Recent advances in cathode engineering to enable reversible room-temperature aluminium–sulfur batteries

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The rigorous requirements, such as high abundance, cost-effectiveness, and increased storage capacities, pose severe challenges to the existing Li-ion batteries' long-term sustainability. Room-temperature aluminum–sulfur (Al–S) chemistry, in particular, is gaining importance due to its high theoretical energy density (1700 W h kg$^{-1}$). Al–S battery technology is one of the emerging metal–sulfur candidates that can surpass current Li-ion chemistries. When coupled with sulfur, aluminum metal brings a cheap and energy-rich option to existing battery technologies. Owing to the unique virtues of the Al–S battery, it has garnered increasing interest among scientific communities. Al–S chemistry has been investigated for quite some time, yet the cell performance remained in its infancy, which poses a challenge to this technology's viability. Besides stabilizing the Al metal anode, the most important challenge in the practical development of Al–S batteries is the development of a suitable sulfur cathode material. Owing to the complexity of this multivalent system, numerous factors have been taken into account, but the best sulfur cathode is yet to be identified. A detailed exploration of sulfur cathodes and their implications on the battery performance are discussed in this mini-review article. We present a detailed picture of cathode materials that may serve as the reference guide for developing more practical cathode materials. Also, fundamental principles and challenges encountered in the development of the sulfur cathodes are highlighted. Through the knowledge disseminated in this mini-review, the development in the multivalent post-Li-ion battery can be accelerated. A glimpse of the future outlook on the Al–S battery system with different potential solutions is also discussed.

1. Introduction

In the era of rapid technological development, the demand for portable energy has grown multi-fold in recent years. Among various known energy storage devices, electrochemical energy storage through Li-ion batteries has become increasingly popular. However, conventional Li-ion batteries are gradually reaching their performance plateau due to the limited capacity of electrode materials, relatively low energy density, and high cost. Nobel laureate Prof. Goodenough highlights the need for a new battery technology that can safely perform tasks, and offer competitive prices and performance with respect to the well-established fossil fuel technologies. To meet the high energy demand, alkaline metal-based batteries (for instance, lithium–sulfur (Li–S)) attracted enormous attention recently due to its high theoretical energy density of about 2510 W h kg$^{-1}$. Despite the high performance, environmental-friendliness, and low cost of sulfur, the commercialization of alkaline-metal-based batteries is plagued seriously by various intrinsic issues, which include the limited specific capacity, low utilization of electrode materials, limited cycling stability, limited rate-performance, and safety concerns due to the flammable electrolyte and metal dendrites. Rechargeable batteries with high energy density, environmental-friendliness, and cost-effectiveness are in demand to power the modern society.

Unlike the Li or Na metal, aluminum (Al) is relatively stable in various aqueous or non-aqueous liquid media. Al is at least 100 times less expensive than Li metal, and its high natural abundance accounts for its cost-effectiveness. By weight, Al has a storage capacity of 2980 mA h g$^{-1}$, which is lower than that of Li (3860 mA h g$^{-1}$). However, due to the higher density of Al, its volumetric storage capacity is substantially greater than the Li metal anode. Besides being the most abundant metallic element on earth and having a safe and environmental benignancy, it has a theoretical gravimetric capacity of 2980 mA h g$^{-1}$, which is very close to that of lithium. Besides, Al has a negative redox potential, i.e., $\sim -1.7$ V vs. SHE, which is more positive than other metals, viz., lithium and sodium, making it relatively safer than those of other alkaline metal anodes in rechargeable batteries. In the construction of a typical Al–S battery, an aluminum foil with an ultra-smooth surface serves as an anode, and the elemental sulfur mixed with conductive
fillers acts as a cathode. The choice of electrolytes can also aid the kinetics of the reaction in a battery. Al has a tendency to form a passivation layer that increases its inertness towards electrolytes. Generally, stripping/plating in an Al–S battery can be enhanced with efficiencies greater than 98% using a room temperature non-aqueous electrolyte. Accordingly, non-aqueous ionic liquid (imidazolium salts + aluminium chloride) electrolytes have been utilized due to their high reaction kinetics towards the Al metal surface. Chloroaluminate ionic liquids are known to be the first generation ionic liquids’, owing to their high ionic conductivity and low volatility. Recently, various other aqueous electrolytes have also been investigated, which provide a safer and cost-effective platform for the development of an Al–S battery. As an illustration, the anode and cathode electrodes are separated by an [EMIM]Cl/AlCl$_3$-type ionic liquid and glass-fiber separator (Fig. 1). During discharge, the following reactions are expected to take place at the anode and cathode, respectively.

