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1. Introduction

Supercapacitors are advanced electrochemical energy storage devices that have attracted tremendous scientific interest in recent years. Supercapacitors are known to have potential advantages in meeting the demand for future electronic devices because of their enhanced charge storage mechanism, higher cycling stability, high-rate capability, and increased power density.¹ As the demand for clean energy sources and energy storage, and conversion technologies increased, supercapacitors have emerged as a promising technology to store renewable energy resources. Supercapacitors are excellent recovery systems due to their shorter charging and discharging times and extended life cycles compared to conventional batteries. They can sustain millions of cycles of charge storage, but suffer from limited energy densities and low potential windows. Supercapacitors store electric energy in electric double layers between electrodes, unlike batteries that experience swelling.2

Micro supercapacitors (MSC) are developing quickly with potential uses in microelectronics, on-chip energy storage, integrated sensors, portable and wearable electronics, and photonic devices with a focus on versatility, miniaturization,

Manipulating charge carrier interactions at solid electrolyte interfaces for enhanced micro-supercapacitor performance†

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Manipulation of ionic charges in the solid electrolyte interface can result in unprecedented device characteristics for energy conversion and storage applications. An ultrathin gel polymer electrolyte film in a stacked device architecture has been proposed to offer a crucial method for exhibiting the characteristics of electric double layer capacitance (EDLC) at interfaces. A thorough understanding and control of ionic transport channels within the electrode/electrolyte interface are essential for the development of suitable design concepts and the fabrication of micro-scale energy storage devices. A stacked device with nanostructured thin gel polymer electrolyte (GPE) – polyvinyl alcohol (PVA) with lithium perchlorate (LiClO₄), and additive lithium sulphate (Li_2SO_4) were studied for micro-supercapacitors (MSCs). The electrochemical properties of supercapacitor devices were studied to confirm how the anions and cations are separated at electrode/electrolyte interfaces utilizing an electrometive force. Significantly, the charge migration and separation of cations and anions at the electrode/electrolyte interfaces were also studied.

and shape-tuning capabilities.^{3–6} Despite having several names, supercapacitors still serve the same basic purpose of storing energy between an electrolyte and a solid electrode. Despite their advantages, supercapacitors face challenges like high self-discharge and low energy density, necessitating extensive research on electrode materials and electrolyte development to enhance their competitiveness.^{7–12} But in addition to the electrode materials, the performance of supercapacitors is also influenced by the properties of the electrolyte. The development of dendrites on the metal anode surface during repeated charge/discharge cycles in liquid electrolytes is a significant issue that affects the performance of supercapacitors.¹³ This can lead to short circuiting and the subsequent loss of capacity.

Numerous research teams have worked very hard in recent years to prevent dendritic growth. Several innovative methods have been suggested, such as incorporating an artificial solidelectrolyte interphase (SEI) or adding additives to the electrolyte solution. Structurally confined gel polymer electrolyte (GPE) offers several advantages over other electrode types in the creation of supercapacitors without altering the electrode's characteristics.14,15 Gel-based electrolytes are influenced by polymer-based electrolytes, which is used to overcome the electrolyte leakage issue and make it simple to fabricate supercapacitor devices.¹⁶⁻²⁰ The polymer electrolytes based on polyvinyl alcohol (PVA) are some of the most commonly reported ones, with 5 to 67 Wh kg^{-1} as their stated energy densities.21-23 In general, polymer electrolytes have ionic conductivity lower than 10^{-3} S cm⁻¹, which is lower than that of aqueous electrolytes (10⁻¹ S cm⁻¹).²⁴ To boost its ionic

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conductivity and electrochemical activity, PVA is frequently employed as an electrolyte in combination with H_2SO_4 , KOH, LiCl, TiO₂, LiClO₄, and H_3PO_4 .^{25,26} Neutral GPEs were shown to have a larger operating potential window than basic and acidic GPEs. This is due to the fact that neutral salts have fewer hydrogen and hydroxide ions than acidic or alkaline substances.²⁷

