamanickam, M. Heydari Gharahcheshmeh, A. Xu, E. Galstyan, L. Delgado and C. Cantoni, *Appl. Phys. Lett.*, 2015, **106**, 032601.

215 M. H. Gharahcheshmeh, G. Majkic, E. Galstyan, A. Xu, Y. Zhang, X. Li and V. Selvamanickam, *Phys. C*, 2018, **553**, 26–32.

216 E. Galstyan, M. H. Gharahcheshmeh, L. Delgado, A. Xu, G. Majkic and V. Selvamanickam, *IEEE Trans. Appl. Supercond.*, 2014, **25**, 1–5.

217 P. Kovacik, G. del Hierro, W. Livernois and K. K. Gleason, *Mater. Horiz.*, 2015, **2**, 221–227.

218 R. Schmuck, R. Wagner, G. Hörpel, T. Placke and M. Winter, *Nat. Energy*, 2018, **3**, 267–278.

219 M. Li, J. Lu, Z. Chen and K. Amine, *Adv. Mater.*, 2018, **30**, 1800561.

220 C. Wang, C. Yang and Z. Zheng, *Adv. Sci.*, 2022, **9**, 2105213.

221 G. Zubi, R. Dufo-López, M. Carvalho and G. Pasaoglu, *Renewable Sustainable Energy Rev.*, 2018, **89**, 292–308.

222 S. Park, S. Y. Jeong, T. K. Lee, M. W. Park, H. Y. Lim, J. Sung, J. Cho, S. K. Kwak, S. Y. Hong and N.-S. Choi, *Nat. Commun.*, 2021, **12**, 838.



223 Y. Lin, M. Xu, S. Wu, Y. Tian, Z. Cao, L. Xing and W. Li, *ACS Appl. Mater. Interfaces*, 2018, **10**, 16400–16409.

224 D. Das, S. Manna and S. Puravankara, *Batteries*, 2023, **9**, 193.

225 Y. Chen, Y. Kang, Y. Zhao, L. Wang, J. Liu, Y. Li, Z. Liang, X. He, X. Li, N. Tavajohi and B. Li, *J. Energy Chem.*, 2021, **59**, 83–99.

226 J. Deng, C. Bae, A. Denlinger and T. Miller, *Joule*, 2020, **4**, 511–515.

227 M. Weiss, R. Ruess, J. Kasnatscheew, Y. Levartovsky, N. R. Levy, P. Minnmann, L. Stoltz, T. Waldmann, M. Wohlfahrt-Mehrens, D. Aurbach, M. Winter, Y. Ein-Eli and J. Janek, *Adv. Energy Mater.*, 2021, **11**, 2101126.

228 L.-L. Lu, Y.-Y. Lu, Z.-X. Zhu, J.-X. Shao, H.-B. Yao, S. Wang, T.-W. Zhang, Y. Ni, X.-X. Wang and S.-H. Yu, *Sci. Adv.*, 2022, **8**, eabm6624.