$$2\text{Al} + 14\text{AlCl}_4^- \rightarrow 8\text{Al}_2\text{Cl}_7^- + 6e^- \quad (1)$$

$$8\text{Al}_2\text{Cl}_7^- + 6e^- + 3\text{S} \rightarrow \text{Al}_2\text{S}_3 + 14\text{AlCl}_4^- \quad (2)$$

Upon discharge, the electrolyte dissociates to generate an Al$_2$Cl$_7^-$ species that move towards the sulfur cathode. It then receives electrons to release AlCl$_4^-$ anions in the electrolyte. The kinetics of these reactions is extremely slow due to the tendency of the Al anode to develop a thin passivation layer that limits its reversibility over a while (a few tens of cycles). Room-temperature ionic liquids are demonstrated to promote electrochemical reversibility in Al–S batteries. However, the [EMIM] Cl/AlCl$_3$-type ionic liquid electrolytes are moisture-sensitive and highly corrosive. Hence, they provide additional impediments to the development of a safe and cost-effective Al–S battery.

Given the advantages offered by Al, Al batteries have attracted increasing attention. In combination with the Al metal anode, various cathode materials have been examined to realize a high-energy and highly stable battery system. Some examples include graphite, vanadium oxides (VO$_2$, $V_2O_5$ (ref. 41)), titanium dioxide, and conductive polymers. Although the abovementioned cathode materials have brought obvious advantages, certain disadvantages have been brought by the traditional insertion-type cathode compounds. For instance, the energy density of such batteries fades quickly due to depletion in the electrolyte concentration. Contrary to the traditional liquid electrolytes, ionic liquids are identified to enhance the Al-ion batteries’ cycling stability, but at the cost of slow reaction kinetics. This is inevitable due to the substantial size of the chloroaluminate species $[(\text{AlCl}_3)_2]$ in ionic liquids.

2. Sulfur cathode

Besides the conventional intercalation-type cathodes, sulfur, as an emerging conversion-type cathode coupled with Al, provides prospects to tackle these challenges with respect to the intercalation reactions. Sulfur is abundant in nature, cost-effective, and environmentally benign. Owing to the extraordinarily high theoretical specific (1675 mA h g$^{-1}$) and volumetric (3459 mA h cm$^{-3}$) capacities, sulfur as an electrode material provides intriguing prospects in enhancing the energy density, as well as maintains the electrolyte concentration constant when used in Al–S batteries. However, the Al–S batteries’ limitations, such as the slow charge transfer kinetics, sluggish reversible reactions, and low compatibility with the electrolyte, remained an obstacle for the commercialization of this chemistry. To address these challenges, we provide a comprehensive insight into sulfur cathodes. This review provides comprehensive studies on the intrinsic and underlying principles of the sulfur cathode and its composites for Al–S batteries. Other cathode materials are also covered in brief to shed light on their basic properties. Through the knowledge disseminated in this mini-review, the development in the multivalent battery can be accelerated. It may pave the way for a practical solution for an efficient Al–S battery. Fig. 2 depicts a detailed framework.
of the progress made in cathode materials for reversible Al–S batteries over time.

2.1. Elemental sulfur/carbon composite for Al–S batteries

With the inception of the interest in Al–S batteries, Licht and co-workers investigated the concentrated polysulfide catholyte and an alkaline aluminium anode for possible application in Al–S batteries. 46 However, the energy densities were about 170 W h kg\(^{-1}\) based on dry materials (excluding water) and 110 W h kg\(^{-1}\) for the total battery materials with an open-circuit voltage of 1.3 V, which is comparable to that of Li-ion batteries. Likewise, Peramunage et al. 52 studied a novel Al–S battery that consists of a concentrated polysulfide catholyte and an alkaline aluminum anode. Among various catholyte solutions, low polarization losses and high storage capacity at the thin-film of CoS were observed for the cathodic polysulfide half-cell of the Al–S battery. An open circuit voltage of up to 1.3 V and specific energy of 110 W h kg\(^{-1}\) was achieved for the complete Al–S cell (the theoretical open-circuit voltage of the cell is about ~1.8 V, and theoretical specific energy based on potassium salts is about ~647 W h kg\(^{-1}\)).