Krishnan et al. studied high-performance supercapacitors, revealing that changes in ionic charges at the solid-electrolyte border create new energy conversion and storage features.³ They demonstrated EDLC properties using a planar device with a thin film, revealing dynamic supercapacitance characteristics. J. Zhao et al. found that adding polysulfide to a carbonate electrolyte significantly increased the number of polyethylene oxide (PEO) polymers, inhibiting the growth of lithium dendrites, which enhanced the stability of lithium metal batteries during extended cycling. The polysulfide formed a durable solid electrolyte interphase film, suppressing lithium dendrite development and enhancing the battery's long cycling.²⁸ F. Wan found that the Na₂SO₄ additive inhibits Zn dendrite and NVO nanobelt dissolution, resulting in highperformance aqueous ZIBs with NVO nanobelts as positive electrodes, resulting in negligible redox reactions, long cycle life, and high-capacity retention.²⁹ X. Guo et al. found that adding lithium chloride (LiCl) stabilizes the Zn metal anode and inhibits dendrite growth. Li⁺ cations form a shield on the Zn surface, while Cl⁻ anions reduce zinc polarization and aid in ion transport. Cells with LiCl electrolyte additive showed greater stability during cycling, decreased side reactions, and increased capacity.³⁰ K. Krishnan et al. studied the ionic charges in the interfacial domination of ITO/polyvinyl alcohol-KOH, focusing on electrical double-layer capacitance (EDLC) characteristics. They found that planar MSC cells showed significant differences in volumetric capacitance and retention. The study also examined the distribution of ionic charges at interfaces with thin and thick PVI-KOH in planar device arrangements. Limiting the persistence of PVI-KOH to 28 nm improved capacitance retention, clarifying the confinement effect of MSC bias.³¹ Zhang et al. in the year of 2015 reported that Li₂SO₄ could be readily added to PVA aqueous solution in large quantities to address the issue of low ionic conductivity of polymer gel supercapacitors. $LiCF_3SO_3$, $LiN(SO_2CF_3)_2$, $LiPF_6$, $LiBF_4$, and LiClO₄ are among the various lithium salts that have also been investigated as conducting salts in PVA hydrogel.32,33 There are several studies investigating how the structural, electrochemical, and mechanical characteristics of gel electrolyte are affected by Li₂SO₄ and how it changes the ionic conductivity of the gel, enhancing ionic conduction via interactions with the polymer matrix, and raising the concentration of free lithium and sulfate ions as charge carriers.34-36

The use of Li_2SO_4 in PVA–LiClO₄ gel electrolyte is a significant advancement in supercapacitor electrolyte design, addressing limitations of single-ion additive systems. Li_2SO_4 can provide Li^+ cations and SO_4^{2-} anions, enhancing bulk electrolyte properties and electrode–electrolyte interface dynamics. Traditional PVA gel electrolytes use single-ion additives for conductivity enhancement, achieving ionic

conductivities up to 48 mS cm⁻¹.³⁷ ClO₄⁻ anions in these systems show limited interfacial activity, while H₂SO₄ containing PVA hydrogels show high conductivity but face challenges with hydrogen evolution reactions at high voltages.38 The main limitation of these additives is their monofunctional nature, focusing on enhancing cation mobility or anion effects but not simultaneously. However, when Li2SO4 is added to PVA-LiClO4 gel matrix, Li⁺ ions maintain high conductivity, while large divalent SO42- anions adsorb on electrode surfaces, modifying the electric double layer structure.³⁹ Hybrid devices using Li₂SO₄ gel electrolytes showed higher areal capacitance and energy density compared to LiCl-based electrolytes, with 76.8% retention after 1000 cycles. The stability of Li₂SO₄ in aqueous environments addresses the corrosive nature of LiClO₄ electrolytes. Thus PVA-based gel electrolytes incorporating a variety of electrolyte ions exhibit high conductivity and good chemical stability over a wide pH range.^{39,40} The dual-ion approach in $SS || PVA-LiClO_4 + Li_2SO_4 || SS particularly benefits hybrid super$ capacitors requiring balanced capacitive and faradaic performance, while its compatibility with industrial-scale PVA processing methods makes Li2SO4 as a transformative additive for next-generation energy storage systems.

