

Showcasing a review on the materials design and charge storage perspective of organic electrodes from Dr. Amitava Banerjee at Department of Metallurgical and Materials Engineering, Indian Institute of Technology Jodhpur, India and the group of Prof. Rajeev Ahuja, at the Matter Theory Division, Department of Physics and Astronomy, Uppsala University, Sweden.

Promise and reality of organic electrodes from materials design and charge storage perspective

Critical overview on the advancement of the materials design along with the charge storage mechanisms of organic battery electrodes of monovalent to multivalent alkali ions.





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# Promise and reality of organic electrodes from materials design and charge storage perspective

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Organic electrode materials are becoming increasingly important as they reduce the C-footprint as well as the production cost of currently used and studied rechargeable batteries. With increasing demand for high-energy-density devices, over the past few decades, various innovative new materials based on the fundamental structure–property relationships and molecular design have been explored to enable high-capacity next-generation battery chemistries. One critical dimension that catalyzes this study is the building up of an in-depth understanding of the structure–property relationship and mechanism of alkali ion batteries. In this review, we present a critical overview of the progress in the technical feasibility of organic battery electrodes for use in long-term and large-scale electrical energy-storage devices based on the materials designing, working mechanisms, performance, and battery safety. Specifically, we discuss the underlying alkali ion storage mechanisms in specific organic batteries, which could provide the designing requirements to overcome the limitations of organic batteries. We also discuss the promising future research directions in the field of alkali ion organic batteries, especially multivalent organic batteries along with monovalent alkali ion organic batteries.

#### 1. Introduction

Organic electrode materials have drawn significant attention in the last few years in the search of suitable electrode materials to meet the requirements of the rapidly growing markets of consumer electronics, electric vehicles, and grid integration of renewable energy.1 Organic electrodes are endowed with several merits over conventional transition metal based inorganic electrodes, such as sustainable production with low carbon footprint, low cost, excellent stability, high flexibility, as well as resource sustainability.2-5 In terms of sustainability, many organic electrode materials can be synthesized from biomass using green-chemistry techniques that have negligible environmental foot print. For example, the organic electrode dilithium rhodizonate (Li<sub>2</sub>C<sub>6</sub>O<sub>6</sub>) can be prepared from the natural sugar myo-inositol, which exists in corn plants in the hexaphosphate form.6 Another useful redox active molecule, polyguinone can be prepared by the polycondensation of malic acid, which is a natural product found in apples.<sup>7</sup> On the other hand, the diversity of organic functional groups and flexibility

of organic molecules allow the molecular level modifications of

traced back to as early as the 1960s, when a tricarbonyl compound was first used as a cathode for lithium-ion batteries (LIBs).12 However, a rapid decay in capacity due to the high solubility of small carbonyl compounds in aprotic electrolytes limited their further development. Subsequently, in the 1970s, many chemists explored conductive polymers as LIB electrode materials due to their low solubility in electrolytes. 13-16 In this category, the most widely studied conductive polymers are polyacetylene, polyaniline, polypyrrole, polythiophene, and poly(p-phenylene). <sup>13,16,17</sup> In the late 1980s, coin-type batteries <sup>18</sup> with polyaniline cathodes and LiAl alloy anodes were introduced but quickly disappeared from the market as they never showed their ideal performance. The main concern with the conductive polymers is low doping level limits, typically below 50%, which means in the charging/discharging process less than half of the redox-active groups participate. Due to this reason the practical capacities usually do not exceed 150 mA h g<sup>-1</sup>, which motivated the search for alternative chemistry for cathode materials. 13,16 In the development of alternative chemistry to obtain high capacity cathode materials, attention was paid to organosulfur compounds where the S-S bond formation (cleavage) takes place during the charge (discharge) process. Organosulfur compounds exhibit high capacities. 19 But there are other concerns, such as the sluggish kinetics of S-S bond formation in the charging process, poor cycling stability, and a serious shuttle effect (where dissolved

organic electrodes to alter their alkali ion storage capacity. 1,8-11 The history of organic electrode materials (Fig. 1) can be

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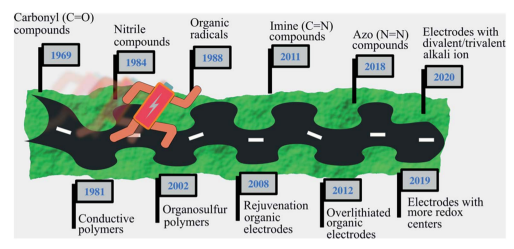


Fig. 1 Schematic of the historical development of organic electrodes. Here each year represents the reporting<sup>6,7,12,14,19,21,22,31–35</sup> of new developments either in the form of new chemistry or new redox mechanisms.

species diffuse between the anode and cathode).20 Along with this development, electrode materials based on nitriles, electron-poor molecules with highly reversible redox chemistry, were also investigated.21 Besides the above-mentioned materials, organic radicals were tested as a promising organic electrode material due to their attractive rapid redox compatibility (where bond rearrangements during charging and discharging are almost nil).22 In fact, they possessed a higher electrontransfer rate compared to organosulfur compounds; for example, in nitroxyl, the self-exchange electron transfer (RNO<sup>+</sup> + RNO'  $\rightleftharpoons$  RNO' + RNO') rate constant is  $\sim 10^{-1}$  cm s<sup>-1</sup>, which is higher than those of organosulfur compounds ( $\sim 10^{-8}$  cm s<sup>-1</sup>).<sup>23,24</sup> However, after 1991, the research and development on organic electrode materials received less attention due to the successful commercialization of inorganic electrode materialsbased (cathode: LiCoO2; anode: graphite) high voltage LIBs,25 and only a few studies were published up to early 2000. From 2002 organic electrode materials again began to gain more attention, when organic radicals were also reported as high voltage cathodes for LIBs.22 The search for organic electrode materials got a boost around 2008, when Armand and Tarascon<sup>7</sup> pointed out the environmental importance and bright prospect of organic electrode materials. In the last few years, along with the advancement of Li-chemistry, search for organic materials is also extended beyond the Li chemistry, such as other monovalent (e.g. K), divalent (e.g. Mg, Ca, Zn) and trivalent cation (e.g. Al) chemistry.26,27 Although there are many reviews of organic electrode materials, 2,16,20,28-30 it is noteworthy that a comprehensive review focusing on the alkali ion storage mechanism along with the structural design strategies to improve the electrochemical performance of various metal-ion batteries in parallel to the practical requirements is still missing. In this review, we first present the general charge storage mechanism of organic electrode materials. We then discuss more insight into the materials chemistry specific mechanism, and the strategies applied to increase the electrochemical performance (such as output voltage, capacity, cycling stability, and rate performance), and the designing

requirements to further enhance the performance to reach the practical application goal. We also discuss the key challenges of existing organic electrode materials and the future prospects of this field beyond the monovalent alkali ion organic batteries.

## General charge/discharge mechanism

In general, the charge/discharge mechanism of organic electrodes is based on the redox reactions of the electrode materials, where the charge state of the electroactive groups changes. In contrast to transition metal-based electrodes, usually an organic polymer is processed in the uncharged form.<sup>28</sup> Due to

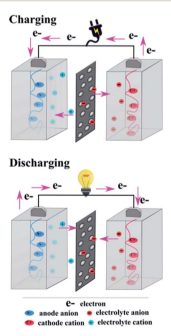


Fig. 2 Schematic representation of the general charge—discharge process in all organic batteries.

this reason, initially the cathode is oxidized and anode is reduced. The general charge/discharge process is shown in Fig. 2, where the electrodes are placed in an ion-conducting electrolyte and an ion-permeable membrane is used to separate them. This ion-conducting electrolyte plays an important role in both charge and discharge processes. In the charge process, electrolyte ions are compensated by the charge formed in the electrode, whereas in the discharge process, the counter ions migrate from the polymer into the electrolyte. There are few fundamental differences between inorganic and organic electrode materials. During the charge/discharge process, an inorganic electrode undergoes alloying or conversion reactions; as a result, the electrode experiences a large structural change.36 On the other hand, the redox reactions of most of the organic electrodes manifest very minimal structural changes with limited bond rearrangement. 37,38 But there are some exceptions as well in the case of organic materials, e.g. disodium rhodizonate and Na<sub>2</sub>(2,5-dihydroxy-1,4-benzoquinone) show relatively large structural changes during the charge/discharge process.<sup>39-41</sup> Another important difference between inorganic and organic electrode materials is the characteristic of the counterion. In the latter, the redox reaction chemistry is not influenced by the radius and charge of the counterions, which is a great advantage of organic electrode materials, so that the same organic electrodes could be used in different aqueous or non-aqueous metal-ion batteries. For example, perylene dianhydride is utilized as an electrode for Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, and Ca<sup>2+</sup> counter ions, where the radius and charge of the counter ions are different. 42-45 Based on the charge-state change of the active ion, overall the organic electrode materials are classified into three different categories:28 n-type, p-type and bipolar type. The general mechanism of the redox chemistry of various organic electrode materials is shown in Fig. 3. The redox reaction in the n-type electrode takes place between the neutral and negatively charged states, whereas in the p-type electrode the reaction occurs between the neutral and positively charged states. Mostly n-type organics are first reduced and then they bind with metal counterions. Conversely, p-type organics are first oxidized before binding with the anions from the electrolyte. Organic electrodes belonging to the bipolar type can be oxidized or reduced first depending on the applied voltage. It is noteworthy that the particular redox chemistry of many organic electrodes is still unclear and in-depth investigation is needed. For instance, during the lithiation process some organic materials (n-type and bipolar type) are believed to exist as radical intermediates, but it is very difficult to identify these types of intermediates due to their poor stability in batteries.46 Also, the reversibility of the over-lithiation process is still under debate.34,47

### Classification of organic electrode materials

#### 3.1 Carbonyl group

In the context of energy storage, carbonyl compounds were first explored sporadically in the 1970s and 80s but they were studied

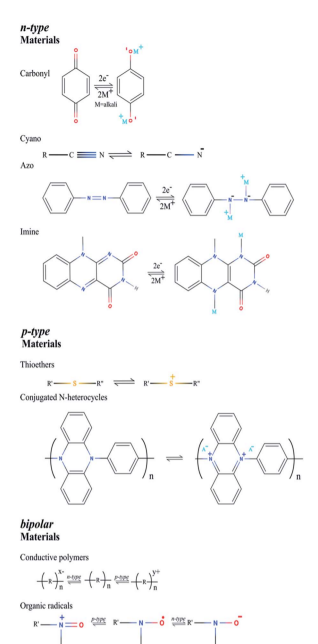


Fig. 3 General mechanism of the redox chemistry of various organic electrode materials.

intensively over the past few decades. 48-52 One of the reasons could be that chemists can extract them by a variety of synthetic means as they are one of the common organic functionalities. Also, they can show oxidative stability as well as reversible reduction capability. The major structural changes (which are eventually the driving force of their electrochemical mechanism) that occur upon reduction are (A) the vicinal carbonyls form stable enolates, (B) whereas aromatic carbonyl groups disperse the negative charge over the delocalized system. Depending on their practical working potentials, they can serve as cathodes as well as anodes. As electrode materials, carbonyl compounds (quinones, carboxylates, anhydrides, imides, and

ketones) are mostly used in the form of small molecules and non-conjugated polymers. A detailed review about carbonyls for battery applications including polymers as well as small molecules is available.<sup>53</sup> Here we will first summarize the potentiality as well as the inferiority of the small molecule form then the non-conjugated polymer form.

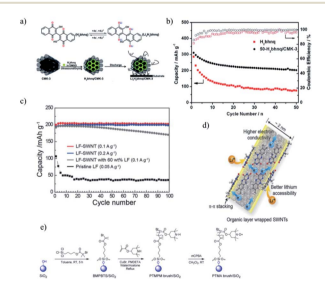
3.1.1 Quinone. Small carbonyl molecules, particularly quinones, are widely used as cathode materials in organic metal-ion batteries because of their high capacity (sometimes beyond the conventional inorganic electrode) and fast reversible electrochemistry (or fast kinetics). For example, 1,4-benzoquinone can exhibit a discharge voltage of 2.7 V (vs. Li<sup>+</sup>/Li) and a theoretical capacity of 496 mA h g<sup>-1</sup>, which are better than those of the conventional cathode LiCoO<sub>2</sub> (155 mA h g<sup>-1</sup>).<sup>54</sup> However, small carbonyl compounds generally suffer from severe dissolution in the electrolyte (causing capacity fadeout during cycling in aqueous and non-aqueous metal-ion batteries), inferior electronic conductivity (restricting the rate performance), and low voltage limit. In the following section, we will discuss the possible remedies that have been suggested to enhance their conductivity, cycling stability (or to decrease the solubility) and reduction potentials, especially to critically justify their advantage of high capacity.

3.1.1.1 Fusing multiple carbonyl groups in the compound. Zou et al. 55 found that increasing the number of aromatic rings with the carbonyl group could enhance the solubility and capacity of the small molecule, quinone. The tetrahydrohexaquinone synthesized by them retained 54.1% and 26.5% of its theoretical capacity (628 mA h g<sup>-1</sup>), respectively, at 200 mA g<sup>-1</sup> and 800 mA g<sup>-1</sup>. Although the presence of large aromatic structures decreases the solubility (compared to its parent anthraquinone) and increases the yield of theoretical capacity, the presence of redox groups in close proximity could also cause electrostatic repulsion during reduction. This could lead to weak crystal packing and, finally, dissolution and capacity fading. Eventually, Zou et al.55 observed 41% loss of initial capacity after 40 cycles.