The Al–S battery system is a promising platform for realizing high energy density batteries with a scale-up potential with the minimum requirement of the infrastructure, unlike alkali metal–sulfur batteries that require a sophisticated infrastructure for large-scale production. Many efforts have been made to manipulate the battery chemistry to advance this technology. For instance, Cohn et al. explored a primary non-aqueous Al–S battery with chloroaluminate ionic liquid as the electrolyte (EMICI/AlCl\(_3\)). 31 Owing to the high surface area, Ketjen black carbonaceous material was used as the sulfur host for the cathode. This was attributed to the homogeneous dispersion of sulfur particles and strong adhesion with the current collector. The cathode showed a remarkably high specific capacity of more than 1400 mA h g\(^{-1}\) (S) at 30 mA g\(^{-1}\) (more than 80% of the theoretical capacity of sulfur, i.e., 1675 mA h g\(^{-1}\)). However, the capacity of the cell was noticed to fade sharply due to the dissolution of the discharged products, which will affect the rechargeability of the cell. As a result, a relatively low coulombic efficiency was recorded. The specific energy density of about 1700 W h kg\(^{-1}\) was estimated based on the first discharge.

Therefore, it is apparent that due to the low charge transfer kinetics, the design and demonstration of a reversible Al–S battery are highly challenging. Gao et al. 50 was among the first few to repost a reversible Al–S battery in the presence of an ionic liquid as an electrolyte. Besides the inclusion of an ionic liquid in the cell, the design of the sulfur cathode was also altered. For instance, a microporous carbon coating was employed to block the dissolution of the discharge product. The specific loading of S was estimated to be about 58% in the cathode. The highest specific capacity of about 1230 mA h g\(^{-1}\) was obtained at a high current density of 50 mA g\(^{-1}\). Due to the nearly perfect encapsulation of sulfur, an improved reversibility of Al–S could be demonstrated. The microporous carbon with pore size <2 nm was identified to be favorable, leading to an enlarged interfacial area to accommodate the active species, 45,55 thus improving the electron access by reducing the diffusion length for Al\(^{3+}\). As depicted in Fig. 3a, a remarkably high specific capacity of about 1000 mA h g\(^{-1}\) (S) was achieved, which leads to a high energy density of 650 W h kg\(^{-1}\). However, the cell could survive only for 20 charges/discharge cycles (Fig. 3b). One of the possible causes of the poor reversibility of the electrochemical reactions in Al–S batteries could lie in the dissociation reaction from Al\(_2\)Cl\(_6\) to free Al\(^{3+}\). Yang et al. replaced all Al\(_2\)Cl\(_6\) anions with Al\(_2\)Cl\(_6\)Br\(^-\) to alleviate the kinetics of the electrochemical reaction. Owing to the ease of dissociation of the Al\(_2\)Cl\(_6\)Br\(^-\) anions, a high sulfur utilization of about 82% could be obtained using NBMPBr/AlCl\(_3\) as the electrolyte. To understand this improvement, DFT calculations were performed, where they evaluated and compared the stability for both Al\(_2\)Cl\(_6\)Br\(^-\) and Al\(_2\)Cl\(_7\)\(^-\) anions. The higher formation energy for Al\(_2\)Cl\(_6\)Br\(^-\) showed a higher stability of these anions. In addition, a smaller LUMO–HOMO gap for Al\(_2\)Cl\(_6\)Br\(^-\) was found to be accountable for its higher dissociation kinetics (at least 15 times) compared to Al\(_2\)Cl\(_7\)\(^-\). This resulted in an initial discharge capacity as high as 1300 mA h g\(^{-1}\), which could maintain over 400 mA h g\(^{-1}\) even after 20 cycles. 56 Although non-aqueous electrolytes promote reversibility better than that of aqueous electrolytes, they invite several inevitable issues, such as corrosion and environmental stability. To harness the aqueous electrolytes’ benefits, while preserving the reversibility of the redox reactions, water-in-salt electrolytes (such as [Al][Al(OTf)\(_3\)] + LiTFSI + HCl/[S/C]) were utilized to demonstrate a rechargeable Al–S battery. 24 They identified that the passivation of the Al anode plays a key role in achieving long-term stability.

The sulfur cathode, which comprises ZIF-67 as the host for sulfur, exhibited an initial specific capacity of 1410 mA h g\(^{-1}\) (for sulfur loading of 0.2 mg cm\(^{-2}\) only). It retained a reversible capacity of about 420 mA h g\(^{-1}\) after 30 cycles at a high current density of 200 mA g\(^{-1}\), and the coulombic efficiency was recorded to be about 97% (Fig. 3c). In addition to the improved
reversibility, the cell voltage could be extended to 3.0 V vs. Al/Al$^{3+}$ in the presence of water-in-salt electrolytes. Moreover, a trace amount (0.02 M) of HCl additive in the water-in-salt electrolyte inhibits the hydrolysis of AlS$_x$, and leads to promoting the smooth stripping/plating of the Al metal (Fig. 3f).