Gel electrolytes for supercapacitors often suffer from low ionic conductivity and poor cycling stability due to lithium dendrite formation. This study addresses these issues by proposing a new gel electrolyte system in which Li₂SO₄ is introduced into a PVA-LiClO₄ matrix, resulting in enhanced ionic conductivity and effective dendrite suppression on a stainless-steel substrate. Using a two-electrode cell design, electrochemical performances such as cyclic voltammetry (CV), galvanostatic charge-discharge cycles (GCD), electrochemical impedance spectroscopy (EIS), and ionic conductivity before and after cycles of stack MSCs were studied. This simple yet effective modification achieves an ultra-high volumetric capacitance of 11.8 F cm $^{-3}$ and maintains stability over 2000 cycles, outperforming conventional PVA-based electrolytes. Li2SO4 additive enhanced the performance of the supercapacitor device by acting as a safer and more efficient gel electrolyte for high-performance supercapacitors by improving the overall volumetric capacitance and cycling stability of the stack MSC.

2. Materials and methods

2.1 Fabrication of gel polymer electrolyte – based stacked devices

2.1.1 Preparation of polymer electrolyte for SS||PVA-LiClO₄||SS (without additive). PVA, (molecular weight = 7×10^4 g mol⁻¹) was used for the thin film. 1 g of PVA was dissolved in 16 ml of distilled water. It was then stirred continually for half an hour (@90 °C). A homogeneous solution was acquired. Then, 0.2 g of LiClO₄ salt solution was prepared in 1 ml of distilled water. The above two solutions were mixed to form a homogeneous mixture. The final solution was stirred for ten minutes at room temperature. At last, 200 microliters of this solution were spin-coated on the planar SS disc.

2.1.2 Preparation of polymer electrolyte for SS||PVA-LiClO₄ + Li₂SO₄||SS (with additive). First, 0.2 g of lithium perchlorate



Scheme 1 Schematic representation of device fabrication (a and b) and coin cell assembly (c).

salt was dissolved separately in 1 ml of distilled water. Then 1 g of polyvinyl alcohol was mixed in 16 ml of distilled water and stirred continuously for half an hour (at 90 °C) to get a homogeneous mixture. After that, as an additive 0.1 g of Li₂SO₄ was separately dissolved in 1 ml of distilled water, then both the LiClO₄ solution and Li₂SO₄ solution (with additive salt) were mixed together to form a uniform solution; this mixture was stirred for ten minutes at room temperature. Finally, 200 µl of this solution were spin coated on the planar SS disc to make the stack device.

2.1.3 Fabrication of stacked micro-super capacitor. 200 microliters of PVA-based electrolyte solution were spin-coated on the SS disc. In this device fabrication, the spin-coating method is considered as an essential technique to achieve the self-assembled film morphology (Scheme 1). The prepared PVA-based film thickness was found to be 246.7 nm using atomic force microscopy. A SS||PVA-LiClO₄||SS device and SS||PVA-LiClO₄ + Li₂SO₄||SS in a stacked configuration were fabricated, in which the SS disc was placed on both sides of the electrolyte. The active area of the device is 1.53 cm².