3.1.1.2 Functionalization of quinones with ionic groups. Addition of ionic groups is another strategy to improve the stability as well as the working potential of the electrode. Mono and disodium sulfate anthraquinones produce a high specific capacity, respectively, on the order of 130 and 150 mA h g<sup>-1</sup> at 0.2C.<sup>59</sup> Also, the presence of an additional electron-withdrawing group in disodium sulfate anthraquinone helps to provide better cycling stability, ~92% after 100 cycles at 0.1C.<sup>59</sup> Yokoji and co-workers<sup>60</sup> observed a 600 meV increase in the reduction potential of benzoquinones by employing a fluorinated electron-withdrawing group. They also observed a slight increase in stability, which could be due to the stable intermediate formation but it did not fully suppress the rapid capacity fading issue due to dissolution. Kim et al.61 provided a DFT based design strategy to understand the effect of halogen substitution on quinone. Their designed tetra-chloroquinone for the sodium-ion battery cathode exhibited a capacity of 150 mA h  $g^{-1}$  at 10 mA  $g^{-1}$  with high voltage plateaus at 2.6 and 2.9 V vs. Na/Na<sup>+</sup>. Unfortunately, it lost 95% of its initial capacity 20 cycles. The positive effect of ionic group

functionalization on voltage is impressive as it also enhances the stability slightly in a few cases. But it also has a negative effect on theoretical capacity due to the addition of inactive mass to the compound without enhancing its electron acceptance capability. Therefore, a clear trade-off is necessary between the theoretical capacity and reduction potential to apply this strategy.

3.1.1.3 Entrapping within the insoluble substrate. The cycling stability of the small molecule could be improved by impregnating the redox small molecule within an insoluble substrate, such as various carbon materials, SiO<sub>2</sub> nanoparticles, etc. as shown in Fig. 4. The thought behind this strategy is that strong interaction (which could be physical, chemical, or both) between the substrate and the organic molecule could reduce the dissolution. Li et al. 57 applied this strategy by impregnating bis-naphthoquinone within mesoporous carbon CMK-3 to improve the cycling stability of bis-naphthoguinone. The observed capacity was  $\sim$ 100% of its theoretical capacity at 0.1C and it retained  $\sim$ 66% of the initial capacity after 50 cycles. This strategy no doubt improves the cycling stability and active material usage but cannot relieve the dissolution problem completely. Another example of this strategy is the lumiflavine (LF) composite with single-walled carbon nanotubes (SWCNTs), where pi-pi interaction (physical interaction) occurs between the aromatic structured LF and nanotubes.<sup>56</sup> This approach could be extended to anthraquinone also. The composite with 50 wt% of LF retains a high capacity of 99.7% after 100 cycles but increasing the LF loading beyond 60 wt% causes poor cycle performance, which may be due to the lower pi-pi interaction



(a) Schematic of the electrochemical redox mechanism of bisnaphthoquinone and electrode reaction of the bis-naphthoquinone/ CMK-3 nanocomposite, (b) cycling performance and the corresponding coulombic efficiency of the bis-naphthoquinone/CMK-3 nanocomposite at a rate of 0.1C, (c) capacity retention of LF-SWNT electrodes and pristine LF electrodes, (d) pi-pi interaction of the organic nanolayer of LF on SWNTs, (e) synthetic route of PTMA-based brush/SiO<sub>2</sub> composites. The figures are reproduced from ref. 56–58 with permission. © Royal Society of Chemistry, John Wiley and Sons, Elsevier.

with excess LF on SWCNTs. Therefore, one of the drawbacks of this strategy is the limited loading of active mass, which certainly limits the energy density. In addition to carbon materials, Lin et al. employed SiO2 nanoparticles to make a composite with poly(2,2,6,6-tetramethylpiperidin-1-oxyl-4-yl methacrylate) (PTMA).58 Although this is not an example of a single molecule, we are only discussing this example here to emphasize the similar strategy. Here PTMA is grafted covalently on the nanoparticles. Thanks to this strong (chemical) interaction, which controlled the dissolution of active PTMA, it showed a long cycle life of 300 cycles. So, impregnating (grafting) organic molecules on conductive insoluble substrates could enhance the capacity and cycling stability, which may need to improve further for practical use. A combination of this strategy with other unique strategies will be required to deal with the solubility issue of small molecule cathodes.

3.1.1.4 Partial replacement of carbonyl to thiocarbonyl. It was observed that partial replacement of the carbonyl group in anthraquinone by a thio group (from thiocarbonyl) could enhance its cycling stability and working potential.62 This material retained 33.8% of the initial capacity after 40 cycles and showed two discharge plateaus at 2.7 and 2.1 V vs. Li/Li<sup>+</sup>. Although it improved the cycling stability, working potential, and charge carrier mobility of anthraquinone, the stability is far from ideal due to the side reaction and dissolution.

3.1.1.5 Introducing nitrogen-containing heterocyclics. It is another way to improve the voltage of organic carbonyl cathodes without having a negative effect on the theoretical capacity. Shimizu et al.63 investigated this effect on anthraquinone and phenanthrenequinone in lithium-ion batteries as shown in Fig. 5. They found that introducing "N" at 1, 4, 5, and 8 positions enhances the reduction potential of phenanthrenequinone from 2.52 V to 2.73 V and 2.94 V vs. Li/Li<sup>+</sup>. But the capacity fadeout issue of those molecules still persists. Therefore, combining this strategy with other strategies, such as porous carbon entrapping or the addition of an ionic group (EWG) could improve the small molecule electrode performance.

3.1.1.6 Crystalline nanostructure. Besides the molecular engineering approach, this is another method to tune the cycling stability and electrochemical performance of the small organic molecule. Crystalline nanowires of croconic acid disodium salt (CADS) not only overcome the barriers of low

2.13 V 9,10-anthraquinone 2.73 V 4,5-diaza-9,10-phenanthrenequinone

Fig. 5 The average voltages of nitrogen-containing heterocyclics, anthraquinone and phenanthrenequinone.

electronic conductivity of CADS and lithiation-induced strain but also exhibit much better electrochemical performance than their crystal bulk material and microwire counterpart.64 As shown in Fig. 6, the crystalline nanowire could retain ~100% of its initial capacity after 110 cycles at 0.2C.

3.1.1.7 Aromaticity tuning. To improve the voltage of the small organic molecule cathodes, Wu and co-workers<sup>65</sup> studied the correlation between the aromaticity (electron delocalization) and the discharge potential of carbonyl-containing polycyclic aromatics. Their study revealed that high discharge potential could be achieved if the aromaticity increases after lithiation. This aromaticity change is expressed by the average change of Clar sextet numbers ( $\Delta C_{2Li}$ ):

$$\Delta C_{2Li} = \Delta C/(0.5 \times \Delta Li)$$

where  $\Delta C_{2Li}$  is the average change of Clar sextet numbers after two Li atoms are absorbed,  $\Delta C$  is the change of Clar sextet numbers during lithiation or the difference in aromaticity in the neutral and reduced compound, and  $\Delta Li$  is the number of adsorbed Li atoms. Therefore, the idea is to have higher numbers of Clar sextet to improve the reduction voltage (Fig. 7). With this approach, the authors designed the quinone to have an average voltage of 2.77 V vs. Li/Li<sup>+</sup>.

3.1.1.8 Impact of phase change. Phase transformation during the charge-discharge cycle could also affect the capacity of organic materials. For example, during the sodiation process the alpha-phase of disodium rhodizonate (Na<sub>2</sub>C<sub>6</sub>O<sub>6</sub>) transforms into the gamma-phase, but the reverse phase transformation during desodiation is kinetically suppressed as observed by Bao et al.39 This irreversible phase transformation results in the low practical capacity of Na2C6O6. Bao and co-workers revealed that without molecular design, reducing the particle size and optimizing the electrolyte could deliver a capacity value (484 mA h  $g^{-1}$ ) very close to the theoretical capacity (501 mA h  $g^{-1}$ , 4Na). A similar report of capacity fadeout and cycling instability on dilithium rhodizonate (Li2C6O6) due to crystal structure change was published by Kim and coworkers.41 Therefore, not only molecular design but also crystal structure design and electrode morphology are crucial factors in enhancing the performance of carbonyl systems.

Simple quinone compounds usually show serious capacity fading and as a consequence generate a shuttle effect. This is due to the organic species dissolution in the electrolyte. As we noticed, several strategies are applied to tackle this problem, such as molecular design, loading with advanced carbon mass,

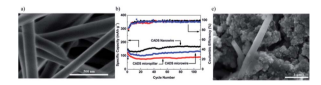


Fig. 6 (a) SEM images of CADS nanowires, (b) cycle life of CADS micropillars, CADS microwires, and CADS nanowires, (c) SEM images of CADS nanowires after 100 cycles. The figures are reproduced from ref. 64 with permission. © American Chemical Society.

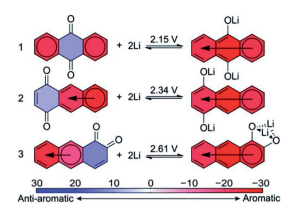


Fig. 7 Correlation between the change in aromaticity and the reduction potential of the small molecule anthracene with carbonyl groups where the arrow signifies the migration of Clar sextets. The figure is reproduced from ref. 65 with permission. © Royal Society of Chemistry.

and phase change. In addition to these, there are other possible strategies such as increasing the viscosity of the electrolyte, increasing the polarity of the salt, and polymerization. In the following section, we will show how polymerization of the small electrode molecule could prevent the dissolution of the electrode material.

**3.1.2** Carboxylate. Molecules with carboxylate groups are mostly considered as anode materials. Out of the many carboxylate group molecules, terephthalates (and their salts) are widely used as anodes for metal-ion organic batteries. Similar strategies as discussed before could also be applied (in a complementary fashion) to these molecules to enhance their capacity and cycling stability. Methods to enhance this group molecules' performance are summarized below.

3.1.2.1 Position of the carbonyl group: ortho vs. para. Within the various molecular engineering approaches, Gottis et al. 67 introduced a smart way to increase the voltage of carbonyl group-containing molecules without adding EWGs or electronegative atoms. So, this approach will not affect the theoretical capacity of the parent molecule. The authors found that the presence of carbonyl groups in the ortho position of the lithiated enolates delivers a voltage of 300 mV over its para position (as shown in Fig. 8(a)) due to the favourable coulombic interaction. Also, they observed that this enolate retains  $\sim 100\%$  of its initial capacity after 30 cycles.

3.1.2.2 Electrode processing methods. The electrochemical performance of the carboxylate group molecules could be enhanced by changing the processing methods. Zhang et al.  $^{66}$  reported that synthesized lithium terephthalate porous microspheres (via a spray drying method) with an N-doped carbon layer coating could improve not only the cycling stability and rate capability of the resultant electrode but also the electronic conductivity and diffusion of lithium ions. At the lowest rate of 0.05C, it gives the highest specific capacity of 259 mA h g $^{-1}$ , and 67.9% of this capacity is retained after 50 cycles (Fig. 8). Zhang et al.  $^{66}$  observed that these performance values are higher compared to standard electrode processing and formulations.

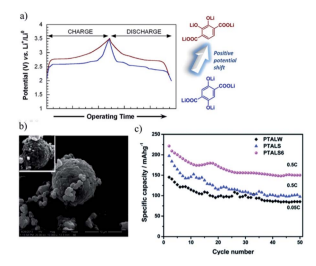


Fig. 8 (a) Charge—discharge curves of dihydroxyterephthaloyl derivatives with two—OLi in *para* and *ortho* positions, (b) SEM image of the N-doped C-coated sphere of lithium terephthalate, (c) the cycling performance of the bulk lithium terephthalate (PTALW), porous spraydried lithium terephthalate (PTALS) and carbon-coated spray-dried lithium terephthalate (PTALS6). The figures are reproduced from ref. 66 with permission. © Royal Society of Chemistry.

3.1.2.3 Conjugated carboxylates: linear vs. cyclic. Lee and coworkers observed an interesting effect of conjugation in linear as well as cyclic carboxylate compounds.68 In the case of the cyclic system, dilithium terephthalate, Li<sub>2</sub>TP, during discharge, shows one plateau at 0.81 V and another sloping plateau, from 0.8 to 0.0 V. It delivers a specific capacity (522 mA h  $g^{-1}$  at 30 mA  $g^{-1}$ ) ~73% higher than its theoretical capacity based on one lithium insertion per carboxylate group. The thiophene derivative of this family, dilithium thiophene-2,5-dicarboxylate, Li2TDC, also delivers a very high specific capacity of 850 mA h  $g^{-1}$ . However, its linear system,  $Li_2C_6H_6O_4$ , does not deliver the excess capacity. By performing the C13 isotope labeling experiment, Lee et al.68 concluded that the excess capacity is coming from the internal alkene of the cyclic system, which can accommodate the extra lithium ions. But the linear system breaks the conjugation of the structure. The chargedischarge mechanism of these molecules with the corresponding reduction potentials is shown in Fig. 9. Based on this mechanism, the "super-lithiated" compound was also explored,47 where the obtained capacity was very high. But its long sloping discharge plateau limited its application in practical use, so the charge storage was limited. Therefore "conjugation" could be an efficient way to improve the capacity of small molecule electrodes.

3.1.2.4 Extension of conjugation. We mentioned in the previous section that conjugation could improve the capacity of small molecules. Now we will see the effect on the performance if we extend the conjugation. It is observed that the extension of electronic conjugation between the carboxylate can increase the rate capability of 2,6-naphthalene dicarboxylate compared to  ${\rm Li_2TP.^{69}}$  It gives a specific capacity of 200 mA h g<sup>-1</sup> at 0.1C and 176 mA h g<sup>-1</sup> at 1C and retains  ${\sim}65.3\%$  capacity after 50 cycles

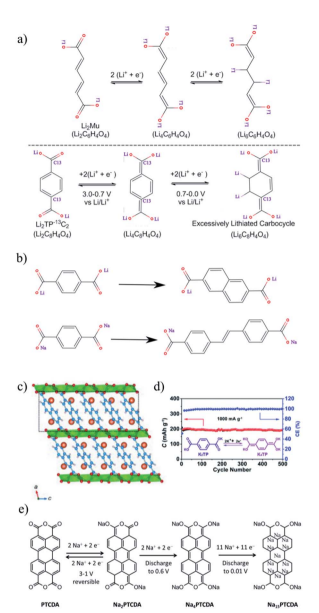


Fig. 9 (a) Mechanism of extra capacity in conjugated carboxylates of the cyclic Li<sub>2</sub>TP system compared to the linear Li<sub>2</sub>C<sub>6</sub>H<sub>6</sub>O<sub>4</sub> system, (b) extended conjugation of Li<sub>2</sub>TP to the 2,6-naphthalene dicarboxylate molecule and SBDC to SSDC, (c) lithium-intercalated crystal structure of 2,6-Naph(COOLi)<sub>2</sub> and the structure of Li-doped naphthalene rings with  $\pi$ -stacking (intercalated Li = orange, Li = yellow, O = red, C = blue, and H= white), (d) cycling performance of  $K_2TP$  at 1 A  $g^{-1}$ , (e) schematic diagrams for the proposed electrochemical reactions during sodiation/desodiation of PTCDA. The figures are reproduced from ref. 45, 72, 73 and 76 with permission. © Royal Society of Chemistry, American Chemical Society, WILEY-VCH.

at 1C. Perylene tetracarboxylate is another example in this context of achieving high rate capability, by extending the aromatic core between carboxylate groups.70 This compound could achieve 95% of its theoretical capacity (233 mA h g<sup>-1</sup>) at 1.25C and could retain  $\sim$ 47% of it at 1.25C after 100 cycles. Therefore, the authors improved the electrode rate capabilities by using a large aromatic core at the cost of cycling stability. In this effort, another possibility is formation of a highly

crystalline electrode, 2,6-naphthalene dicarboxylate dilithium. Here the metal-organic framework (Fig. 9) helps to pack the aromatic ring closely, which eventually enhances the electronic and ionic conductivity of the electrode. The metal-organic framework of this compound retains 100% capacity after 10 cycles at 0.1C.71,72 However, the number of cycles in the experiment is too low to comment on the high cycle stability.