Guo and co-workers further improved the stability and reversibility of the charge/discharge reactions in the Al–S battery. They fabricated a metal–organic framework (MOF) derived microporous carbon decorated with Cu nanoparticles (HKUST-1-C) as a possible host for sulfur particles (Fig. 3g). The presence of Cu nanoparticles in the host improves the cathode’s performance by eliminating the possibility of polysulﬁde dissolution. As a result, a high initial discharge capacity of about 1200 mA h g$^{-1}$ could be achieved at a current density of 1000 mA g$^{-1}$. The S@HKUST-1-C retained a reversible capacity of about 600 mA h g$^{-1}$ and 460 mA h g$^{-1}$ after the 75th and 500th cycles, and the coulombic efficiency was about 90% and 95%, respectively. It has been shown that the presence of the Cu nanoparticles promotes the reversibility of sulfur due to its ability to form an ionic cluster with polysulfide. In addition to that, the conductive character of the Cu nanoparticles significantly minimizes the kinetic barrier during the electrochemical conversion of sulfur.$^{57}$

Xia et al. investigated a rechargeable Al–S battery comprising a S–C cathode and dichloromethane (DCM) electrolytes. It could exhibit an initial discharge capacity of 113.64 mA h g$^{-1}$ (pure ionic liquid – 85.23 mA h g$^{-1}$), which retained its highest value of about 131.67 mA h g$^{-1}$ at the 3rd cycle and 104.69 mA h g$^{-1}$ at the 40th.$^{39}$

In addition to MOF-derived microporous carbon as the host for sulfur particles, a carbon nanofibers (CNF)-based assembly was also examined as the possible host materials for the sulfur cathode. Among the widely studied materials, carbon nanotubes (CNTs)$^{58-60}$ and carbon nanofibers (CNFs)$^{54,61,62}$ are commonly used carbonaceous cathode materials for lithium–sulfur batteries. They have high conductivity, good mechanical flexibility and strength, large surface area, and a porous structure. For instance, Yu et al. coated single-wall carbon nanotubes (SWCNT) on the separator of a reversible room temperature Al–S battery.$^{63}$ This, in turn, behaved as a significant barrier to mitigate the diffusion of the polysulfide species, and thereby showed a reduction in the cell’s polarization. A freestanding CNF paper as the host for sulfur was also fabricated for the cathode (Fig. 4).

Likewise, Smajic et al. designed a composite of sulfur with single-walled carbon nanotubes (SWCNT) as the cathode, and AlCl$_3$:EMIM as the electrolyte.$^{59}$ Besides the improved charge transfer kinetics, the addition of CNTs also improves the Al–S battery capacity with minimal decay per cycle. They unraveled the unique attributes of CNTs that minimize the capacity decay by enhancing the interaction between chloroaluminate ions and sulfur. However, despite the above-mentioned advantages offered by the SWCNT–S cathode, their studies have shown that the formation of short-chain polysulfides could hinder the diffusion of electrons (Fig. 5a),
resulting in capacity decay. A freestanding CNF paper as the host for a sulfur electrode with Li$^+$ ions-mediated electrolyte was designed by Yu and his group to promote reversibility in the Al–S battery. With the addition of the Li-ions mediated electrolytes, the cell’s performance could be enhanced significantly. For instance, the initial discharge capacity of about 1000 mA h g$^{-1}$ was achieved, and the specific capacity as high as 600 mA h g$^{-1}$ could be retained after 50 cycles (Fig. 5c). The study showed that the Li-ions initiate the chemical reactivation of the Al polysulfo and during cycling (Fig. 5b). Furthermore, the presence of the Li$^+$ ion in the electrolyte (Al[EMI]Cl$_4$ ionic liquid) causes the kinetic barrier to become lower by the formation of soluble polysulfide intermediates. It also suppressed the formation of Al=S upon full discharge of the composite sulfur cathode.$^{49}$ DFT calculations were performed to understand the thermodynamic behavior of Li$_3$AlS$_3$ upon mixing of Li, Al and S atoms, where they have studied the relationship of the relative thermodynamic stability for Li$_3$AlS$_3$ and its phase segregated products as expressed in the following eqn (3):

$$\text{Li}_3\text{AlS}_3 \rightarrow \frac{3}{2}\text{Li}_2\text{S} + \frac{1}{2}\text{Al}_2\text{S}_3$$