3. Characterizations

The electrochemical characteristics of the EDLC devices were measured with a two-electrode configuration. The cyclic voltammetry (CV), galvanostatic charge–discharge (GCD), and impedance were conducted using Bio-Logic BCS-810. In the CV measurements, the sweep rate varied from 5 to 100 mV s⁻¹ and the potential window varied in the range of 0–1 V. In the GCD studies, the device was charged and discharged with a current of 0.5 and 1 microampere. By using CV curves, the volumetric capacitance (C_V) was calculated. By using GCD profiles, the discharge capacitances of the devices were estimated. The surface morphology and thickness of the polymer film were observed using AFM.

4. Results and discussions

The electrochemical performances of SS $||PVA-LiClO_4||SS$ stack MSC were studied in the potential window of 0–1.0 V. CV measurements of SS $||PVA-LiClO_4||SS$ at several sweep rates

varying from 5–100 mV s⁻¹ were plotted. CV plots of MSC (Fig. 1a) correspond to quasi-rectangular shapes, which indicates electric double layer behaviour. In addition, the quasirectangular behaviour of the CV curve was maintained throughout all sweep rates, indicating that the SS||PVA-LiClO₄||SS has both excellent rate capability and optimal capacitive behaviour. As the sweep rate was increased from 10– 100 mV s⁻¹, the volumetric capacitance dropped from 5.8 F cm⁻³ until it reached 1.9 F cm⁻³ (Fig. 1c). The volumetric capacitance (C_V) was estimated by the following relation:

$$C_{\mathrm{Vol}} = rac{1}{2 imes \mathrm{vol} imes \mathbf{v} imes \Delta V} \int i(t) \mathrm{d}t$$

Here, ν is the scan rate, ΔV is the potential difference, *i* is the measured current in CV, and *t* is time. Using electrochemical impedance spectroscopy (EIS), the charge transfer properties of the SS||PVA-LiClO₄||SS device were analysed. From the Nyquist plot (Fig. 1e), the increased ionic conductivity of the gel polymer electrolyte is further supported by the absence of a half circle pattern in the high frequency zone. The ionic conductivity (σ) of the thin film was measured using the following relationship:

$$\sigma_{\rm AC} = \frac{t}{R_{\rm ct} \times A}$$

where *d* is the distance between the electrodes, $R_{\rm ct}$ is the resistance value obtained directly from EIS measurement, and *A* is the area of the electrode. According to the calculations, the pristine device has an ionic conductivity of about 7.6 × 10^{-7} S cm⁻¹, and after EC studies, the ionic conductivity was found to be 7.6 × 10^{-7} S cm⁻¹.

Using a two-electrode cell design, we studied the electrochemical performance of SS||PVA-LiClO₄ + Li₂SO₄||SS. Cyclic voltammetry (CV) measurements were performed in a potential window of 0–1.0 V at several scan rates (5–100 mV s⁻¹). The cyclic voltammetry graphs (Fig. 1b) showed a quasi-rectangular shape across all sweep rates, representing the pseudocapacitive behaviour. The quasi-rectangular behaviour of the CV curve was maintained throughout the sweep rates, indicating the good capacitance behaviour of SS||PVA-LiClO₄ + Li₂SO₄||SS and its rate capability. According to calculations, the maximum CV of the SS||PVA-LiClO₄ + Li₂SO₄||SS is found to be 10.5 F cm⁻³ at

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Fig. 1 (a) CV curves at different scan rate of device SS||PVA-LiClO₄||SS (b) CV curves at different scan rate of device SS||PVA-LiClO₄ + Li₂SO₄||SS, (c) scan rate dependent C_{VOI} of SS||PVA-LiClO₄ + Li₂SO₄||SS, (d) scan rate dependent C_V of SS||PVA-LiClO₄ + Li₂SO₄||SS (e) EIS measurement of SS||PVA-LiClO₄ + Li₂SO₄||SS before and after cycling (f) EIS measurement of SS||PVA-LiClO₄ + Li₂SO₄||SS before and after cycling.