Extended  $\pi$  conjugation is also important to enhance the intermolecular interactions and layer-by-layer packing, which could be a fast route for alkali ion diffusion between two layers as well as charge transport. Introducing a carbon-carbon double bond linkage in sodium benzene-dicarboxylate (SBDC) could extend its  $\pi$ -conjugated system to form a sodium 4,4'stilbene-dicarboxylate (SSDC) compound<sup>73</sup> (Fig. 9). SBDC exhibits a specific capacity of 260 mA h g<sup>-1</sup> at 50 mA g<sup>-1</sup> and 72 mA h  $g^{-1}$  at 10 A  $g^{-1}$  and SSDC shows 192 mA h  $g^{-1}$  at 50 mA  $g^{-1}$  and 22 mA h  $g^{-1}$  at 10 A  $g^{-1}$ . The cyclability performance under a high current density of 1 A g<sup>-1</sup> showed that the capacity is still higher than 112 mA h  $\rm g^{-1}$  even after 400 cycles with retention higher than 70%. These results imply that the extension of the  $\pi$ -conjugated system is an efficient strategy to improve the high rate performance.

3.1.2.5 Degree of deprotonation. In the effort to enhance the cycle stability, the degree of deprotonation is another alternative way. For example, the degree of deprotonation of the carboxylic acid (COOH) groups in 4,4'-biphenyldicarboxylate (bpdc) sodium salts affects their electrochemical performance and corresponding reaction mechanisms.74 Choi et al.74 observed that bpdc needs to be fully deprotonated to exhibit a promising electrochemical performance with a reversible capacity of about 200 mA h g<sup>-1</sup> at ca. 0.5 V vs. Na/Na<sup>+</sup>, stable cycle performance over 150 cycles, and excellent rate performance of 100 mA h g<sup>-1</sup> even at a 20C rate.<sup>74</sup> The electrochemical performance is better in the fully deprotonated bpdc-sodium salt than in the partially deprotonated bpdc-sodium salt and complete deprotonation is required for good stability during cycling.

3.1.2.6 Choice of counter ion. It is observed that the choice of the counter ion itself is also important to tune the cycle stability of the small molecule electrode. For instance, the carboxylate derivative, benzene diacrylate as an anode for the sodium-ion battery shows the capacity fading problem. It delivers a specific capacity of 177.7 mA h g<sup>-1</sup> at 0.025C with a 91% coulombic efficiency, which decreases to 40 mA h g<sup>-1</sup> after 40 cycles. 75 This decrease in capacity is due to the dissolution of the material in the electrolyte. But its lithiated form does not show severe capacity fading. The authors suspected that this is because the lithium salts easily form a polymer network, which restricts the dissolution of the active material.

3.1.2.7 Salification. In this strategy, organic materials form organic salts by introducing high-polarity groups to enhance the cycling stability of the materials by limiting their dissolution in the aprotic electrolyte. Organic salts are mostly used in metalion batteries, such as Li2TP, a widely considered anode in LIBs. 10 Similarly, Na<sub>2</sub>TP and K<sub>2</sub>TP are used for SIBs and KIBs. In fact, the high polarity of -COOK in K2TP was responsible for the high capacity retention of 94.6% over 500 cycles at 1 A g $^{-1}$  with a coulombic efficiency of 100% (Fig. 9(d)). $^{76,77}$ 

**3.1.3 Anhydride.** Anhydride is another typical category of carbonyl compound cathode materials. Anhydride organic materials are promising candidates for alkali-ion batteries due to their reversible insertion and de-insertion capability of alkali ions on the conjugated carbonyl structure. We have summarized below various observations related to the prime issues of anhydride-based electrodes, such as rapid capacity decay, high solubility, *etc.* 

3.1.3.1 Capacity enhancement without tuning unsaturated bonds. As an organic semiconductor-based electrode material, perylene-3,4,9,10-tetracarboxylic dianhydride (PTCDA) has been investigated for both Na and Li-ion batteries. 45,78,79 Here we will discuss another strategy to enhance the capacity without affecting the unsaturated bond. Along with acetylene black, NaPTCDA yields a specific capacity (361 mA h g<sup>-1</sup>) higher than its theoretical capacity (273 mA h g<sup>-1</sup>).<sup>78</sup> This extra capacity-gain could suggest a similar mechanism to that observed in the case of Li and aromatic anodes, an additional reaction between the unsaturated carbon and alkali metal. Surprisingly, the same mechanism is not observed in NaPTCDA. The authors suggested that the extra capacity is attributed to the formation of a solid electrolyte interphase (SEI) layer resulting from the irreversible reaction with the electrolyte.78 Wang et al. showed that the PTCDA electrode could retain a reversible capacity of 250.5 mA h g<sup>-1</sup> and coulombic efficiency of 100% after 140 cycles, corresponding to a capacity retention ratio of 40.4%.<sup>78</sup> On the other hand, Wei et al. showed that PTCDA could adjust the amount of Na-insertion by fixing the discharge voltage as shown in Fig. 9. At the discharge voltage of 0.01 V, PTCDA could insert 15 Na ions and delivered a capacity of 1017 mA h g<sup>-1</sup>.45 Authors exhibited high electrochemical reversibility in forming  $Na_{2-x}PTCDA$  (0  $\leq x < 2$ ) only without any modification commercial PTCDA sub-micrometer rods.45 PTCDA is a prime candidate for the SIB anode compared to most SIB anodes, including hard carbon, alloys, and metal oxides or sulfides. But the cycling stability needs to improve further for it to be a highly promising practical anode material for SIBs.

3.1.3.2 3D hybrid structures for retention of capacity. Yuan et al. so suggested nanoengineered ultralight cathode materials based on a PTCDA and graphene aerogel composite, namely, the PGC40 composite cathode, where PTCDA molecules were confined in a 3D hybrid architecture. This composite cathode material involves a two-step redox reaction and delivers a specific capacity of 202 mA h g $^{-1}$  at 50 mA g $^{-1}$  and 64 mA h g $^{-1}$  at a high current density of 2000 mA g $^{-1}$ . Also, it still exhibited a reversible capacity of 78 mA h g $^{-1}$  after 500 cycles even at a high current density of 1000 mA g $^{-1}$  with a capacity retention of 66%, which is far better than the pristine PTCDA. These enhanced electrochemical properties of PGC40 come from its special 3D structure, which not only improves the conductivity of both electrons and Li-ions, but also effectively prohibits the dissolution of the active material into the electrolyte.

3.1.3.3 Increasing the molecular weight. Wang et al. 81 studied four anhydride organic compounds to reveal the essential reason for the rapid capacity attenuation. They found that the

reason was the loss of active material due to the formation of an easily dissolvable dilithium compound during the discharge reaction. Out of the four anhydride compounds of different molecular weights, the lowest molecular weight compound showed faster attenuation of the discharge capacity and poorer electrochemical cycle performance. This observation implies that the increasing molecular weight of the active material could reduce its dissolution in the electrolyte.

**3.1.4 Imide.** In the small molecule category, diimides are highly attractive cathode materials because of their relatively low solubility along with low cost and easy-to-functionalize nature. Problems with the diimides are decomposition, low voltage, and limited theoretical capacity. As we can see in Fig. 10, a maximum of two electrons can be reversibly accommodated by arylene diimide. More than two electrons can create electrostatic repulsion and steric hindrance, which can lead to decomposition.

3.1.4.1 Influence of the active ion. Disodium pyromellitic diimidate shows poor capacity retention with a steady decrease in capacity during cycling.<sup>83</sup> Interestingly, a similar effect is not observed in the case of its Li-homologue. Brandell and coworkers suggested that this decomposition occurs due to the highly concentrated negative charge on the molecules during cycling instead of the solubility of the active material in the electrolyte.<sup>84</sup> This kind of negative charge accumulation is not observed in the case of its Li-homologue as the high-affinity of Li<sup>+</sup> ions can shield and well compensate the negative charge.

3.1.4.2 Effect of substituents and functionalization. Different substituent groups are used to enhance the voltage of naphthalene diimides (NDIs). Vadehra and coworkers showed the effect of this strategy on NDIs with and without functionalization of diimide nitrogens with hexyl groups. <sup>85</sup> In both cases, they observed a change in voltage. Out of all considered substituents (Me<sub>2</sub>N, F, CN), cyanide could increase the voltage from 2.5 V to 2.9 V. But it was also reported that with hexyl functionalization NDIs lose the capacity rapidly due to dissolution. This work also reported that the performance of unsubstituted NDIs was poor due to an unfavorable crystal packing. The performance could be increased further by reducing the domain size of the active material and increasing the homogeneity. Bhosale et al. showed that the performance of hydrazine-treated benzoic acid-functionalized NDIs could be

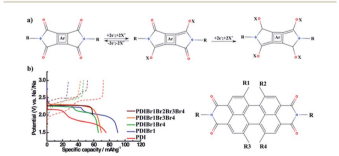


Fig. 10 (a) Reversible and irreversible processes in the diimide system, (b)charge—discharge voltage profile of PDI and its Br derivatives. The figures are reproduced from ref. 82 with permission. © WILEY-VCH.

improved compared to the untreated one.86 It was observed that hydrazine increases the conductivity and rate capabilities with 85 mA h  $g^{-1}$  specific capacity at 1C and 68 mA h  $g^{-1}$  at 10C with 88% capacity retention after 200 cycles at 5C. Therefore, this hydrazine treatment could be a useful strategy to improve the rate capabilities of other NDI analogues having high theoretical capacities.

3.1.4.3 Without excess mass. Cycling stability could be increased without the addition of excess mass in terms of substituents or functionalization. Chen et al. showed that designing a triangular-shaped NDI could lower the solubility and enhance the cycling stability with 60% retention of capacity after 300 cycles at 10C.87 Also, it delivers a specific capacity of 146.4 mA h  $g^{-1}$  at 0.1C and 58.1 mA h  $g^{-1}$  at 100C. Benefits of these triangular arrangements are fast diffusion of lithium-ion through the triangular channel (which ensures high rate performance), and high electronic conductivity due to the electronic coupling of the redox unit. The triangular-shaped NDI is slightly soluble in the neutral and reduced states, which could be improved by replacing NDI with the less soluble perylene diimide (PDI).

3.1.4.4 Molecular engineering to shape the discharge curve. A stable voltage output of organic molecules could be achieved by tuning the shape of the discharge curve by molecular engineering. Banda et al. discovered how the dihedral angles in PDI could affect the shape of the discharge curve.82 They increased the dihedral angles by increasing the number of Br substituents in the molecule. It was observed that (Fig. 10) PDI with one and two Br displays two discharge plateaus, whereas PDI with three and four Br shows one discharge plateau.

3.1.5 Non-conjugated polymer form. The above-discussed small molecule categories comprising the carbonyl group and its derivatives have plenty of advantages, e.g. easy functionalization and high capacity. Also, they have high solubility, which leads to low cycling stability. Without losing their favorable properties, the cycling stability could be improved by incorporating them into a polymer, wherein the solubility could be tuned efficiently to enhance the stability. In the following two sections, first we will discuss non-conjugated polymer and then conjugated polymer electrodes.

3.1.5.1 Suppression of dissolution by di-imide polymers. Arylene di-imide incorporated non-conjugated polymeric materials have been studied as cathodes. In an effort to design a flexible and free-standing binder-free cathode, Wu and co-workers performed an in situ polymerization of pyromellitic dianhydride with ethylenediamine and SWCNTs to form polyimide.88 This material exhibits good rate capabilities with a high  $C_{\rm sp}$  of 226 mA h  $g^{-1}$  at 0.1C and 120 mA h  $g^{-1}$  at 20C and retains 85% of its initial capacity after 200 cycles at 0.5C due to its insolubility in the electrolyte. Wang et al. studied a series of different non-conjugated arylene di-imide polymers with pyromellitic, NDI, and PDI cores with different lengths of diamine spacers.8 A decrease in  $C_{\rm sp}$  was observed with increase in the alkyl spacer length from propyl to butyl with the PDI core. Most importantly, it was reported that the cycling stability increases by increasing the size of the arylene core. The capacity fading issue, especially dissolution could be suppressed by using arylene di-imide

polymers with N-atoms. But the cycling stability still needs to improve to serve the practical purpose. An interesting example of this non-conjugated polymer class is a dendronized polymer. Kim and co-workers synthesized a dendronized polymer with anthraquinone groups pendant to the dendrons as a cathode.89 Although it exhibits high capacity retention (~90-95% after 100 cycles at 0.5C). But electrical conductivity is an issue that may be caused by large polymer domains that are electrically isolated from the conductive pathway of the carbon black particles.