The predicted mixing enthalpy $\Delta H_{\text{mix}}$ for Li$_3$AlS$_3$ showed a positive value of about 0.5 eV. Despite a substantial decrease in $\Delta H_{\text{mix}}$ and complete miscibility, it could not be achieved for the segregated phases, i.e., Li$_2$S and AlS$_3$. This was credited to the amorphization of Li$_2$S and AlS$_3$-like phases. It was predicted that upon full discharge of the sulfur cathode, the Li$^+$ ions mediated electrolyte could enhance the reactivation of the reduced sulfur products by suppressing the formation of the segregated phases, resulting in the improved performance of the Al–S batteries.$^{49}$

2.2 Metal sulfide/carbon composite for the Al–S batteries

The development in the metal–sulfur chemistry is critically impeded by various obstructions related to the sulfur cathode, such as its serious polysulfide solubility, shuttling, and slow kinetics. Massive efforts have been made to investigate other materials systems for the sulfur cathode. Among various known sulfur compounds, the high theoretical specific capacity of transition metal sulfides paves the way for enhancing the chemical stability and ionic/electrical conductivity. For
example, LiS64–66 and NaS67–71 have previously been examined as the potential candidate for the sulfur cathode in metal–sulfur batteries. Although the electronic conductivity of the majority of metal–sulfides is limited72–74 TiS2 (30–50 S cm−1),74 Co9S8 (290 S cm−1),75 their polar character provides a tremendous binding effect. Hence, this offers a strong chemical interaction to mitigate the polysulfide dissolution or shuttling.67,76,77 Besides the strong chemical binding with polysulfide, such cathodes offer another significant intrinsic benefit of a substantial sulfphilic property.68,78

The molybdenum sulfide polymorph, such as Mo6S8, is one of the oldest known cathode materials that is being examined for various metal anode systems, for instance, Li–S and Mg–S. The unique crystal structure of Mo6S8 has stacks of Mo6S8 blocks composed of the sulfur anionic cubic cell with an octahedral cluster of Mo atoms. Due to availability of two different types of intercalation sites in Mo6S8 structure, it can easily accommodate cations, such as Li+, Cu+, and Mg2+.51,79,80 Inspired by its unique crystal structure, Aurbach and co-workers used Mo6S8 as a cathode to demonstrate a rechargeable Mg-ion battery.81 Geng et al. examined the same materials system, i.e., Mo6S8, for their possible applications in room-temperature Al–S batteries. The initial discharge capacity of about 40 mA h g−1 and 25 mA h g−1 at current densities of 60 mA g−1 and 120 mA g−1, respectively, could be achieved.82 The reversibility of the cell for over 50 cycles could be achieved due to the formation of Al3Mo6S8 after Al intercalated Mo6S8 at two different sites in the crystal lattice. Further investigations are needed towards understanding the Al intercalating and trapping mechanism. Over the past few years, numerous efforts have been made towards various issues encountered by the sulfur cathode for Al–S batteries Wang et al. examined metal–sulfur composite materials.33 A cathode comprising Ni3S2/graphene micro flakes was designed for a rechargeable Al-ion battery. The bond length of Ni in the heazlewoodite Ni3S2 crystalline structure is 2.50 Å. It shows a trigonal crystal lattice orientation with the R32 space group.83,84 The metal sulfide composite cathode showed an initial discharge capacitance of 350 mA h g−1 and a discharge capacity of 60 mA h g−1 at the 100th cycle when the current density was 100 mA g−1. A reversible capacity of about 50 mA h g−1 could still be achieved at 200 mA g−1 after 300 cycles with the coulombic efficiency of about 99% (Fig. 6a and b). Yu et al.86 synthesized hexagonal NiS nanobelts with nickel(II) acetate and sodium thiosulfate as the precursors through a temperature hydrothermal method (Fig. 6c). As illustrated in Fig. 6d, the obtained NiS nanobelts exhibited a specific capacity of about 100 mA h g−1 at 200 mA g−1 after 100 cycles with good cyclic stability of 90% capacity retention. Fig. 6e shows the performance of the composite cathode up to current densities of 300 mA g−1, exhibiting high cycling stabilities. After 10 cycles, 111.7 mA h g−1 discharge capacity was observed at 150 mA g−1.
which decreased to 83.6 mA h g\(^{-1}\) at 300 mA g\(^{-1}\) and 96.8 mA h g\(^{-1}\) at 150 mA g\(^{-1}\). Although the Al\(^{3+}\) diffusion was enhanced, the battery suffered from low voltage plateaus (1.15 V vs. Al\(^{3+}/\)Al). Unveiling the mechanism of the charge/discharge reactions in sulfur-based cathodes is of great importance. Characterization tools (for instance XRD, TEM and XPS) could offer approximate information about the structural and chemical integrity of the electrode material upon charge/discharge reactions. However, it is utterly complicated to obtain the exact information about the chemical processes due to the inherent limitations of these techniques. Besides that, these techniques suffer from voltage fluctuations or oxidation on exposure to air, which results in inaccurate information. Contrary to the conventional characterization techniques, in situ characterization or operando methods have become increasingly important, as it has become possible to monitor the instant activities that may occur in the vicinity of the electrode. In addition, such methods provide discernment to the intermediate reactions and corresponding phases. Thus, they can be more precise and reliable. Recently, Li et al.\(^85\) evaluated the surface chemistry of the Co\(_3\)S\(_4\) electrode upon charge/discharge reaction using in situ Raman spectroscopy. Unlike the conventional spectroscopic techniques, in situ Raman spectroscopy could unveil the localized concentration variation of the tetrachloroaluminate ions on the surface of the electrode, which possibly could be due to intercalation of the dissociated products of Al\(_2\)Cl\(_7\) into the Co\(_3\)S\(_4\) structure during discharge. However, during charging, the ions could not be fully extracted. The Co\(_3\)S\(_4\) cathode exhibited an initial discharge capacity of about 287 mA h g\(^{-1}\) at a current density of 50 mA g\(^{-1}\), and the cathode retained a capacity of 90 mA h g\(^{-1}\) after 150 cycles. The Co\(_3\)S\(_4\) cathode exhibited an initial discharge capacity of about 287 mA h g\(^{-1}\) at a current density of 50 mA g\(^{-1}\), and the cathode retained a capacity of 90 mA h g\(^{-1}\) after 150 cycles.\(^85\)