10 mV s⁻¹. Fig. 1d shows a volumetric capacitance of a minimum value of 2.8 F cm⁻³ at 100 mV s⁻¹ as the sweep rate was increased. Using electrochemical impedance spectroscopy (EIS), the charge transfer properties of additive-based SS||PVA-LiClO₄ + Li₂SO₄||SS were analysed. Fig. 1f shows a Nyquist plot, where the low-frequency area is represented by a straight line with a small semicircle, which is typically associated with a better capacitance. In addition, the gel polymer electrolyte's increased ionic conductivity is inferred from the lack of a semicircle pattern in the high-frequency area.

The ionic conductivity of SS||PVA-LiClO₄ + Li₂SO₄||SS is about 1.18×10^{-6} S cm⁻¹, while the ionic conductivity of the device after EC tests increased to 1.1×10^{-6} S cm⁻¹. After 1000 continuous CV cycling of SS||PVA-LiClO₄ + Li₂SO₄||SS at 100 mV s⁻¹, additional EIS measurements were taken to analyse the ion transport property of the device. Later in the CV experiment, the cell's conductivity returned, showing that the device maintains its performance over time. It is also important to know the amount of current that is carried by each mobile charged species than the overall conductivity of a polymer when the polymer contains more than one mobile charged species. So, by considering the ionic resistance (*R*_{ele}), *t*_{ion} (the mean transfer number) can be estimated from the equation given below:

$$V_{\text{cell}} = \frac{R_{\text{i}}^{-1}}{(R_{\text{i}}^{-1} + R_{\text{ele}}^{-1})} \times V_{\text{emf}} = t_{\text{ion}} \times V_{\text{emf}}$$

Using the V_{cell} to t_{ion} relation, t_{ion} values were estimated to be 0.51 and 0.61 for SS||PVA-LiClO₄||SS and SS||PVA-LiClO₄ +

 Li_2SO_4 ||SS, respectively. This indicates that the gel polymer electrolyte of SS||PVA-LiClO₄ + Li_2SO_4 ||SS enhances ionic transport efficiency, possibly resulting in superior electrochemical performance. SS||PVA-LiClO₄ + Li_2SO_4 ||SS may provide enhanced performance, including greater capacitance, energy density, improved ionic mobility, and reduced hindrance within the polymer matrix when compared to SS||PVA-LiClO₄||SS.

Comparison of SS PVA-LiClO₄ SS and SS PVA-LiClO₄ + Li₂- $SO_4 \|SS$ is shown in Fig. 2a. Due to the presence of the electrical double layer at the interface, the CV plots of the devices, which is thickness dependent, displayed a quasi-rectangular behaviour. Quasi-rectangular shape in the CV curve was maintained throughout all scan rates, indicating that the polymer- Li ion integrated devices have excellent capacitance properties and a high-rate capability. The CV plot was considered to measure the volumetric capacitance $(C_{\rm V})$ of both devices. The compared volumetric capacitance of the devices based on the role of the additive is displayed in Fig. 2b. CV studies were performed according to the device capacity to evaluate the cycle characteristics of the constructed stack MSC device. As in Fig. 2c, the cycling performance of the SS||PVA-LiClO₄||SS stack MSC was inferior, with 67% of capacitance retention after 1000 cycles. It was found that at a specific characteristic nanoscale, the cycle performance and capacitance retention both improved by adding Li₂SO₄. Compared to the other device, the SS||PVA- $LiClO_4 + Li_2SO_4 ||SS|$ demonstrated superior cycling performance, with 1000 cycles on continuous use and considerable capacitance retention of 90%.

Similarly, using a two-electrode cell design, galvanostatic charge–discharge (GCD) characteristics of SS||PVA-LiClO₄||SS



Fig. 2 (a) Stability performance of device A (SS||PVA-LiClO₄||SS) and B (SS||PVA-LiClO₄ + Li₂SO₄||SS), (b) C_V comparison of device A and B, (c) C_V comparison at 10 mV s⁻¹ scan rate of device SS||PVA-LiClO₄ + Li₂SO₄||SS (A) and SS||PVA-LiClO₄ + Li₂SO₄||SS (B).

and SS $||PVA-LiClO_4 + Li_2SO_4||SS$ stack MSC were studied at currents varying from 0.5 to 1 micro ampere in the potential window of 0–1.0 V as shown in Fig. 3a and b.