229 Y. Zhang, J. C. Kim, H. W. Song and S. Lee, *Nanoscale*, 2023, **15**, 4195–4218.

230 D. Lin, Y. Liu and Y. Cui, *Nat. Nanotechnol.*, 2017, **12**, 194–206.

231 Y. Han, B. Liu, Z. Xiao, W. Zhang, X. Wang, G. Pan, Y. Xia, X. Xia and J. Tu, *InfoMat*, 2021, **3**, 155–174.

232 Z. Luo, X. Qiu, C. Liu, S. Li, C. Wang, G. Zou, H. Hou and X. Ji, *Nano Energy*, 2021, **79**, 105507.

233 C. Fang, X. Wang and Y. S. Meng, *Trends Chem.*, 2019, **1**, 152–158.

234 Y. Xiao, Y. Wang, S.-H. Bo, J. C. Kim, L. J. Miara and G. Ceder, *Nat. Rev. Mater.*, 2020, **5**, 105–126.

235 W. Lyu, X. Yu, Y. Lv, A. M. Rao, J. Zhou and B. Lu, *Adv. Mater.*, 2024, **36**, 2305795.

236 Y. Lu, L. Li, Q. Zhang, Z. Niu and J. Chen, *Joule*, 2018, **2**, 1747–1770.

237 X. Yi, H. Fu, A. M. Rao, Y. Zhang, J. Zhou, C. Wang and B. Lu, *Nat. Sustainability*, 2024, **7**, 326–337.

238 A. Wang, S. Kadam, H. Li, S. Shi and Y. Qi, *npj Comput. Mater.*, 2018, **4**, 15.

239 S. Gao, F. Sun, N. Liu, H. Yang and P.-F. Cao, *Mater. Today*, 2020, **40**, 140–159.

240 G. Ma, Z. Wen, Q. Wang, C. Shen, J. Jin and X. Wu, *J. Mater. Chem. A*, 2014, **2**, 19355–19359.

241 S. Pan, J. Han, Y. Wang, Z. Li, F. Chen, Y. Guo, Z. Han, K. Xiao, Z. Yu and M. Yu, *Adv. Mater.*, 2022, **34**, 2203617.

242 S. Ahn, C. Zor, S. Yang, M. Lagnoni, D. Dewar, T. Nimmo, C. Chau, M. Jenkins, A. J. Kibler, A. Pateman, G. J. Rees, X. Gao, P. Adamson, N. Grobert, A. Bertei, L. R. Johnson and P. G. Bruce, *Nat. Chem.*, 2023, **15**, 1022–1029.

243 P. Li, G. Zhao, X. Zheng, X. Xu, C. Yao, W. Sun and S. X. Dou, *Energy Storage Mater.*, 2018, **15**, 422–446.

244 M. Salah, P. Murphy, C. Hall, C. Francis, R. Kerr and M. Fabretto, *J. Power Sources*, 2019, **414**, 48–67.

245 H. Wu and Y. Cui, *Nano Today*, 2012, **7**, 414–429.

246 H. Wu, G. Yu, L. Pan, N. Liu, M. T. McDowell, Z. Bao and Y. Cui, *Nat. Commun.*, 2013, **4**, 1943.

247 Y. Yao, N. Liu, M. T. McDowell, M. Pasta and Y. Cui, *Energy Environ. Sci.*, 2012, **5**, 7927–7930.

248 S. Bai, Y. Ma, X. Jiang, Q. Li, Z. Yang, Q. Liu and D. He, *Surf. Interfaces*, 2017, **8**, 214–218.

249 I.-H. Ko, S.-J. Kim, J. Lim, S.-H. Yu, J. Ahn, J.-K. Lee and Y.-E. Sung, *Electrochim. Acta*, 2016, **187**, 340–347.

250 J. Chen, Y. Liu, A. I. Minett, C. Lynam, J. Wang and G. G. Wallace, *Chem. Mater.*, 2007, **19**, 3595–3597.

251 T. Kennedy, E. Mullane, H. Geaney, M. Osiak, C. O'Dwyer and K. M. Ryan, *Nano Lett.*, 2014, **14**, 716–723.

252 W. Bao, G. Qian, L. Zhao, Y. Yu, L. Su, X. Cai, H. Zhao, Y. Zuo, Y. Zhang and H. Li, *Nano Lett.*, 2020, **20**, 8832–8840.

253 Q. Gan, N. Qin, Y. Zhu, Z. Huang, F. Zhang, S. Gu, J. Xie, K. Zhang, L. Lu and Z. Lu, *ACS Appl. Mater. Interfaces*, 2019, **11**, 12594–12604.

254 Y. Cao, X. Qi, K. Hu, Y. Wang, Z. Gan, Y. Li, G. Hu, Z. Peng and K. Du, *ACS Appl. Mater. Interfaces*, 2018, **10**, 18270–18280.

255 Q. Liu, Y.-T. Liu, C. Zhao, Q.-S. Weng, J. Deng, I. Hwang, Y. Jiang, C. Sun, T. Li, W. Xu, K. Du, A. Daali, G.-L. Xu, K. Amine and G. Chen, *ACS Nano*, 2022, **16**, 14527–14538.