The Zheng group fabricated a graphene/CoS\(_2\)/S composite cathode for a rechargeable Al–S battery.\(^87\) The composite with a high S-content of about 48% led to a high discharge capacity of 1145 mA h g\(^{-1}\) for the current density of 50 mA g\(^{-1}\) and stable cycle life for over 38 cycles. The surface of the as-fabricated composite sulfur cathode was in direct contact with the rGO-coated separator, which behaved as a barrier to mitigate the diffusion of polysulfides and lowered the polarization. With the addition of a small amount of CoS\(_2\) of 9 mg cm\(^{-2}\) into the composite structure, the enhancement of the polysulfide electrochemical reaction could be expedited due to the strong adsorption sites provided by the CoS\(_2\) particles. Zhang et al. prepared 2D layered materials (e.g., MoS\(_2\), WS\(_2\), and BN) as sulfur fixers during repeated charge/discharge processes for achieving high-performance rechargeable Al–S batteries. BN/S/C, when used as the cathode, exhibited the highest capacity of 532 mA h g\(^{-1}\) @ 100 mA g\(^{-1}\). In addition, a remarkable life span of up to 300 cycles with a high coulombic efficiency of 94.3% and discharge plateaus at ~1.15 V vs. AlCl\(_4\)\(^-/\)Al was displayed for the Al–S battery.\(^88\)

### 2.3 Covalently bonded sulfur composite for Al–S batteries
As discussed in previous sections, the elemental sulfur cathode or sulfur–carbon composite cathode or metal-sulfide-based
sulfur cathode was found to suffer low discharge capacity and low coulombic efficiency, which are directly related to polysulfide diffusion. A polymeric sulfur host, such as sulfurized polyanacrylonitrile (SPAN) has widely been used in Li–S99–101 and Na–S batteries92,93 as the cathode due to its strong covalent bonding. PAN constitutes the nitrile group, which on heating with elemental sulfur, gets destabilized to form SPAN. It is a polymeric sulfur composite composed of pyrrolidine rings constituting C–C, C–N, and S–C bonds in the molecular structure of the SPAN composite.94–96 The electrochemical reactivity of SPAN with monovalent cations is well established with good electronic conductivity of 10−4 S cm−1.97 Wang et al. studied the reactivity of the SPAN cathode with Al3+-ions. Owing to the high covalent binding in SPAN, significant control over the high irreversibility of AlSx could be achieved. Unlike intercalation or conversion type cathode, the S–S bond peaks upon binding of Al3+ ions with active sites of S/N atoms, as confirmed by the cyclic voltammograms (CVs) depicted in Fig. 7a. It should be noted that the increase in discharge capacity from 320 mA h g−1 to 605 mA h g−1 remains at 210 mA h g−1 after a subsequent number of cycles (Fig. 7b). Fig. 7c depicts the low voltage plateau (1st discharge cycle) with the low reduction potential curve (1st scan in the cyclic voltammetry curve). The SPAN composite demonstrated high rate capabilities of 343, 258, 160, 93 and 54 mA h g−1 @ 0.025, 0.05, 0.1, 0.2, and 0.5 A g−1, respectively (see Fig. 7d).98 Detailed information on the sulfur materials for room-temperature Al–S batteries is summarized in Table 1.