By using GCD plots, the volumetric capacitances of SS $\|PVA-LiClO_4\|SS$ were found to be 6.821 and 3.76 F cm⁻³ at 0.5 to 1 microampere, respectively. The volumetric capacitances of SS $\|PVA-LiClO_4 + Li_2SO_4\|SS$ were found to be 11.8 and 4.43 F cm⁻³ at different currents of 0.5 and 1 microampere respectively, using the following equation:

$$C_{\rm Vol} = 2 \left(\frac{j \times \Delta t}{\Delta V} \right)$$

where *j* is the current density, Δt is the discharge time, and ΔV is the potential window of the cell.

In order to investigate the cycling performance of fabricated stack MSC devices, the GCD measurement was tested up to 2000 continuous cycles at a constant current of 1.0 μ A (Fig. 3d). It infers that the electrolyte PVA-LiClO₄ + Li₂SO₄ can give excellent device characteristics, which are comparable to the previously reported devices as mentioned in Table 1.

The stability of SS||PVA-LiClO₄||SS started fading at 300 cycles (Fig. 3c), whereas SS||PVA-LiClO₄ + Li₂SO₄||SS (Li₂SO₄ as additive) was stable even after 2000 cycles. The shape of the GCD profile remained almost unchanged (Fig. 3b) and exhibited good capacitance retention of 83% even after 2000 continuous cycles (Fig. 3e). The device with additive, showed excellent cycling characteristics, which indicate its potential for use in future energy-storage devices. Despite exhibiting a slightly lower



Fig. 3 (a) GCD 1st cycle at 0.5 microampere (b) 1^{st} cycle at 1 microampere, (c) cyclic stability of SS||PVA-LiClO₄||SS (d) cyclic stability SS||PVA-LiClO₄ + Li₂SO₄||SS (e) comparison of C_V retention of both devices (f) comparison with previous literature.

Table 1 Comparison of different electrolyte additive

SI no.	Electrolyte	Capacitance	Voltage	Capacitance retention/cycle life	Power density	Energy density	Ref. no.
1	PVA-KOH	5.39 F cm ⁻³	0-1 V	98%/11 000 cycles	300 mW cm ⁻³	2.36 mW h cm ⁻³	3
2	PVI-KOH	0.128 F cm ⁻³	0-1 V	92%/1400 cycles	6.89 mW cm ⁻³	0.056 mW h cm ⁻³	41
3	PAM (polyacrylamide)-LiCl	0.341 F cm ⁻³	0-0.6 V	99.12%/30 000	257.07 mW cm ⁻³	17.07 mW h cm ⁻³	42
4	PVA + LiClO₄+ Li₂SO₄	11.8 F cm⁻³	0-1 V	83% after 2000 cycles	6.58 mW cm ⁻³	1.6 mW h cm⁻³	This Work

coulombic efficiency than SS||PVA-LiClO₄||SS (Fig. S1(b and d)),† the inclusion of Li_2SO_4 is advantageous overall, since it promotes pseudocapacitive redox reactions at the electrode/ electrolyte interface, enhancing specific capacitance and cycling stability, hence enabling the device to maintain its performance over a greater number of cycles. Although these faradaic processes may induce minor side reactions that reduce coulombic efficiency per cycle,^{39,43,44} the overall energy storage and long-term dependability are significantly enhanced, making the Li_2SO_4 -based device the superior option for highperformance supercapacitors.

Microscopic analysis to find the thickness of the thin film coating was conducted using AFM. The AFM was measured by spin coating the SS disc with the electrolyte and then masked the film edge with respect to the substrate then the step height of the edge was measured (Fig. 4b). From the corresponding depth profile (Fig. 4c), the thickness of the film was found to be 246.7 nm.