The non-conjugated polymer form is also considered to improve the anode performance. For example, 1,4,5,8-naphthalenetetracarboxylic dianhydride (NTCDA)-derived polyimide has been proposed as the anode material for aqueous rechargeable lithium-ion or sodium-ion batteries.90 The polymeric nature undoubtedly aids in the cycling stability by preventing dissolution. In the case of a lithium electrolyte, it has a 160 mA h  $g^{-1}$   $C_{\rm sp}$  at 100 mA  $g^{-1}$  and as a full cell with a LiCoO<sub>2</sub> cathode, it delivers 71 mA h  $g^{-1}$   $C_{\rm sp}$  with an operating potential of 1.12 V. With a sodium electrolyte, the  $C_{\rm sp}$  is 165 mA h  ${\rm g}^{-1}$ , and in a full device with NaVPO4 as the cathode, the  $C_{\rm sp}$  is 40 mA h g<sup>-1</sup>. It is also observed that the capacity decay is faster in the NaVPO₄F electrode, ~30% capacity drop within 20 cycles.

In general, non-conjugated polymers are used less often as anode materials for metal-ion batteries. This is because the backbone is susceptible to decomposition at low voltages and therefore, they are mostly used as aqueous battery anodes.

**3.1.6 Conjugated polymer form.** In general, the inherent conductivity and redox activity of conjugated polymers are conducive for the fabrication of metal-ion battery electrodes with high rate performance and low amounts of inactive fillers, such as carbon additives and binders. However, the most common issue, the charge repulsion from delocalized polarons and bipolarons on the backbone, could be tackled by designing conjugated polymers with redox-active groups having localized charges and rapid-reversible electrochemistry. Overall, the polymerization could enhance the stability as shown below.

3.1.6.1 Polymerization with and without sulfur. Chloroanilic acid and dilithium chloranilate, when polymerized with sulfur, exhibit, respectively, 214 mA h  $g^{-1}$  and 247 mA h  $g^{-1}$   $C_{\rm sp}$  at 50 mA g<sup>-1</sup>, whereas without polymerization, they exhibit 119 mA h g<sup>-1</sup> and 193 mA h g<sup>-1</sup>  $C_{\rm sp}$  at 50 mA g<sup>-1</sup>, respectively.<sup>91</sup> It is also observed that the polymerization increases the cycling stability, e.g. the lithiated derivative retained 90% of the capacity after 1500 cycles at 500 mA g<sup>-1</sup>. In a follow-up study, Song and co-workers polymerized different isomers of anthraquinone with either sulfur or by condensation polymerization.92 Out of all synthesized polymers, they observed that the 1,4isomer of anthraquinone formed by condensation polymerization is most promising with 263 mA h  $\rm g^{-1}$   $C_{\rm sp}$  at 0.2C and 98.3% capacity retention after 100 cycles at 0.2C. Poly(anthraquinonyl sulfide) (PAQS) shows similar cycling stability and 98.4% capacity retention after 100 cycles at 0.2C. But the discharge capacity is low, 213.8 mA h  $\rm g^{-1}$   $C_{\rm sp}$  at 0.2C. They also reported 93 that poly(benzoquinonyl sulfide) as a sodium-ion battery cathode, has 268 mA h  $\mathrm{g}^{-1}$   $C_{\mathrm{sp}}$  at 50 mA  $\mathrm{g}^{-1}$  and 68% capacity retention after 100 cycles at 500 mA g<sup>-1</sup>. Therefore, these reports suggest that polymerization with sulfur is an attractive

strategy to design stable and high-performance organic cathodes.

3.1.6.2 Donor-acceptor copolymers. Liang and co-workers introduced donor-acceptor copolymers for use in lithium-ion battery electrodes. 94 Donor-acceptor copolymers such as poly {[N,N'-bis(2-octyldodecyl)-1,4,5,8-naphthalenedicarboximide-2,6-diyl]-alt-5,5'-(2,2'-bithiophene)} or (P(NDI2OD-T2)) and poly  $\{[N,N']$ -bis(2-octyldodecyl)-1,4,5,8-naphthalenedicarboximide-2,6-diyl]-*alt*-5,5'-[2,2'-(1,2-ethanediyl)bithiophene]} or (P(NDI2OD-TET)) have also been studied as ultrafast lithiumion battery cathodes. The authors observed 96% capacity retention after 3000 cycles for P(NDI2OD-T2). Low theoretical capacities of 54.2 and 52.7 mA h g<sup>-1</sup> were observed for P(NDI2OD-T2) and P(NDI2OD-TET), respectively. They also observed that interruption of polymer conjugation with a saturated ethylene linker lowers the rate capability. Therefore, the main drawback here is  $C_{\rm sp}$ , which could be enhanced by removing the solubilizing alkyl chains.

#### 3.2 Porous organic polymers

Organic electrode materials have attracted considerable interest mainly owing to their outstanding features, namely remarkable electrochemical capabilities, low costs as well as versatility of the geometrical structure. Notably, the conjugated molecules have been recently promoted to serve as prospective electrodes for upcoming advanced rechargeable batteries with high-performance electrochemical properties. PoPs, relying on rigid covalent lattices providing high stability with wide pores allowing easier ion transfer from the electrolyte, represent exciting prospects for rechargeable battery electrodes. PoPs, relying on rigid covalent lattices providing high stability with wide pores allowing easier ion transfer from the electrolyte, represent exciting prospects for rechargeable battery electrodes.

Furthermore, the functional POPs displayed enhanced Li/Na/K<sup>+</sup> conductivity with enormous prospects for serving solid-state electrolyte materials. In this section, we will discuss the recent developments in the application of porous organic polymers in various advanced batteries, including K-ion and Li–S batteries. Table 1 summarizes the electrochemical performances of select POP electrodes and gives a comparison with other electrodes.

We have already mentioned that the implementation of organic materials as cathode batteries has some advantages such as low-cost and chemically tuneable characteristics, but they still face some disadvantages, most notably, dissolution of the redox-active material in the electrolytes, suppression of the electrolyte-ion transport in the cathodes, as well as very lower conductivity. To date, POP materials as battery cathodes offer significant strengths as they feature well-ordered open pores for ion transport from the electrolyte, small volume expansion upon the insertion of Na<sup>+</sup>, as well as robust stability. Notwithstanding the fact that POPs provide several benefits as cathode materials in Li-ion rechargeable batteries, they mostly suffer from low redox potentials and low conductivity thus constraining their applicability in LIBs. 105

To tackle the above concerns, a variety of approaches have been investigated, such as the introduction of redox-active monomers, insertion of redox-active modules by means of post-synthetic functionalization of POP materials, boosting their conductivity behavior, and introducing various active materials to hybridize POP materials.

**3.2.1 Redox-active monomers to enhance the redox activity.** Initially, Wu *et al.*<sup>106</sup> successfully designed and synthesized a redox-active covalent organic framework-*graft*-polysulfide (COF-*graft*-PS), illustrated in Fig. 11(a), through the

Table 1 Selected electrochemical properties of some organic electrode materials for metal-ion and Li-S batteries<sup>a</sup>

Rechargeable battery type	Selected electrodes	$S_{ m BET}$ (m <sup>2</sup> g <sup>-</sup>	$^{1}$ ) Capacity (mA h g $^{-1}$	Current energy ) density	Long-term performance	Voltage (V)
Li-ion batteries	${\rm Li_2TP^{69}}$	_	200	0.1C	50 cycles	0.88
	Benzoic-NDI <sup>86</sup>	_	85	1C	200 cycles	2.9
	mp-COF-graft-PS <sup>106</sup>	14.8	425	$250 \ { m mA g^{-1}}$	500 cycles	1.7-3.0
	DTP-ANDI-COF@CNTs <sup>107</sup>	478	67	2.4C	700 cycles	1.5-3.5
	NT-COF <sup>99</sup>	1276	124	$10~\mathrm{mA~g^{-1}}$	100 cycles	2.7
	TFPB-COF <sup>99</sup>	458	968	$100 \text{ mA g}^{-1}$	300 cycles	0.005-3
	E-TFPB-COF/MnO <sub>2</sub> (ref.	345	1359	$100 \; { m mA g^{-1}}$	300 cycles	0.005-3
	100)			-	•	
	ADALS <sup>35</sup>	_	190	0.5C	100 cycles	0.61-1.3
	$PBQS^{93}$	_	275	$50 \; { m mA g^{-1}}$	1000 cycles	1.5-4.0
Na-ion batteries	SBDC <sup>73</sup>	_	260	50	400 cycles	0.18 - 2.5
	$\mathrm{SSDC}^{73}$	_	192	50	400 cycles	0.1 - 2.5
	PTCDA <sup>78</sup>	_	361	$25~\mathrm{mA~g}^{-1}$	140 cycles	0-3.0
	CTF-HUSTs <sup>109</sup>	764	467	$50 \text{ mA g}^{-1}$	500 cycles	0.05 - 2.0
	TFPB-TAPT COF <sup>108</sup>	120	246	$30 \text{ mA g}^{-1}$	500 cycles	0.001 - 3.0
	ANDASS <sup>141</sup>	_	170	0.2C	1000 cycles	0.5 - 2.5
	PBQS <sup>93</sup>	_	268	$50 \; { m mA g^{-1}}$	100 cycles	1.0-3.5
K-ion batteries	PyBT CMP <sup>121</sup>	493	428	$30 \text{ mA g}^{-1}$	500 cycles	0.1-3
	SHPNC <sup>122</sup>	693-1798	318	$25 \text{ mA g}^{-1}$	370 cycles	0.01-3
Li-sulfur batteries	CTF-1 (ref. 123)	789	848	0.2C	50 cycles	1.1-3.0
	Por-COF <sup>124</sup>	1095	633	0.5C	200 cycles	1.8-2.7

 $<sup>^{</sup>a}$  —:  $S_{\text{RET}}$  data are not available in the literature.

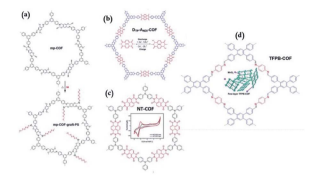


Fig. 11 (a) Chemical configuration of mp-COF & mp-COF-graft-PS structures; (b) DTP-ANDI-COF structure, inset: redox process of NDI; (c) NT-COF structure, inset: CV of NT-COF in the presence and absence of light; (d) TFPB-COF structure; inset: schematic-illustration of the chemical-exfoliation of COFs. Figures are reproduced from ref. 98-100 with permission. © 2018 WILEY-VCH, 2019 WILEY-VCH, 2020 WILEY-VCH.

functionalization of a conventional covalent organic framework (COF) as a new cathode material for high-energy rechargeable Li-ion batteries and demonstrated that it can yield a remarkable capacity of about 425 mA h g<sup>-1</sup> at a rate of 250 mA h g<sup>-1</sup> with a significantly higher rate capability. 106 Noticeably, COF-graft-PS-based Li-ion batteries exhibited excellent cycling stability up to 500 cycles, which was much higher compared to that of numerous Li- S batteries, which is derived from the high structural stability of mp-COF-graft-PS connected by carbonsulfur covalent bonds, as was reflected in the high coulombic performance of over 98% within the initial 10-cycles. The novel strategy proposed in this study to functionalize COFs provides a pathway towards the preparation of new robust cathode materials for sustainable and high-performance Li-ion batteries based on abundant and low-cost materials.

3.2.2 Addressing the low electrical conductivity. Despite the fact that COF-based electrode materials for LIBs have reached very high starting discharge capacities with robust cycling stability, their high rate discharge has been inhibited due to the very low electrical conductivity of the COF materials. For the purpose of improving the electrical conductivity of these materials, Xu et al.107 introduced another strategy in the quest for energy storage by developing and exploring the possibility of using a new redox-active, crystalline, mesoporous covalent organic framework (COF) on carbon nanotubes (Fig. 11(b)) as an electrode for LIBs. 107 DTP-ANDI-COF@CNTs as the cathode for Li-ion batteries showed excellent coulombic efficiency and high rate capability with a very stable cycle performance owing to the synergistic effect of both DTP-ANDI-COF and CNTs. Under such conditions, the DTP-ANDI-COF contributed to improving the redox activity, cycle performance, and ion transport, while the CNTs contributed by improving the electrical conductivity. 107

3.2.3 Effect of intra-molecular charge transfer in COFs. It is noteworthy that energy loss occurs inevitably when power sources are connected with energy storage systems. Recently, Jiangquan et al.99 have presented a COF integrating naphthalene diimide and triphenylamine units (NT-COF) shown in

Fig. 11(c) as moderate cathode materials together with Li-anode for direct solar to electrochemical energy conversion and storage. LIBs based on NT-COF as the cathode showed promising electrochemical performance, including an initial capacity of 124 mA h  $g^{-1}$  and a reversible capacity of about 100 mA h  $g^{-1}$ after the first 100 cycles. The improved discharge voltage profile resulted from the synergetic effect of intra-molecular charge transfer with a reversible electrochemical process. Additionally, during the discharge process, NT-COF as the cathode under irradiation produces the TPA+-NDI- complex, while in the charging process, the irradiation strengthens the conversion of the redox complex TPA+NDI- to TPA-NDI in the NT-COF cathode material. This study presents a series of samples that can be used to efficiently convert solar energy into electrochemical energy.99

3.2.4 Influence of 2D heterostructures. More recently, the Ajayan group reported a set of E-TFPB-COF and E-TFPB-COF/ MnO<sub>2</sub> heterostructures shown in Fig. 11(d) through a chemical exfoliation pathway as cathodes for LIBs. 100 They demonstrated that these new 2D heterostructure cathodes present moderate electrochemical properties, such as ion and electron kinetics, and provide a good redox-active site for improved reversible Listorage capacity, which leads to remarkable capacity with excellent cycling efficiency of about 1359 and 968 mA h  $g^{-1}$  after the first 300 cycles.100

On the other hand, Na-ion batteries based on COFs as battery electrodes have also been investigated in recent years. K. Wang et al. 109 have introduced a new strategy to construct covalent triazine frameworks (CTF-HUSTs) through the polycondensation method with many interesting properties such as layered structures with a wide inter-layer distance of about 3.9 Å, large surface areas, and adjustable functions. Particularly, CTF-HUST-4 has been revealed to be a prospective electrode material with an excellent discharge capacity of approximately 467 mA h  $g^{-1}$  for sodium-ion batteries. <sup>109</sup> Furthermore, Pradhan et al. have reported a new COF material made up of 1,3,5-tris-(4formylphenyl)benzene (TFPB) and 1,3,5 tris-(4-amino phenyl)triazine (TAPT) with  $C_3$ - $C_3$  symmetrical topology, <sup>108</sup> denoted as TFPB-TAPT COF, as a sustainable anode material for sodium-ion batteries. As shown in Fig. 12, the TFPB-TAPT COF revealed an outstanding electrochemical performance including a good reversible capacity of 246 mA h  $g^{-1}$ , excellent cycle stability up to 500 cycles and a very high Na+storage capability at several current rates owing to its wide-open ordered porous framework.108 The outcomes of this study suggest that the redox-active layered 2D TFPB-TAPT COF shows high promise not only as an anode for SIBs but also for diverse alkali metal batteries.