3. Outlook and future perspectives

As summarized in this mini-review, more progress has been made towards the development of a stable sulfur cathode for the long-life and reversible Al–S batteries. The focus has been laid on the latest advances of sulfur cathode components.99 The positive and negative aspects have been well considered. Research on rechargeable Al–S batteries is booming, and the bottleneck for this battery system can thus be further eliminated. However, owing to the polysulfide shuttling effect of the sulfur electrode and the underlying factors, the Al–S batteries go through complex electrochemical mechanisms rendering their sluggishness in the kinetics of the entire cell. In addition to the conventional challenges encountered by the sulfur cathode, the volumetric strain and the shuttle effect due to the inclusion of Al3+ need to be studied in detail. The aim towards high energy density batteries could be achieved by redefining the problems as they claim, “New challenges are redefined under these boundaries regarding low ionic conductivity of high-concentrated Al-PSs electrolyte, saturation and premature precipitation of Al-PSs, and rapid failure of the metal anode”. The aforementioned challenges can be mitigated through modifications in the chemical and physical structure of the sulfur cathode. These approaches can be investigated to provide comprehensive knowledge, and applied towards improving the kinetics of Al batteries (Fig. 8).

For example, the inclusion of a core–shell structure has made tremendous progress in lithium and sodium–sulfur battery systems.91,100 They exhibited stronger physical
| S. no. | Material system | Synthesis route | Initial discharge capacity (mA h g⁻¹) | Discharge (mA h g⁻¹) at ~n cycles | Advantage | Disadvantage |
|--------|-----------------|-----------------|----------------------------------------|----------------------------------|-----------|-------------|
| 1      | Ketjen black carbon/S¹¹ | Solution-processed slurry based method | 1200 @ 120 mA g⁻¹ | <50 after 4 cycles | High energy density (1700 W h kg⁻¹) | Inability to oxidize Al₂S₃, fast capacity decay |
| 2      | Activated carbon cloth/S⁵³ | Melt diffusion method | 1320 @ 50 mA g⁻¹ | 1000 @ 20th cycles | Improved reversibility | Large voltage hysteresis, slow solid-state sulfur conversion reaction, and low cycle efficiency (20 cycles) |
| 3      | S@CMK-3 (ref. 56) | Melt diffusion method | 1390 @ 251 mA g⁻¹ | >400 after 20 cycles | Enhanced charge/discharge kinetics and high S-utilisation of 82% | Short cycle life (20 cycles) and polysulfide diffusion |
| 4      | Carbonised-ZIF/S⁴⁴ | Solvothermal synthesis | 1410 @ 200 mA g⁻¹ | 420 after 30 cycles | Passivation of Al anode and inhibition of hydrolysis of Al₂S₃ | Large voltage hysteresis and low S-loading (0.2 mg cm⁻²) |
| 5      | S@HKUST-1-C⁵⁷ | Hydrothermal method | 1200 @ 1000 mA g⁻¹ | 460 @ 50th cycles | Improved reversibility and enhanced cycle efficiency | Reduced utilization of sulfur |
| 6      | S³⁹ | Commercial S-powder | 113.64 @ 15 mA g⁻¹ | 104.69 after 40 cycles | Improved performance of the electrolyte | Low discharge capacity and loss of sulfur |
| 7      | CNF/S as cathode & SWCNT coated separator⁴⁸ | Vacuum filtration and vacuum drying | >1200 @ 0.05C | >450 after 10 cycles | Reduced polarization and alleviate polysulfide diffusion | Low S-loading (1 mg cm⁻²) |
| 8      | SWCNT/S⁵⁰ | Nondestructive sublimation–deposition method | >900 @ 100 mA g⁻¹ | 1024 @ 2nd cycle | Improved charge transfer kinetics | Capacity decay |
| 9      | CNF/S as cathode & SWCNT-coated GF separator⁴⁹ | Vacuum filtration and vacuum drying and activation process | ~1000 @ 0.05C | 600 after 50 cycles | Enhanced reversibility and suppressed formation of Al=≡S | Low sulfur-loading (1 mg cm⁻²) |
| 10     | Mo₆S₈ (ref. 82) | Chemical intercalation process | 148 @ 12 mA g⁻¹ | 70 after 50 cycles | Good cyclic stability | Low discharge capacity |
| 11     | Ni₃S₂/graphene micro flakes³³ | Mixing | 350 @ 100 mA g⁻¹ | 60 after 100 cycles | High discharge voltage plateau (≈ 1.0 V vs. Al/AlCl₄⁻) | Side reactions due to the dissociation process of complexes (Al₂Cl₇⁻) |
| 12     | NiS⁸⁶ | Hydrothermal method | 104.7 @ 200 mA g⁻¹ | 100 after 100 cycles | Facilitation of the electrolyte immersion and enhancement of Al¹⁺ diffusion, high storage capacity, good cyclability and low overpotential | Low cell voltage plateau (~1.15 V vs. Al⁴⁺/Al) |
| 13     | Graphene/CoS₄/S as cathode & rGO coated separator⁸⁷ | Melt diffusion method and hydrothermal method | 680 @ 50 mA g⁻¹ | 1145 after 37 cycles | Suppress polysulfide diffusion and decrease polarisation | Low sulfur loading of 0.2–0.4 mg cm⁻² |
| 14     | SPAN⁹⁸ | Thermal polymerization | 343 @ 0.025 mA g⁻¹ | 201 after 20 cycles | Superior rate capacities | Low cycle life and low S-loading (1.5 mg cm⁻²) |
| 15     | Co₃S₁ (ref. 85) | Solvothermal process | 287.9 @ 50 mA g⁻¹ | 90 after 150 cycles | Good rate performance | Rapid capacity loss and limited current density |
confinement and high chemical adsorption, resulting in improved electrolyte penetration and prevention of diffusion of polysulfide. For instance, a core–shell structure of the sulfur nanospheres@ultrathin δ-MnO₂ composite for Li–S batteries with tailorability high S-mass ratio of 82 weight% was investigated by Li et al. (Fig. 9a). The composite showed a remarkable specific capacity of about 846 mA h g⁻¹ at 1C with an areal capacity of 502 mA h cm⁻² @ 1 mA h cm⁻². Another group
(Xiao Liang and Linda) worked on the in situ reaction assembly of the core–shell sulfur–MnO2 cathode of the Li–S battery, where the low capacity fading of 0.039% per cycle for over 1700 cycles was achieved. We expect to achieve similar landmarks. However, the complex reaction chemistry, due to the trivalent nature of Al3+, demands more extensive research on the cathode, anode, and electrolyte separately.