Open circuit potential (OCP), being the crucial parameter, represents the voltage across the terminals of supercapacitors when no current is flowing, indicating energy storage in the device. A higher OCP indicates a higher energy storage capacity of the supercapacitor. Here, the device $SS||PVA-LiClO_4 + Li_2-SO_4||SS$ has a higher OCP than $SS||PVA-LiClO_4||SS$ (Fig. 5). It suggests that the device with the additive can store more energy than the device without the additive. This higher OCP leads to the storage of more power per unit volume compared to a device without an additive. $SS||PVA-LiClO_4 + Li_2SO_4||SS$ device has minimal voltage drop and maintains a stable OCV after the sweep, ensuring stability. Fig. 5 shows the time-dependent cell voltage of the $SS||PVA-LiClO_4||SS$ and $SS||PVA-LiClO_4 + Li_2SO_4||SS$ MSCs measured under a certain scan rate. The measurement was carried out in two steps, first, a positive



Fig. 5 Time-dependent V_{cell} of SS||PVA-LiClO₄||SS and SS||PVA-LiClO₄ + Li₂SO₄||SS measured as a function of time under pristine state v/s after cycling.

cyclic sweep was performed at 0–1 V at certain sweep rates. Then, the voltage was measured as a function of time under an open circuit condition. In case of SS||PVA-LiClO₄ + Li₂SO₄||SS MSC, a gradual increase in cell voltage for more than 300 s was observed because of the equilibration of ionic charges Li⁺ and ClO_4^{-}/SO_4^{2-} at the electrode/electrolyte interface, whereas in SS||PVA-LiClO₄||SS the cell voltage was dropping drastically. It was observed that the increase in scan rate increased the cell voltage for the device with additive, whereas the device without additive showed a decrease in cell voltage when the scan rate was increased due to the difference in ionic charge separated at the electrode/electrolyte interfaces.



Fig. 4 (a) 2D AFM image of the thin film (b), 3D image (c) height profile (thickness) of the film.

The cyclic voltammograms of both devices did not deviate from rectangular shapes even at higher scan rates. These results reveal that the electrolyte plays a crucial role in the capacitive property of both MSCs. The charge storage kinetics is analysed with the help of the power law as given by,

 $i = av^b$

where, *a* and *b* are adjustable parameters, ν is the scan rate of CV, and *i* is the responsive current. When *b* is equal to 1, the voltametric currents show ideal capacitive behaviour, whereas when *b* is closer to 0.5, the electrochemical process will be diffusion controlled.

Fig. 6(a & b) shows the dual logarithm plots for the cyclic voltametric current of both devices from 5 to 100 mV s⁻¹ in CV. From the dual logarithm plots, the slope b for SS||PVA-LiClO₄||SS and SS||PVA-LiClO₄ + Li₂SO₄||SS were found to be 0.66 and 0.61, respectively. The increased ionic transference number in SS||PVA-LiClO₄ + Li₂SO₄||SS (0.66) suggests that the system transitions towards a more diffusion-controlled process, which improves the ionic conductivity and electrode–electrolyte interaction. SO₄²⁻