3.2.5 Role of conjugated microporous polymers for K-ion batteries. In the past few years, there has been a great deal of interest in potassium ion batteries (KIBs) as well, given both the resource accessibility and abundance of the potassium component, along with the fact that K<sup>+</sup>/K offers a somewhat similar redox profile (of 2.93 V versus a standard hydrogen electrode) relative to the redox potential of Li<sup>+</sup>/Li (3.04 V). 110-112 However, the significant volume modification induced upon the adsorption and desorption of K-ions typically leads to poor

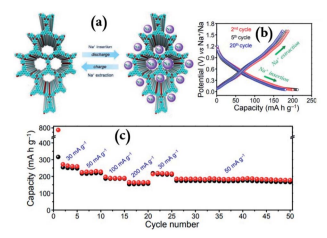


Fig. 12 The electrochemical efficiency of TFPB-TAPT COF organic anode materials in sodium-ion batteries. (a) The model for reversible Na<sup>+</sup>-storage upon charging, (b) charge and discharge process profiles for three different cycles, and (c) rate efficiency at various current rates. Figures are reproduced from ref. 108 with permission. © 2016 Royal Society of Chemistry.

cycling stability.<sup>113,114</sup> Accordingly, achieving viable electrode materials for K-ion batteries with both high-capacity and consistent cycling capability remains a big challenge.<sup>115,116</sup> Here we show conjugated microporous polymers (CMPs) as an example of the host electrode for K-ion batteries, which does not imply any limitation of the application of K-ion as an active alkali metal ion in other organic systems.

Conjugated microporous polymers (CMPs),  $^{117,118}$  which serve as a strong candidate for addressing the growing environmental and energy related concerns, have been extensively probed for various applications. As a consequence of the widespread  $\pi$ -conjugation throughout the polymer skeleton, outstanding physico-chemical stability, highly porous nature along with increased surface area, CMPs have emerged as prospective high-energy storage materials.  $^{119,120}$  Jiang  $et\ al.$  successfully synthesized two groups of CMPs with distinct configurations (Fig. 13(a and b)), and investigated their possible use as negative

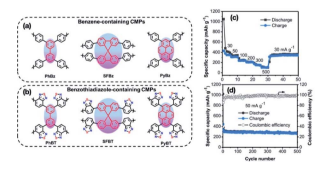


Fig. 13 (a and b) The polymer structures of both benzene-containing and benzothiadiazole-containing conjugated microporous polymers. (c) The electrochemical performance of PyBT at a current density between 30 and 500 mA  $\rm g^{-1}$ . And (d) the cycling stability/coulombic performance of PyBT at 50 mA  $\rm g^{-1}$ . The figures are reproduced from ref. 121 with permission. © 2019, American Chemical Society.

electrodes for K-ion batteries. <sup>121</sup> Some influencing factors such as the electronic structure and the lowest unoccupied molecular orbitals of the polymer skeleton have a great impact on the performance of the K-ion batteries. The de-localized LUMO not only enables remarkable redox activity due to a high degree of charge de-localization across the polymer skeleton, but also simultaneously decreases the charge density of redox-active sites. Consequently, K-ion batteries with PyBT-CMPs have shown increased charge de-localization, and small electronic bandgaps, and have achieved a large specific capacity of about 428 mA h g<sup>-1</sup> along with smooth cycling stability as depicted in Fig. 13(c and d). A strategy for effective conception of CMPs to serve as anodes of high-performance K-ion batteries has been presented within the context of this study. <sup>121</sup>

More recently, a new hierarchically porous nitrogen-doped carbon (SHPNC) has been synthesized via a cocoon-silk chemistry technique as a potential anode material for batteries by Luo and co-workers. Notably, both the hierarchically porous nature and the high n-doping within the SHPNC enable a high rate of ion as well as electron transport, serving a significant purpose in buffering K-ion volume changes, and further extending the electrochemically binding sites. Accordingly, the capacity *versus* cycling number at varying charge–discharge current rates have been probed within a potential voltage window of about 0.01–3.00 V *versus* K/K $^+$ . Additionally, as the current density drops to 25 mA g $^{-1}$ , roughly 93% of the discharge capacity can be recovered with a reversible specific capacity of about 318 mA h g $^{-1}$  achieved over 200 cycles, which indicates the high stability of SHPNC as an anode for KIBs.

3.2.6 COF as the host in Li–S batteries. Li–S batteries are still suffering from several limitations, namely the utilization of electron-insulating sulfur and  ${\rm Li_2S}$  molecule, together with the very low cycle stability resulting from the reduction of soluble lithium-polysulfide intermediates species  $({\rm Li_2S_x})$  during discharge processes. Cathode materials with plenty of nanopores could overcome these limitations. However, the development of many of these materials is frequently associated with many difficulties and multistep synthesis approaches. POPs could serve as host cathode materials for Li–S batteries (LiSBs), whose well-ordered nanopores and a large surface area are suitable for the adsorption of sulfur and lithium-polysulfide species.

Liao *et al.*<sup>123</sup> used COFs for the first time as host materials for S-containing cathodes to address the problem caused by the loss of soluble lithium polysulfide intermediates that occur in the discharge process.<sup>123</sup> The composite covalent triazine-based framework (CTF-1) was made up with the melt – diffusion technique. A high capacity of about 848 mA h g<sup>-1</sup> at a current density of 0.2C has been achieved. Nevertheless, a very small sulfur charge of 34% has been achieved in CTF-1. Furthermore, CTF as the host for the S-cathode shows capacity retention of 64% after 50 cycles at charge/discharge performance of 0.1C.<sup>123</sup> Subsequently, for the purpose of increasing the sulfur loading and cycle stability, they designed another type of COFs, a 2D porous porphyrin-based COF (Por-COF) with a comparatively large pore volume and small pore size distribution. Por-COF as a host-material for sulfur storage in LiSBs has demonstrated

a superior capacity of 670 mA h  $\rm g^{-1}$  at charge/discharge rates of 1.0C with high cycle stability. 124

#### 3.3 Organometallic polymers

Organometallic polymers having organic binders and transition metal ions exhibit bifunctional characteristics of both organic and metal components. They possess distinct electrochemical features such as high capacity and active metal sites of inorganic components. Moreover, most of these materials are not soluble in the electrolyte. Therefore, organo-metallic polymers could serve as potential electrodes for rechargeable batteries, mainly Li- and Na-ion batteries and high-performance Li-S batteries. At present, a wide range of organometallic polymers have been explored as battery electrodes, and they are mainly divided into following groups: metal-organic frameworks, porphyrins, phthalocyanines, ferrocene, and so on.125,126 MOFs consist of porous materials featuring a uniform structure made up of metal-ions or clusters, acting as joints, connected by multi-directional organic ligands, which serve as linkers in the lattice structure. They feature improved characteristics such as high porosity, wide surface area, and numerous metal-centers, therefore, they have been widely employed for electrochemical energy-based technologies. 126,127 Over the years since 2006, MOFs have been consistently presented128 and extensively explored for rechargeable battery applications. In 2013, Satoshi et al. reported the ionic conductivity and transport of MOF materials for batteries and fuel-cells.129 Recently, Pang et al. reviewed MOF-based materials as a potential candidate in energy storage technologies with fascinating electrochemical performances for metal-ion and advanced Li-S batteries. 130 Wang and co-workers have suggested a new strategy to use transition-metal-based MOFs and their derivatives as potential anode materials for Li-ion batteries due to their adjustable structure, large surface area, and highly porous structure. 131 The transition metal in MOFs is susceptible to inducing many different valence variations in redox reactions, thereby leading to increased capacities. Furthermore, MOFs could serve as host materials for sulfur-cathodes by virtue of their tuneable porosity. Also, MOFs as battery electrodes have the ability to sieve the ions which is advantageous for uniform deposition of Li-ion thereby preventing dendrite growth in lithium metal anodes. The porphyrin complexes involve the bonding of metal ions in the middle together with the four nitrogen atoms in the hetero-cycles. In this process, the metal-ions act as electronacceptors while the nitrogen atoms act as electron-donors and thus display redox activity132 as shown in Fig. 14. Within alkali metal-ion batteries, the porphyrin is subjected to a transformation from anti-aromatic into aromatic during the redox processes. 132 Likewise, phthalocyanines consist of a macrocyclic structure with up to eight N-atoms and eight C-atoms. The metal-ions and N-atoms in the middle can absorb lithium-polysulfide, thereby inhibiting the shuttling effect in Li-S batteries. Additionally, ferrocene compounds consist of dicyclopentadiene and ferrous-ions (hereinafter referred to as Fe<sup>2+</sup>). <sup>133</sup> Accordingly, Fe<sup>2+</sup> can be oxidized and inversely reduced in metal-ion batteries. The ferrocene in Li-S batteries can also

14
$$\pi$$
 aromatic

NiNC<sup>2+</sup>

NiNC<sup>2+</sup>

NiNC<sup>2+</sup>

16 $\pi$  antiaromatic

+ e<sup>-</sup>
- e<sup>-</sup>

NiNC

NiNC

18 $\pi$  aromatic

NiNC

Fig. 14 Redox process of antiaromatic dimesitylnorcorrole nickel(II) complex (NiNC) with 16 pi electrons. Reproduced with permission from ref. 132. © 2014, Wiley-VCH.

act as a containment medium for the lithium-polysulfide species owing to its outstanding flexibility, reversibility, and fast diffusion.<sup>133</sup>

Metal-organic frameworks (MOFs), consisting of either metal ions or clusters linked to an organic binder via coordinate bonds, have gained considerable attention in the field of electrochemical energy storage based on their high-porosity structure and pseudo-capacitive capabilities upon the inclusion of redox-active metal cores. 134,135 The MOFs have been extensively implemented for electrochemical energy storage, either as MOFs or MOF-derived carbonaceous materials. 136 Notwithstanding recent progress, 135,136 their applicability in metal-ion and Li-S batteries has been limited due to the low electrical conductivity and poor stability of pristine MOFs. 137-139 Therefore, further investigation and characterization of highly stable MOFs as electrode materials with both high-power density and increased energy density, as well as improved cycling stability and enhanced electrical conductivity for convenient uses, is essential, and it remains a major challenge.

#### 3.4 Azo based salts

Organic battery electrode materials show promise for sustainable and eco-friendly secondary batteries owing to the lightweight, low-cost, durability, and recyclability of organic-based materials. Nevertheless, conventional organic-based electrodes are negatively affected by their low cycling stability and limited power density. In recent years, a new organic-chemical material in which the azo-group (-N=N-) is part of the molecular structure was widely investigated as an organic battery electrode for large-scale electrochemical storage applications. Luo *et al.* reported a class of organic anode materials that contain azo-functional groups, namely azobenzene-4,4'-dicarboxylic acid

lithium salt (ADALS) having an azo-group in the middle of the conjugated structure for lithium and sodium-ion batteries. ADALS has achieved a high capacity of about 190 mA h g $^{-1}$  at an energy density of 0.5C with up to 90%, 71%, and 56% capacity retention as the current density is raised to 2, 10, and 20C, respectively. Additionally, ADALS retains about 89% of its starting capacity even after 5000 cycles at a current energy density of 20C with a very low capacity decay factor of 0.0023% by cycle, which represents the highest performance among all organic-based anode materials.  $^{\rm 35}$ 

Azo compounds as a family of organic anode materials have demonstrated potential performance for Na-ion batteries.140 Luo et al. have also reported an organic anode material based on azobenzene-4,4'-dicarboxylic acid sodium salt (ADASS) as a new strategy to improve the electrochemical performance of Na-ion batteries.141 ADASS organic anode showed a good reversible capacity of about 170 mA h g<sup>-1</sup> at current energy density of 0.2C and retains about 66% and 58% reversible capacities as the current density increases to 10C (1000 cycles) and 20C (2000 cycles), respectively, showing outstanding high-rate capability and long cycle life. Furthermore, it maintains a reversible capacity of 98 mA h  $g^{-1}$  for 2000cycles at 20C and achieves a low capacity decay rate of 0.0067% by cycle. This is attributed to the presence of a carboxyl group which prevents the azo-benzene from dissolving in the liquid electrolyte.141 Based on energy density, high rate capability and cycle-life, the azo-group materials as battery electrodes surpass many other organic electrodes available for Na-ion batteries, exclusively those based on carbonyl-groups (C=O), including Na-terephthalate77,142 and benzo-quinone derivatives,143 as well as imine compounds (C=N), namely Schiff's base derivatives. 144

#### 3.5 Organic radical polymers

Recently, great attention has been paid to organic radical polymers (ORPs), which have led to several outstanding advances in different areas, including electrochemical energy storage applications. The synthesis techniques currently employed for their development have been widely surveyed. 128,140,145-147 The majority of ORPs are made up of two main components: a robust radical pendant and a backbone polymer. In order to be suitable for use in rechargeable batteries, ORPs require long-term cycling stability, insolubility, and affinity in the electrolyte, as well as very high-energy-density.

Firstly, the long-term cycling stability of ORP cathodes in full cells remains to be fully explored to evaluate their practical applicability. In this context, one of the most important and challenging criteria to meet in molecular conception consists of the fact that the presence of unpaired electrons in the polymer have to be sufficiently non-reactive in the electrolyte. The representative structures of the radical counterparts are the alicyclic-nitroxyls (such as TEMPO and PROXYL). For instance, the unpaired electrons of the TEMPO framework are capable of sustaining for over a year within an aprotic medium which contains 1.0 M of LiPF<sub>6</sub>. <sup>148</sup>

In addition, the geometrical structure of the backbone polymer in the ORPs has an impact on the insolubility and affinity in the electrolyte. In order to avoid contact of the radical polymer with the other electrode, the polymer requires to be insoluble in the electrolyte and to have good affinity for the electrolyte in order to ensure good ionic conductivity. So far, a wide range of backbone polymers, including polymethacrylate, 149 polystyrene, 150 polynorbornene, 151,152 polyether, 153 polyacetylene, 154 and cellulose, 155 have been identified and widely reviewed. Also, the polymer to be employed should be dense in the thick electrode in order to provide a high enough capacity. From the perspective of processability and structural stability, polymethacrylate and polyether have proven to be the highly desirable backbone for ORPs.