To further boost the redox kinetics of the sulfur conversion, there has been a recent review by the Xiao group in the introduction of single-atom catalysts (SACs) into metal–sulfur batteries. Recently, single-atom catalysts have gained broad interest due to their interactions in accelerating the transfer of charge between the catalysts and substrates. SACs can further reduce the polarization and mitigate the shuttle effect by accelerating the sulfur conversion mechanisms (Fig. 9b). In 2020, Wang et al. studied an iron single-atom catalyst anchored on a nitrogen-rich metal–organic framework (MOF) to accelerate the conversion mechanism of the sulfur cathode. Their fabricated FeS@CN/S composites exhibited low capacity decay of 0.06% per cycle for 500 cycles at 4C, owing to its confined porous monocage-like structure. The high charge density of Al provides hindrances in the efficiency of the battery, which altogether slows down the reaction kinetics of the sulfur cathode. The introduction of SACs into the Al–S batteries can enhance the sluggishness of the cell due to their high catalytic efficiency. In the future, more efforts are expected towards the development of SACs. Thus, its mechanisms can therefore be further explored. Covalent-organic frameworks have been advanced as an advanced electrode material for lithium-ion and sodium ion batteries due to their highly featured “structural diversity, framework tenability and functional versatility”. They can be constructed via incorporation of redox-active sites or units into a porous organic framework to enhance the storage capacity. Owing to the rich-physiochemical properties of the covalent-organic frameworks, they could be served as the potential host for elemental sulfur in the Al–S battery. Other possible solutions to enhance the cathode’s performance may include the addition of polyimide particles, organic-based electrode materials or nitrogen dopants as the suitable host for the Al–S battery system. Efforts towards the incorporation of metal sulfides or other components with the core–shell architecture to developing favourable cathodic materials for a feasible Al–S battery system.

The development of Al–S batteries with high electrochemical cyclic stability can be explored and carried forward to the next-generation battery system. Even with all these efforts towards this system, the research on Al–S still has a long way to reach its practicality. We believe that our proposed mini-review can influence the research to contribute more towards understanding the cathodic system in Al–S batteries.

Conflicts of interest
There are no conflicts to declare.

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