ions, due to their smaller size and more hydration compared to ClO₄⁻, facilitate enhanced ion mobility in the electrolyte, hence promoting accelerated Li⁺ movement and boosting total ionic conductivity. The combination of monovalent (ClO₄⁻) and divalent (SO42-) anions results in a more effective ion transport network, reducing ionic clustering or ion pairing. The interaction between Li⁺ and SO₄²⁻ reduces ion aggregation and improves the dissociation of ionic species, hence decreasing the internal resistance. Li⁺ ions typically have a smaller hydration radius compared to SO_4^{2-} ions. A smaller hydration radius allows Li⁺ to approach the electrode surface more closely, enabling faster adsorption which leads to lithium deposition in SS||PVA-LiClO₄||SS device (Fig. 6c). When Li_2SO_4 is added SO_4^{2-} ions may compete with Li⁺ for access to the electrode surface as it has higher charge density than ClO₄⁻, which would potentially slow down lithium deposition at the surface as shown in Fig. 6d. SO_4^{2-} is a divalent anion with a higher charge density and forms strong hydration shells this helps stabilize dissociated Li⁺ ions by competing with ClO_4^- or other anions for Li^+ , thus reducing ion pairing. This results in an optimized ionic atmosphere that enhances mobility without excessive pairing. SO4²⁻ ions adsorb at



Fig. 6 (a) Dual logarithm plot of current *versus* scan rate of CV in SS $||PVA-LiClO_4||SS$, (b) dual logarithm plot of current *versus* scan rate of CV in SS $||PVA-LiClO_4||SS$, (c) deposition of Li⁺ in the electrode surface, (d) inhibition of further Li⁺ deposition by SO₄²⁻.

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the electrode–electrolyte interface by weak electrostatic interactions that are driven by the applied electric field. This reduces degradation of the electrode and maintains consistent performance over many cycles. The additional Li⁺ ions increase the overall concentration of charge carriers in the electrolyte, thereby enhancing ionic conductivity within the PVA gel matrix.³⁴ Here, a small amount of Li₂SO₄ is added, in which the sulfate ions participate in forming a passivation layer at the electrode surface. This layer inhibits the deposition of more Li⁺ ions at the electrode surface and further enhances the cycle stability of the device with Li₂SO₄ as an additive, which is evident from the GCD cycling which was stable even after 2000 continuous cycles.

5. Conclusions

The volumetric capacitance and the stability of the device were enhanced by the addition of Li₂SO₄ into the PVA-gel electrolyte. Thus, SS||PVA-LiClO₄ + Li₂SO₄||SS showed better ionic conductivity and the thin film device shows ultra-high volumetric capacitance 11 F cm⁻³ at 5 mV s⁻¹ scan rate. From the GCD plots, the volumetric capacitances of SS||PVA-LiClO₄||SS were found to be 6.8 F cm^{-3} and 3.76 F cm^{-3} at 0.5 μ A and 1 μ A respectively, whereas the volumetric capacitances of SS||PVA-LiClO₄ + Li₂SO₄||SS were found to be 11.8 F cm⁻³ and 4.43 F cm⁻³ at currents ranging from 0.5 μ A to 1 µA respectively. SS||PVA-LiClO₄ + Li₂SO₄||SS showed a good cycling performance, which was stable for 2000 continuous cycles with 83% capacitance retention. The device remarkably achieved an energy density of 1.6 mWh cm⁻³ and a power density of 6.58 mW cm⁻³. The optimized PVA-LiClO₄ + Li₂SO₄ gel electrolyte significantly inhibited the lithium deposition (dendrite development) on the surface of the SS disc during the charging and discharging process, which improved overall capacitance and cycle stability. In addition to the previously mentioned performance enhancements, our findings highlight the promise of Li₂SO₄ as a valuable additive in gel electrolytes for supercapacitor systems. The inclusion of Li₂SO₄ seems to enhance ionic conductivity and electrochemical stability. This study lays the groundwork for future investigations into the effective use of Li2SO4 in different electrolyte formulations and electrode materials, potentially leading to the improvement in energy density, power output, and device lifespan. In addition, understanding the mechanisms by which Li₂SO₄ inhibits dendrite formation could facilitate advancements in the creation of safer and more resilient supercapacitors and associated battery technologies in the future. In summary, our results enhance the development of gel electrolyte formulations, establishing a foundation for advanced supercapacitors that exhibit enhanced performance and stability.

Data availability

The data that support the findings of this study are available on request from the corresponding author.

Conflicts of interest

There are no conflicts to declare.

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