Furthermore, the energy density of a rechargeable battery can be determined by the product of the voltage profile and the specific capacity. The voltage profile of ORPs is mainly related to redox reactions. RPs undergo two types of redox reactions, namely the p-type reaction between a radical and a cation and the n-type reaction between a radical and an anion. He conventional Li-TMO reaction, the potential voltage of the p-type reaction typically tends to be greater than that of the n-type reaction. Hence, employing a p-type reaction is preferable to achieve high energy density.

## 4. Future outlook: multivalent alkali ion and dissolution issue

Despite the various scientific achievements, there is still a long way to go before realizing the large-scale applications of organic electrode materials in metal-ion batteries. As discussed above, the implementation of organic electrode materials in rechargeable batteries has been expected to reduce the energystorage price as well as provide long-term durability. 128 More specifically, multi-valent metal-organic batteries have also recently attracted significant interest owing to their large availability as well as the high volumetric energy density of the multi-valent metals, such as Al, Mg, and Ca.157-159 Besides their geo-political abundance over Li-atom, they also have higher volumetric-energy density by virtue of their multi electronic redox features. 160,161 Furthermore, the electrodes with multivalent metals show lower susceptibility to dendrite-growth, 162 and this is due to the significantly shorter diffusion profile of multi-valent metals. 163 Relative to other active monovalent metals, the multi-valent metals are less reactive in air, thereby substantially reducing the manufacturing cost and could provide better battery stability-safety.164,165 Considering the availability of both organic electrode materials and multi-valent metals, the metal-organic batteries could stand as a sustainable and affordable high-performance rechargeable battery, suitable for large-scale electrochemical energy storage systems.

Despite the fact that over  $\sim 30$  years, these organic batteries have been studied, still significant interest in the practical implementations of organic electrode materials is hindered mainly owing to their low electrical/ionic conductivity, their

dissolution into the electrolyte, their small potential-voltage and their limited specific capacity. For this purpose, several alternative techniques have been derived for improving the electrochemical performance of organic electrode materials in rechargeable batteries through the structural design, polymerization, and amorphization of organic electrode materials. Such approaches were shown to dramatically boost the performance of organic electrode materials in multi-valent metal-organic batteries. 166-168 In addition to the above-discussed strategies, this review prescribes the following guidelines, which could help not only to understand but also to overcome the potential drawbacks of organic batteries of monovalent as well as multivalent alkali ions:

Firstly, the combination of computational design and successful synthesis of high-efficiency organic electrode materials having a high-operating potential voltage with extended lifetime capability remains a primary concern. Ideal organic electrodes require a moderate specific capacity of more than 200 mA h g<sup>-1</sup>, a wide operating potential voltage of over 2.0 V, and an extended lifetime of greater than 1000 cycles in order to be suitable for use in large-scale energy-storage systems. 169 A prominent approach involves reducing the molecular weight of the active materials as well as increasing the concentration of redox active sites in the organic electrode surface. Furthermore, the synthesis approaches that are carried out shall be ecofriendly and highly scalable and cost-effective. From the existing literature overview, the highest mean performance for organic electrode materials is apparently exhibited by the carbonyl-based group materials owing to their strongly reversible multi-electron redox features, 170 although lowering their solubility as well as increasing their structural strength needs to be addressed properly. Polymerization and formation of alkali ion salts could be effective in improving the cycling performance of organic cathodes. Increase of molecular weight could lower the dissolution, but at the same time it actually decreases the theoretical capacity of the cathode material. Therefore, future research should focus not only on designing the organic cathode materials with lower equivalent molecular weight, but also on developing suitable solid electrolytes with high ionic conductivity and low solubility of organic cathode materials. Furthermore, organic cathode materials possess lower redox potentials with a lower stability window compared to conventional inorganic cathodes, which again lowers the energy density. Here, structural modification and molecular functionalization of organic cathode materials could be performed to further improve their redox potential and provide a competitive energy density. It is noteworthy that recent advancements in theoretical modeling and materials informatics such as machine learning, artificial intelligence (AI) kernel and data mining can not only trigger and narrow down the highthroughput search process more effectively but also suggest potential candidates for synthesis.

Secondly, a deeper insight into the electrochemical storage behavior of organic electrode materials in multi-valent metalorganic batteries needs to be provided. The operation principle of organic materials and electrodes in metal-organic batteries is much more challenging as compared to that in LIBs. For

instance, the stored ions on the electrodes (cathode: AlCl<sub>4</sub>-, AlCl<sup>2+</sup> and anode: Al<sup>3+</sup>) can be different in the multi-valent metal-organic batteries with chlorine containing electrolytes compared to typical inorganic systems for aluminum batteries.171 Both the structural flexibility and poor intermolecular interactions provided by organic based materials ensure the reversible diffusion and storage of bulky aluminum carrier ions without undergoing significant structural modifications, as in the case of majority of inorganic-based materials.172,173 In addition, organic electrodes typically sustain a storage process through "ionic-coordination" thereby significantly contrasting with the slow kinetics provided by certain inorganic-based compounds resulting from their severe coulombic interactions with Al-carrier ions. 174 An exciting development over the past few years has been the rise of a broad class of organic materials for high-performance Al-organic batteries. 175-178 Notably, quinones are remarkable for their increased specific capacity, enhanced redox potential and multielectron reactivity. Apart from the para- and/or orthoquinone, the electronic integration of the contiguous carbonyl groups leads to the creation of negative enolates, which can be conjugated with Al ions to maintain the structure. 179,180 The aim of the most recent quinone studies was to increase the number of carbonyl groups within the framework.181 However, studies have revealed that the 2 or 3 adjacent carbonyl groups could be chelated by only a single bulky Al-carrier ion, thus restricting the efficient application of the active sites as well as hindering their electrochemical performance. 182 In particular, the nitrogen-containing hetero-aromatic materials are potentially implicated in the reversible redox reactions with Al-carrier ions, although they exhibit a comparatively limited working potential with a poor battery efficiency.<sup>183</sup> Regardless of all these efforts, so far there are no available reports regarding the development of organic cathode materials with more than one synergistic active core for Al-organic based batteries, as well as exploring the related redox-reactions is still not fully accomplished. Within this framework, heterocyclic conjugated polymer nanoarchitectonics which entails molecular engineering of active organic molecules with both C=O and/or C=N redox active sites along with  $\pi$ -conjugation within a framework has the potential184 to yield a fundamentally novel perspective for designing and developing new innovative electrode-based materials for Al-organic batteries along with enhanced capacitance, long-term cyclability, and superior energy density. Consequently, further investigation efforts are expected to shift towards more sophisticated operando approaches for better insight into the structural modifications of organic electrodes in multi-valent metal-organic batteries.

Thirdly, addition of large amounts (~30-60 wt%) of conductive carbon is considered to balance the low electrical conductivity of organic electrode materials. Addition of a large amount of conductive carbon also lowers the energy density. Therefore, designing organic electrodes with less than 30 wt% of conductive carbon is one of the bottlenecks for successful realization of organic batteries. At the same time, it is also important to consider as future research focus on the thermal

stability of the organic materials, specially the electrode consists of small organic molecules.

In short, research activities of organic cathode materials should focus on the high mass loadings, solid electrolyte design, minimal electrolyte usage, higher potential window, and easy and affordable large-scale production for their commercialization.

#### 5. Conclusions

In summary, this review provides an overview of the most recently explored organic electrode materials for promising mono- and multi-valent alkali-ion based organic batteries with reference to the materials chemistry and the storage mechanisms of alkali metal-ions. Organic electrode materials have attracted considerable focus from the rechargeable battery research community over the last 10–15 years, as evidenced by the wide range of publications on the subject mirrored in this review.

Capabilities of organic batteries could frequently exceed the ones achieved using inorganic materials. One feature that is perhaps less debated about with regard to organic materials is that they possess a smaller specific gravity range from 1.5 to 2.2 g cm<sup>-3</sup> relative to approximately 4.5 g cm<sup>-3</sup> in the case of LiCoO<sub>2</sub>. Substantial use of conductive additives in the organic batteries could lower the energy density, whereas covalent bonding in the organic molecules could also provide stability. So, there are various facts that could either enhance or suppress the efficiency of organic batteries. This review article discussed possible strategies such as fusing multiple carbonyl groups, introducing nitrogen-containing heterocyclics, crystalline nanostructures, tuning of the aromaticity, electrode processing methods, extension of conjugation, degree of deprotonation, choice of counterion, enhancing the molecular weight, the effect of substituents and functionalization, molecular engineering, polymerization, etc. to address the critical issues related to organic electrodes, i.e. dissolution, capacity fading, lower energy density, and electrical conductivity. To conclude, we believe that organic electrode materials still have suitable physical and chemical properties to be implemented in largescale high-efficiency rechargeable batteries, if we can overcome the aforementioned problems. Here, we have critically addressed each strategic descriptor to show its effect on the successful realization of organic batteries. It is expected that this study will bring more insight into this area and will refuel the worldwide attention on sustainable energy storage studies, especially to move from academia to practical applications in the foreseeable future.

#### Conflicts of interest

There are no conflicts to declare.

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#### References

- 1 R. B. Araujo, A. Banerjee, P. Panigrahi, L. Yang, M. Strømme, M. Sjödin, C. M. Araujo and R. Ahuja, *J. Mater. Chem. A*, 2017, 5, 4430–4454.
- 2 S. Lee, G. Kwon, K. Ku, K. Yoon, S. K. Jung, H. D. Lim and K. Kang, *Adv. Mater.*, 2018, **30**(42), 1704682.
- 3 Y. Lu and J. Chen, Nat. Rev. Chem., 2020, 4(3), 127-142.
- 4 J. B. Goodenough and Y. Kim, *Chem. Mater.*, 2010, 22(3), 587-603.
- 5 J. R. Owen, T. Le Gall, K. H. Reiman and M. C. Grossel, *J. Power Sources*, 2003, **119**, 316–320.
- 6 H. Chen, M. Armand, G. Demailly, F. Dolhem, P. Poizot and J. M. Tarascon, *ChemSusChem*, 2008, 1(4), 348–355.
- 7 M. Armand and J. M. Tarascon, *Nature*, 2008, **451**(7179), 652–657.
- 8 H. G. Wang, S. Yuan, D. L. Ma, X. L. Huang, F. L. Meng and X. B. Zhang, Adv. Energy Mater., 2014, 4(15), 1400554.
- 9 P. G. Bruce, S. A. Freunberger, L. J. Hardwick and J.-M. Tarascon, *Nat. Mater.*, 2011, 11, 19–29.
- 10 M. Armand, S. Grugeon, H. Vezin, S. Laruelle, P. Ribière, P. Poizot and J. M. Tarascon, *Nat. Mater.*, 2009, **8**, 120–125.
- 11 K. Nakahara, J. Iriyama, S. Iwasa, M. Suguro, M. Satoh and E. J. Cairns, *J. Power Sources*, 2007, **165**(1), 398–402.
- 12 D. L. Williams, J. J. Byrne and J. S. Driscoll, *J. Electrochem. Soc.*, 1969, **116**(1), 2.
- 13 P. Novák, K. Müller, K. S. V. Santhanam and O. Haas, *Chem. Rev.*, 1997, **97**(1), 207–282.
- 14 D. MacInnes, M. A. Druy, P. J. Nigrey, D. P. Nairns, A. G. MacDiarmid and A. J. Heeger, *J. Chem. Soc., Chem. Commun.*, 1981, (7), 317–319.
- 15 H. Shirakawa, E. J. Louis, A. G. MacDiarmid, C. K. Chiang and A. J. Heeger, *J. Chem. Soc., Chem. Commun.*, 1977, (16), 578–580.
- 16 X. Jia, Y. Ge, L. Shao, C. Wang and G. G. Wallace, *ACS Sustainable Chem. Eng.*, 2019, 7(17), 14321–14340.
- 17 R. B. Araujo, A. Banerjee, P. Panigrahi, L. Yang, M. Sjödin, M. Strømme, C. M. Araujo and R. Ahuja, *Phys. Chem. Chem. Phys.*, 2017, 19(4), 3307–3314.
- 18 T. Matsunaga, H. Daifuku, T. Nakajima and T. Kawagoe, *Polym. Adv. Technol.*, 1990, 1(1), 33–39.
- 19 S. J. Visco and L. C. DeJonghe, *J. Electrochem. Soc.*, 1988, 135(12), 2905.
- 20 D. Y. Wang, W. Guo and Y. Fu, *Acc. Chem. Res.*, 2019, 52(8), 2290–2300
- 21 S. Tobishima, J. Yamaki and A. Yamaji, *J. Electrochem. Soc.*, 1984, 131(1), 57.