256 L. Su, P. M. Smith, P. Anand and B. Reeja-Jayan, *ACS Appl. Mater. Interfaces*, 2018, **10**, 27063–27073.

257 R. Xiong, Y. Pan, W. Shen, H. Li and F. Sun, *Renewable Sustainable Energy Rev.*, 2020, **131**, 110048.

258 S.-M. Bak, K.-W. Nam, W. Chang, X. Yu, E. Hu, S. Hwang, E. A. Stach, K.-B. Kim, K. Y. Chung and X.-Q. Yang, *Chem. Mater.*, 2013, **25**, 337–351.

259 F. Wu, J. Liu, L. Li, X. Zhang, R. Luo, Y. Ye and R. Chen, *ACS Appl. Mater. Interfaces*, 2016, **8**, 23095–23104.

260 P. Rozier and J. M. Tarascon, *J. Electrochem. Soc.*, 2015, **162**, A2490.

261 L. Su, J. L. Weaver, M. Groenenboom, N. Nakamura, E. Rus, P. Anand, S. K. Jha, J. S. Okasinski, J. A. Dura and B. Reeja-Jayan, *ACS Appl. Mater. Interfaces*, 2021, **13**, 9919–9931.

262 Y. Yang, G. Yu, J. J. Cha, H. Wu, M. Vosgueritchian, Y. Yao, Z. Bao and Y. Cui, *ACS Nano*, 2011, **5**, 9187–9193.

263 L. Xiao, Y. Cao, J. Xiao, B. Schwenzer, M. H. Engelhard, L. V. Saraf, Z. Nie, G. J. Exarhos and J. Liu, *Adv. Mater.*, 2012, **24**, 1176–1181.

264 Z. Wang, R. Pan, C. Xu, C. Ruan, K. Edström, M. Strømme and L. Nyholm, *Energy Storage Mater.*, 2018, **13**, 283–292.

265 W. Liu, D. Lin, A. Pei and Y. Cui, *J. Am. Chem. Soc.*, 2016, **138**, 15443–15450.

266 H. Zhang, Y. Cao, H. Yang, S. Lu and X. Ai, *J. Polym. Sci., Part B: Polym. Phys.*, 2013, **51**, 1487–1493.

267 P. R. Das, L. Komsa, O. Osters and G. Wittstock, *J. Electrochem. Soc.*, 2015, **162**, A674.

268 H. Zhong, A. He, J. Lu, M. Sun, J. He and L. Zhang, *J. Power Sources*, 2016, **336**, 107–114.

269 D. Shao, H. Zhong and L. Zhang, *ChemElectroChem*, 2014, **1**, 1679–1687.

270 L. L. Zhang and X. S. Zhao, *Chem. Soc. Rev.*, 2009, **38**, 2520–2531.

271 Y. Meng, K. Wang, Y. Zhang and Z. Wei, *Adv. Mater.*, 2013, **25**, 6985–6990.

272 A. Liu, P. Kovacik, N. Peard, W. Tian, H. Goktas, J. Lau, B. Dunn and K. K. Gleason, *Adv. Mater.*, 2017, **29**, 1606091.

273 M. I. Ahamed, R. Boddula and T. Altalhi, *Flexible Supercapacitor Nanoarchitectonics*, John Wiley & Sons, 2021.



274 J. A. Lee, M. K. Shin, S. H. Kim, H. U. Cho, G. M. Spinks, G. G. Wallace, M. D. Lima, X. Lepró, M. E. Kozlov and R. H. Baughman, *Nat. Commun.*, 2013, **4**, 1970.

275 V. T. Le, H. Kim, A. Ghosh, J. Kim, J. Chang, Q. A. Vu, D. T. Pham, J.-H. Lee, S.-W. Kim and Y. H. Lee, *ACS Nano*, 2013, **7**, 5940–5947.

276 Z. Yang, J. Deng, X. Chen, J. Ren and H. Peng, *Angew. Chem., Int. Ed.*, 2013, **52**, 13453–13457.

277 J. Huang, K. Wang and Z. Wei, *J. Mater. Chem.*, 2010, **20**, 1117–1121.

278 A. J. Heeger, *J. Phys. Chem. B*, 2001, **105**, 8475–8491.

279 D. Villers, D. Jobin, C. Soucy, D. Cossement, R. Chahine, L. Breau and D. Bélanger, *J. Electrochem. Soc.*, 2003, **150**, A747.