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- 22 K. Nakahara, S. Iwasa, M. Satoh, Y. Morioka, J. Iriyama, M. Suguro and E. Hasegawa, Chem. Phys. Lett., 2002, 359(5-6), 351-354.
- 23 T. Suga, Y. J. Pu, K. Oyaizu and H. Nishide, Bull. Chem. Soc. Ipn., 2004, 77(12), 2203-2204.
- 24 M. Liu, S. J. Visco and L. C. De Jonghe, J. Electrochem. Soc., 1990, 137(3), 750.
- 25 R. Van Noorden, *Nature*, 2014, **507**(7490), 26–28.
- 26 T. B. Schon, B. T. McAllister, P. F. Li and D. S. Seferos, Chem. Soc. Rev., 2016, 45(22), 6345-6404.
- 27 R. B. Araujo, A. Banerjee and R. Ahuja, J. Phys. Chem. C, 2017, 121(26), 14027-14036.
- 28 S. Muench, A. Wild, C. Friebe, B. Häupler, T. Janoschka and U. S. Schubert, Chem. Rev., 2016, 116(16), 9438-9484.
- 29 J. Kim, J. H. Kim and K. Ariga, *Joule*, 2017, 1(4), 739-768.
- 30 Z. Song and H. Zhou, Energy Environ. Sci., 2013, 6(8), 2280-
- 31 Y. Lu, X. Hou, L. Miao, L. Li, R. Shi, L. Liu and J. Chen, Angew. Chem., 2019, 131(21), 7094-7098.
- 32 A. Ponrouch, J. Bitenc, R. Dominko, N. Lindahl, P. Johansson and M. R. Palacin, Energy Storage Mater., 2019, 20, 253-262.
- 33 T. Matsunaga, T. Kubota, T. Sugimoto and M. Satoh, Chem. Lett., 2011, 40(7), 750-752.
- 34 X. Han, G. Qing, J. Sun and T. Sun, Angew. Chem., Int. Ed., 2012, 51(21), 5147-5151.
- 35 C. Luo, O. Borodin, X. Ji, S. Hou, K. J. Gaskell, X. Fan, J. Chen, T. Deng, R. Wang, J. Jiang and C. Wang, Proc. Natl. Acad. Sci. U. S. A., 2018, 115(9), 2004-2009.
- 36 J. Meng, H. Guo, C. Niu, Y. Zhao, L. Xu, Q. Li and L. Mai, *Joule*, 2017, **1**(3), 522–547.
- 37 A. Vizintin, J. Bitenc, A. Kopač Lautar, K. Pirnat, Grdadolnik, J. Stare, A. Randon-Vitanova and R. Dominko, Nat. Commun., 2018, 9(1), 1-7.
- 38 Y. Liang and Y. Yao, Joule, 2018, 2(9), 1690-1706.
- 39 M. Lee, J. Hong, J. Lopez, Y. Sun, D. Feng, K. Lim, W. C. Chueh, M. F. Toney, Y. Cui and Z. Bao, Nat. Energy, 2017, 2(11), 861-868.
- 40 X. Wu, S. Jin, Z. Zhang, L. Jiang, L. Mu, Y. S. Hu, H. Li, X. Chen, M. Armand, L. Chen and X. Huang, Sci. Adv., 2015, **1**(8), e1500330.
- 41 H. Kim, D. H. Seo, G. Yoon, W. A. Goddard, Y. S. Lee, W. S. Yoon and K. Kang, J. Phys. Chem. Lett., 2014, 5(17), 3086-3092.
- 42 I. A. Rodríguez-Pérez, Y. Yuan, C. Bommier, X. Wang, L. Ma, D. P. Leonard, M. M. Lerner, R. G. Carter, T. Wu, P. A. Greaney, J. Lu and X. Ji, J. Am. Chem. Soc., 2017, **139**(37), 13031–13037.
- 43 Y. Chen, W. Luo, M. Carter, L. Zhou, J. Dai, K. Fu, S. Lacey, T. Li, J. Wan, X. Han, Y. Bao and L. Hu, Nano Energy, 2015, 18, 205-211.
- 44 X. Han, C. Chang, L. Yuan, T. Sun and J. Sun, Adv. Mater., 2007, **19**(12), 1616–1621.
- 45 W. Luo, M. Allen, V. Raju and X. Ji, Adv. Energy Mater., 2014, 4(15), 1400554.

- 46 A. J. Wain, G. G. Wildgoose, C. G. R. Heald, L. Jiang, T. G. J. Jones and R. G. Compton, J. Phys. Chem. B, 2005, **109**(9), 3971-3978.
- 47 S. Renault, V. A. Oltean, C. M. Araujo, A. Grigoriev, K. Edström and D. Brandell, Chem. Mater., 2016, 28(6), 1920-1926.
- 48 J. S. Foos, S. M. Erker and L. M. Rembetsy, J. Electrochem. Soc., 1986, 133(4), 836.
- 49 B. L. Funt and P. M. Hoang, in Electrochemical Society Extended Abstracts, 1984, vol. 84-1.
- 50 T. Boschi, R. Pappa, G. Pistoia and M. Tocci, J. Electroanal. Chem., 1984, 176(1-2), 235-242.
- 51 T. Ohzuku, H. Wakamatsu, Z. Takehara and S. Yoshizawa, Electrochim. Acta, 1979, 24(6), 723-726.
- 52 H. Alt, H. Binder, A. Köhling and G. Sandstede, Electrochim. Acta, 1972, 17(5), 873-887.
- 53 B. Häupler, A. Wild and U. S. Schubert, Adv. Energy Mater., 2015, 5, 1402034.
- 54 T. B. Reddy, Linden's Handbook of Batteries, McGraw-Hill, New York, 4th edn, 2011.
- 55 Q. Zou, W. Wang, A. Wang, Z. Yu and K. Yuan, Mater. Lett., 2014, 117, 290-293.
- 56 M. Lee, J. Hong, H. Kim, H. D. Lim, S. B. Cho, K. Kang and C. B. Park, Adv. Mater., 2014, 26(16), 2558-2565.
- 57 H. Li, W. Duan, Q. Zhao, F. Cheng, J. Liang and J. Chen, Inorg. Chem. Front., 2014, 1(2), 193-199.
- 58 H. C. Lin, C. C. Li and J. T. Lee, J. Power Sources, 2011, **196**(19), 8098-8103.
- 59 W. Wan, H. Lee, X. Yu, C. Wang, K. W. Nam, X. Q. Yang and H. Zhou, RSC Adv., 2014, 4(38), 19878-19882.
- 60 T. Yokoji, H. Matsubara and M. Satoh, J. Mater. Chem. A, 2014, 2, 19347-19354.
- 61 H. Kim, J. E. Kwon, B. Lee, J. Hong, M. Lee, S. Y. Park and K. Kang, Chem. Mater., 2015, 27, 7258-7264.
- 62 A. Iordache, V. Maurel, J. M. Mouesca, J. Pécaut, L. Dubois and T. Gutel, J. Power Sources, 2014, 267, 553-559.
- 63 A. Shimizu, Y. Tsujii, H. Kuramoto, T. Nokami, Y. Inatomi, N. Hojo and J. I. Yoshida, Energy Technol., 2014, 2, 155-158.
- 64 C. Luo, R. Huang, R. Kevorkyants, M. Pavanello, H. He and C. Wang, Nano Lett., 2014, 14(3), 1596-1602.
- 65 D. Wu, Z. Xie, Z. Zhou, P. Shen and Z. Chen, J. Mater. Chem. A, 2015, 3, 19137-19143.
- 66 H. Zhang, Q. Deng, A. Zhou, X. Liu and J. Li, J. Mater. Chem. A, 2014, 2(16), 5696-5702.
- 67 S. Gottis, A. L. Barres, F. Dolhem and P. Poizot, ACS Appl. Mater. Interfaces, 2014, 6(14), 10870-10876.
- 68 H. H. Lee, Y. Park, K. H. Shin, K. T. Lee and S. Y. Hong, ACS Appl. Mater. Interfaces, 2014, 6(21), 19118-19126.
- 69 L. Fédèle, F. Sauvage, J. Bois, J.-M. Tarascon and M. Bécuwe, J. Electrochem. Soc., 2014, 161(1), A46.
- 70 L. Fédèle, F. Sauvage and M. Bécuwe, J. Mater. Chem. A, 2014, 2(43), 18225-18228.
- 71 T. Yasuda and N. Ogihara, Chem. Commun., 2014, 50(78), 11565-11567.
- 72 N. Ogihara, T. Yasuda, Y. Kishida, T. Ohsuna, K. Miyamoto and N. Ohba, Angew. Chem., 2014, 126(43), 11651-11656.

- 73 C. Wang, Y. Xu, Y. Fang, M. Zhou, L. Liang, S. Singh, H. Zhao, A. Schober and Y. Lei, *J. Am. Chem. Soc.*, 2015, 137(8), 3124–3130.
- 74 A. Choi, Y. K. Kim, T. K. Kim, M. S. Kwon, K. T. Lee and H. R. Moon, *J. Mater. Chem. A*, 2014, 2, 14986–14993.
- 75 V. A. Mihali, S. Renault, L. Nyholm and D. Brandell, RSC Adv., 2014, 4, 38004–38011.
- 76 K. Lei, F. Li, C. Mu, J. Wang, Q. Zhao, C. Chen and J. Chen, *Energy Environ. Sci.*, 2017, **10**(2), 552–557.
- 77 Y. Park, D. S. Shin, S. H. Woo, N. S. Choi, K. H. Shin, S. M. Oh, K. T. Lee and S. Y. Hong, *Adv. Mater.*, 2012, 24(26), 3562–3567.
- 78 H. G. Wang, S. Yuan, Z. Si and X. B. Zhang, Energy Environ. Sci., 2015, 8, 3160–3165.
- 79 M. Veerababu, U. V. Varadaraju and R. Kothandaraman, *Int. J. Hydrogen Energy*, 2015, **40**, 14925–14931.
- 80 C. Yuan, Q. Wu, Q. Li, Q. Duan, Y. Li and H. G. Wang, *ACS Sustainable Chem. Eng.*, 2018, **6**, 8392–8399.
- 81 J. Wang, X. Wang, H. Li, X. Yang and Y. Zhang, J. Electroanal. Chem., 2016, 773, 22-26.
- 82 H. Banda, D. Damien, K. Nagarajan, A. Raj, M. Hariharan and M. M. Shaijumon, *Adv. Energy Mater.*, 2017, 7, 1701316.
- 83 R. Rajagopalan, Y. Tang, C. Jia, X. Ji and H. Wang, *Energy Environ. Sci.*, 2020, **13**, 1568–1592.
- 84 S. Renault, V. A. Mihali, K. Edström and D. Brandell, *Electrochem. Commun.*, 2014, 45, 52–55.
- G. S. Vadehra, R. P. Maloney, M. A. Garcia-Garibay and
   B. Dunn, *Chem. Mater.*, 2014, 26, 7151–7157.
- 86 M. E. Bhosale and K. Krishnamoorthy, *Chem. Mater.*, 2015, 27, 2121–2126.
- 87 D. Chen, A.-J. Avestro, Z. Chen, J. Sun, S. Wang, M. Xiao, Z. Erno, M. M. Algaradah, M. S. Nassar, K. Amine, Y. Meng, J. Fraser Stoddart, D. Chen, A. Avestro, J. Sun, Z. Erno, Z. Chen, K. Amine, S. Wang, M. Xiao, Y. Meng, M. M. Algaradah and M. S. Nassar, *Adv. Mater.*, 2015, 27, 2907–2912.
- 88 H. Wu, S. A. Shevlin, Q. Meng, W. Guo, Y. Meng, K. Lu, Z. Wei and Z. Guo, *Adv. Mater.*, 2014, **26**, 3338–3343.
- 89 J. Kim, J. Kim, J. Lee, H. K. Song and C. Yang, *ChemElectroChem*, 2014, 1(10), 1618–1622.
- 90 H. Qin, Z. P. Song, H. Zhan and Y. H. Zhou, *J. Power Sources*, 2014, **249**, 367–372.
- 91 Z. Song, Y. Qian, X. Liu, T. Zhang, Y. Zhu, H. Yu, M. Otani and H. Zhou, *Energy Environ. Sci.*, 2014, 7(12), 4077–4086.
- 92 Z. Song, Y. Qian, M. L. Gordin, D. Tang, T. Xu, M. Otani, H. Zhan, H. Zhou and D. Wang, *Angew. Chem., Int. Ed.*, 2015, 54, 13947–13951.
- 93 Z. Song, Y. Qian, T. Zhang, M. Otani and H. Zhou, *Adv. Sci.*, 2015, 2(9), 1500124.
- 94 Y. Liang, Z. Chen, Y. Jing, Y. Rong, A. Facchetti and Y. Yao, J. Am. Chem. Soc., 2015, 137, 4956–4959.
- 95 B. Dunn, H. Kamath and J. M. Tarascon, *Science*, 2011, 334(6058), 928-935.
- 96 Y. Xu, S. Jin, H. Xu, A. Nagai and D. Jiang, *Chem. Soc. Rev.*, 2013, 42(20), 8012–8031.
- 97 Z. Cheng, H. Pan, H. Zhong, Z. Xiao, X. Li and R. Wang, *Adv. Funct. Mater.*, 2018, **28**(38), 1707597.

- 98 X. Liu, C. F. Liu, W. Y. Lai and W. Huang, Adv. Mater. Technol., 2020, 5, 2000154.
- 99 J. Lv, Y. X. Tan, J. Xie, R. Yang, M. Yu, S. Sun, M. De Li, D. Yuan and Y. Wang, *Angew. Chem., Int. Ed.*, 2018, 57, 12716–12720
- 100 X. Chen, Y. Li, L. Wang, Y. Xu, A. Nie, Q. Li, F. Wu, W. Sun, X. Zhang, R. Vajtai, P. M. Ajayan, L. Chen and Y. Wang, Adv. Mater., 2019, 31(29), 1901640.
- 101 B. Kang and G. Ceder, Nature, 2009, 458(7235), 190-193.
- 102 P. S. Herle, B. Ellis, N. Coombs and L. F. Nazar, *Nat. Mater.*, 2004, 3(3), 147–152.
- 103 X. Zhan, Z. Chen and Q. Zhang, J. Mater. Chem. A, 2017, 5(28), 14463-14479.
- 104 E. Jin, M. Asada, Q. Xu, S. Dalapati, M. A. Addicoat, M. A. Brady, H. Xu, T. Nakamura, T. Heine, Q. Chen and D. Jiang, Science, 2017, 357(6352), 673–676.
- 105 G. Zhou, E. Paek, G. S. Hwang and A. Manthiram, *Nat. Commun.*, 2015, **6**(1), 1–11.
- 106 Y. Wu, Z. Zhang, S. Bandow and K. Awaga, *Bull. Chem. Soc. Jpn.*, 2017, **90**(12), 1382–1387.
- 107 F. Xu, S. Jin, H. Zhong, D. Wu, X. Yang, X. Chen, H. Wei, R. Fu and D. Jiang, *Sci. Rep.*, 2015, 5(1), 1–6.
- 108 B. C. Patra, S. K. Das, A. Ghosh, A. K. Raj, P. Moitra, M. Addicoat, S. Mitra, A. Bhaumik, S. Bhattacharya and A. Pradhan, J. Mater. Chem. A, 2018, 6(34), 16655–16663.
- 109 K. Wang, L. M. Yang, X. Wang, L. Guo, G. Cheng, C. Zhang, S. Jin, B. Tan and A. Cooper, *Angew. Chem., Int. Ed.*, 2017, 56(45), 14149–14153.
- 110 L. Fan, Y. Hu, A. M. Rao, J. Zhou, Z. Hou, C. Wang and B. Lu, *Small Methods*, 2021, 5, 2101131.
- 111 S. Liu, L. Kang and S. C. Jun, Adv. Mater., 2021, 33, 2004689.
- 112 J. Ge, L. Fan, A. M. Rao, J. Zhou and B. Lu, *Nat. Sustain.*, 2021, 5(3), 225–234.
- 113 J. Y. Hwang, S. T. Myung and Y. K. Sun, *Adv. Funct. Mater.*, 2018, **28**, 1802938.
- 114 C. Han, K. Han, X. Wang, C. Wang, Q. Li, J. Meng, X. Xu, Q. He, W. Luo, L. Wu and L. Mai, *Nanoscale*, 2018, 10, 6820–6826.
- 115 X. Lu, D. Zhang, J. Zhong, L. Wang, L. Jiang, Q. Liu, G. Shao, D. Fu, J. Teng and W. Yang, *Chem. Eng. J.*, 2022, 432, 134416
- 116 Y. Meng, C. Nie, W. Guo, D. Liu, Y. Chen, Z. Ju and Q. Zhuang, *Mater. Today Energy*, 2022, **25**, 100982.
- 117 H. Ma, Y. Chen, X. Li and B. Li, *Adv. Funct. Mater.*, 2021, **31**, 2101861.
- 118 K. Amin, N. Ashraf, L. Mao, C. F. J. Faul and Z. Wei, *Nano Energy*, 2021, **85**, 105958.
- 119 M. Gamal Mohamed, T. Hassan Mansoure, M. Mohamed Samy, Y. Takashi, A. A. K. Mohammed, T. Ahamad, S. M. Alshehri, J. Kim, B. M. Matsagar, K. C.-W. Wu and S.-W. Kuo, *Mol*, 2022, 27, 2025.
- 120 S. Luo, Z. Zeng, H. Wang, W. Xiong, B. Song, C. Zhou, A. Duan, X. Tan, Q. He, G. Zeng, Z. Liu and R. Xiao, *Prog. Polym. Sci.*, 2021, 115, 101374.
- 121 C. Zhang, Y. Qiao, P. Xiong, W. Ma, P. Bai, X. Wang, Q. Li, J. Zhao, Y. Xu, Y. Chen, J. H. Zeng, F. Wang, Y. Xu and J. X. Jiang, ACS Nano, 2019, 13, 745–754.

- 122 H. Luo, M. Chen, J. Cao, M. Zhang, S. Tan, L. Wang, J. Zhong, H. Deng, J. Zhu and B. Lu, Nano-Micro Lett., 2020, 12, 1-13.
- 123 H. Liao, H. Ding, B. Li, X. Ai and C. Wang, J. Mater. Chem. A, 2014, 2(23), 8854-8858.
- 124 H. Liao, H. Wang, H. Ding, X. Meng, H. Xu, B. Wang, X. Ai and C. Wang, J. Mater. Chem. A, 2016, 4(19), 7416-7421.
- 125 H. Zhang, Y. Gao, X. H. Liu, Z. Yang, X. X. He, L. Li, Y. Qiao, W. H. Chen, R. H. Zeng, Y. Wang and S. L. Chou, Adv. Funct. Mater., 2022, 32, 2107718.
- 126 D. Y. Wang, R. Liu, W. Guo, G. Li and Y. Fu, Coord. Chem. Rev., 2021, 429, 213650.
- 127 X. F. Lu, Y. Fang, D. Luan and X. W. D. Lou, Nano Lett., 2021, 21(4), 1555-1565.
- 128 F. Xie, W. Hu, L. Ding, K. Tian, Z. Wu and L. Li, Polym. Chem., 2017, 8, 6106-6111.
- 129 S. Horike, D. Umeyama and S. Kitagawa, Acc. Chem. Res., 2013, 46(11), 2376-2384.
- 130 M. Du, Q. Li, Y. Zhao, C. Sen Liu and H. Pang, Coord. Chem. Rev., 2020, 416, 213341.
- 131 T. Li, Y. Bai, Y. Wang, H. Xu and H. Jin, Coord. Chem. Rev., 2020, 410, 213221.
- 132 J. Y. Shin, T. Yamada, H. Yoshikawa, K. Awaga and H. Shinokubo, Angew. Chem., Int. Ed., 2014, 53, 3096-3101.
- 133 J. L. Qin, B. Q. Li, J. Q. Huang, L. Kong, X. Chen, H. J. Peng, J. Xie, R. Liu and Q. Zhang, Mater. Chem. Front., 2019, 3, 615-619.
- 134 R. Zhao, Z. Liang, R. Zou and Q. Xu, Joule, 2018, 2, 2235-
- 135 T. Sun, J. Xie, W. Guo, D. S. Li and Q. Zhang, Adv. Energy Mater., 2020, 10, 1904199.
- 136 X. Li, S. Zheng, L. Jin, Y. Li, P. Geng, H. Xue, H. Pang and Q. Xu, Adv. Energy Mater., 2018, 8, 1800716.
- 137 W. Zhu, A. Li, Z. Wang, J. Yang and Y. Xu, Small, 2021, 17, 2006424.
- 138 S. Bai, X. Liu, K. Zhu, S. Wu and H. Zhou, Nat. Energy, 2016, **1**, 1–6.
- 139 Y. Zheng, S. Zheng, H. Xue and H. Pang, J. Mater. Chem. A, 2019, 7, 3469-3491.
- 140 K. A. Hansen and J. P. Blinco, *Polym. Chem.*, 2018, 9(13), 1479-1516.
- 141 C. Luo, G. L. Xu, X. Ji, S. Hou, L. Chen, F. Wang, J. Jiang, Z. Chen, Y. Ren, K. Amine and C. Wang, Angew. Chem., *Int. Ed.*, 2018, 57(11), 2879–2883.
- 142 L. Zhao, J. Zhao, Y. S. Hu, H. Li, Z. Zhou, M. Armand and L. Chen, Adv. Energy Mater., 2012, 2(8), 962–965.
- 143 C. Luo, J. Wang, X. Fan, Y. Zhu, F. Han, L. Suo and C. Wang, Nano Energy, 2015, 13, 537-545.
- 144 M. López-Herraiz, E. Castillo-Martínez, J. Carretero-González, J. Carrasco, T. Rojo and M. Armand, Energy Environ. Sci., 2015, 8(11), 3233-3241.
- 145 H. Roghani-Mamaqani, V. Haddadi-Asl and M. Salami-Kalajahi, *Polym. Rev.*, 2012, **52**(2), 142–188.
- 146 C. Hua, A. Rawal, T. B. Faust, P. D. Southon, R. Babarao, J. M. Hook and D. M. D'Alessandro, J. Mater. Chem. A, 2014, 2(31), 12466-12474.

- 147 Y. Xie, K. Zhang, Y. Yamauchi, K. Oyaizu and Z. Jia, Mater. Horiz., 2021, 8(3), 803-829.
- 148 K. Nakahara, K. Oyaizu and H. Nishide, Chem. Lett., 2011, 40, 222-227.
- 149 H. Nishide, S. Iwasa, Y. J. Pu, T. Suga, K. Nakahara and M. Satoh, *Electrochim. Acta*, 2004, **50**(2–3), 827–831.
- 150 T. Suga, S. Sugita, H. Ohshiro, K. Oyaizu and H. Nishide, Adv. Mater., 2011, 23(6), 751-754.
- 151 J. Qu, T. Katsumata, M. Satoh, J. Wada and T. Masuda, Polymer, 2009, 50(2), 391-396.
- 152 T. Suga, H. Konishi and H. Nishide, Chem. Commun., 2007, (17), 1730-1732.
- 153 K. Oyaizu, T. Suga, K. Yoshimura and H. Nishide, Macromolecules, 2008, 41(18), 6646-6652.
- 154 K. Ovaizu, T. Sukegawa and H. Nishide, Chem. Lett., 2011, 40(2), 184-185.
- 155 J. Qu, R. Morita, M. Satoh, J. Wada, F. Terakura, K. Mizoguchi, N. Ogata and T. Masuda, Chem.-Eur. J., 2008, 14(11), 3250-3259.
- 156 H. Hishide and T. Suga, Electrochem. Soc. Interface, 2005, **14**, 32–36.
- 157 C. Li, C. C. Hou, L. Chen, S. Kaskel and Q. Xu, EnergyChem, 2021, 3, 100049.
- 158 R. Cang, Y. Song, K. Ye, K. Zhu, J. Yan, J. Yin, G. Wang and D. Cao, J. Electroanal. Chem., 2020, 861, 113967.
- 159 N. Khossossi, D. Singh, A. Banerjee, W. Luo, I. Essaoudi, A. Ainane and R. Ahuja, ACS Appl. Energy Mater., 2021, 4, 7900-7910.
- 160 J. Chen, Q. Zhu, L. Jiang, R. Liu, Y. Yang, M. Tang, J. Wang, H. Wang and L. Guo, Angew. Chem., Int. Ed., 2021, 60, 5794-5799.
- 161 F. Wu, H. Yang, Y. Bai and C. Wu, Adv. Mater., 2019, 31, 1806510.
- 162 D. Y. Wang, C. Y. Wei, M. C. Lin, C. J. Pan, H. L. Chou, H. A. Chen, M. Gong, Y. Wu, C. Yuan, M. Angell, Y. J. Hsieh, Y. H. Chen, C. Y. Wen, C. W. Chen, B. J. Hwang, C. C. Chen and H. Dai, Nat. Commun., 2017, 8(1), 1-7.
- 163 M. Jäckle, K. Helmbrecht, M. Smits, D. Stottmeister and A. Groß, Energy Environ. Sci., 2018, 11, 3400-3407.
- 164 K. Zhang, K. O. Kirlikovali, J. M. Suh, J. W. Choi, H. W. Jang, R. S. Varma, O. K. Farha and M. Shokouhimehr, ACS Appl. Energy Mater., 2020, 3, 6019-6035.
- 165 T. T. Wei, P. Peng, S. Y. Qi, Y. R. Zhu and T. F. Yi, J. Energy Chem., 2021, 57, 169-188.
- 166 B. Pan, J. Huang, Z. Feng, L. Zeng, M. He, L. Zhang, J. T. Vaughey, M. J. Bedzyk, P. Fenter, Z. Zhang, A. K. Burrell and C. Liao, Adv. Energy Mater., 2016, 6, 1600140.
- 167 Y. Wang, Z. Liu, C. Wang, Y. Hu, H. Lin, W. Kong, J. Ma and Z. Jin, Energy Storage Mater., 2020, 26, 494-502.
- 168 D. Lu, H. Liu, T. Huang, Z. Xu, L. Ma, P. Yang, P. Qiang, F. Zhang and D. Wu, J. Mater. Chem. A, 2018, 6, 17297-17302.
- 169 M. Mao, C. Luo, T. P. Pollard, S. Hou, T. Gao, X. Fan, C. Cui, J. Yue, Y. Tong, G. Yang, T. Deng, M. Zhang, J. Ma, L. Suo,

- O. Borodin and C. Wang, *Angew. Chem.*, 2019, **131**, 17984–17990.
- 170 B. Häupler, A. Wild and U. S. Schubert, *Adv. Energy Mater.*, 2015, 5(11), 1402034.
- 171 H. Yang, H. Li, J. Li, Z. Sun, K. He, H. M. Cheng and F. Li, *Angew. Chem., Int. Ed.*, 2019, **58**, 11978–11996.
- 172 J. Zou, K. Fan, Y. Chen, W. Hu and C. Wang, *Coord. Chem. Rev.*, 2022, **458**, 214431.
- 173 C. Wang, Energy Environ. Mater., 2020, 3, 441-452.
- 174 K. Qin, J. Huang, K. Holguin and C. Luo, *Energy Environ. Sci.*, 2020, **13**, 3950–3992.
- 175 X. Peng, Y. Xie, A. Baktash, J. Tang, T. Lin, X. Huang, Y. Hu, Z. Jia, D. J. Searles, Y. Yamauchi, L. Wang and B. Luo, *Angew. Chem., Int. Ed.*, 2022, e202203646.
- 176 D. J. Kim, D. J. Yoo, M. T. Otley, A. Prokofjevs, C. Pezzato, M. Owczarek, S. J. Lee, J. W. Choi and J. F. Stoddart, *Nat. Energy*, 2018, 4, 51–59.
- 177 J. Chen, Q. Zhu, L. Jiang, R. Liu, Y. Yang, M. Tang, J. Wang, H. Wang and L. Guo, *Angew. Chem., Int. Ed.*, 2021, **60**, 5794– 5799.

- 178 F. Guo, Z. Huang, M. Wang, W. L. Song, A. Lv, X. Han, J. Tu and S. Jiao, *Energy Storage Mater.*, 2020, 33, 250–257.
- 179 J. Zhou, X. Yu, J. Zhou and B. Lu, *Energy Storage Mater.*, 2020, **31**, 58–63.
- 180 J. Bitenc, N. Lindahl, A. Vizintin, M. E. Abdelhamid, R. Dominko and P. Johansson, *Energy Storage Mater.*, 2020, 24, 379–383.
- 181 Y. Li, L. Liu, Y. Lu, R. Shi, Y. Ma, Z. Yan, K. Zhang and J. Chen, *Adv. Funct. Mater.*, 2021, **31**, 2102063.
- 182 Y. T. Kao, S. B. Patil, C. Y. An, S. K. Huang, J. C. Lin, T. S. Lee, Y. C. Lee, H. L. Chou, C. W. Chen, Y. J. Chang, Y. H. Lai and D. Y. Wang, ACS Appl. Mater. Interfaces, 2020, 12, 25853–25860.
- 183 M. Mao, C. Luo, T. P. Pollard, S. Hou, T. Gao, X. Fan, C. Cui, J. Yue, Y. Tong, G. Yang, T. Deng, M. Zhang, J. Ma, L. Suo, O. Borodin and C. Wang, *Angew. Chem., Int. Ed.*, 2019, 58, 17820–17826.
- 184 J. Kim, J. H. Kim and K. Ariga, Joule, 2017, 1, 739-768.