280 S. Hashmi and H. M. Upadhyaya, *Solid State Ionics*, 2002, **152**, 883–889.

281 D. P. Dubal, S. H. Lee, J. G. Kim, W. B. Kim and C. D. Lokhande, *J. Mater. Chem.*, 2012, **22**, 3044–3052.

282 G. Yu, L. Hu, N. Liu, H. Wang, M. Vosgueritchian, Y. Yang, Y. Cui and Z. Bao, *Nano Lett.*, 2011, **11**, 4438–4442.

283 Y. Y. Smolin, K. L. V. Aken, M. Boota, M. Soroush, Y. Gogotsi and K. K. S. Lau, *Adv. Mater. Interfaces*, 2017, **4**, 1601201.

284 M. Liu, Y.-E. Miao, C. Zhang, W. W. Tjiu, Z. Yang, H. Peng and T. Liu, *Nanoscale*, 2013, **5**, 7312–7320.

285 W. Wang, Q. Luo, B. Li, X. Wei, L. Li and Z. Yang, *Adv. Funct. Mater.*, 2013, **23**, 970–986.

286 Z. Yang, J. Zhang, M. C. W. Kintner-Meyer, X. Lu, D. Choi, J. P. Lemmon and J. Liu, *Chem. Rev.*, 2011, **111**, 3577–3613.

287 A. Khor, P. Leung, M. R. Mohamed, C. Flox, Q. Xu, L. An, R. G. A. Wills, J. R. Morante and A. A. Shah, *Mater. Today Energy*, 2018, **8**, 80–108.

288 N. Goujon, N. Casado, N. Patil, R. Marcilla and D. Mecerreyres, *Prog. Polym. Sci.*, 2021, **122**, 101449.

289 A. Dinesh, S. Olivera, K. Venkatesh, M. S. Santosh, M. G. Priya, I. Inamuddin, A. M. Asiri and H. B. Muralidhara, *Environ. Chem. Lett.*, 2018, **16**, 683–694.

290 X. Wang, M. Ding and C. Jia, *J. Phys. D: Appl. Phys.*, 2024, **57**, 423002.

291 A. Ashok and A. Kumar, *Green Chem. Lett. Rev.*, 2024, **17**, 2302834.

292 K. Gong, F. Xu, J. B. Grunewald, X. Ma, Y. Zhao, S. Gu and Y. Yan, *ACS Energy Lett.*, 2016, **1**, 89–93.

293 H. Zhang and C. Sun, *J. Power Sources*, 2021, **493**, 229445.

294 R. Wang, Y. Li, Y. Wang and Z. Fang, *Appl. Energy*, 2020, **261**, 114369.

295 R. Wang, Y. Li and Y.-L. He, *J. Mater. Chem. A*, 2019, **7**, 10962–10970.

296 H. Wang, J. Lin and Z. X. Shen, *J. Sci.: Adv. Mater. Devices*, 2016, **1**, 225–255.

297 X. Lu, W. Zhang, C. Wang, T.-C. Wen and Y. Wei, *Prog. Polym. Sci.*, 2011, **36**, 671–712.

298 J. Winsberg, T. Hagemann, T. Janoschka, M. D. Hager and U. S. Schubert, *Angew. Chem., Int. Ed.*, 2017, **56**, 686–711.

299 Y. Ding, C. Zhang, L. Zhang, Y. Zhou and G. Yu, *Chem. Soc. Rev.*, 2018, **47**, 69–103.

300 L. Zeng, T. S. Zhao, L. Wei, H. R. Jiang and M. C. Wu, *Appl. Energy*, 2019, **233–234**, 622–643.

301 M. Vagin, C. Che, V. Gueskine, M. Berggren and X. Crispin, *Adv. Funct. Mater.*, 2020, **30**, 2007009.

302 E. K. Radwan, R. A. Omar, A. M. Ali, A. S. Elsayed and E. N. El Sawy, *Sustainable Mater. Technol.*, 2024, e01069.

