Hydrogel-polymer electrolytes for electrochemical capacitors: an overview

N. A. Choudhury, a S. Sampath b and A. K. Shukla a*c

Received 1st July 2008, Accepted 24th October 2008
First published as an Advance Article on the web 1st December 2008
DOI: 10.1039/b811217g

Electrochemical capacitors are electrochemical devices with fast and highly reversible charge-storage and discharge capabilities. The devices are attractive for energy storage particularly in applications involving high-power requirements. Electrochemical capacitors employ two electrodes and an aqueous or a non-aqueous electrolyte, either in liquid or solid form; the latter provides the advantages of compactness, reliability, freedom from leakage of any liquid component and a large operating potential-window. One of the classes of solid electrolytes used in capacitors is polymer-based and they generally consist of dry solid-polymer electrolytes or gel-polymer electrolyte or composite-polymer electrolytes. Dry solid-polymer electrolytes suffer from poor ionic-conductivity values, between $10^{-8}$ and $10^{-2}$ $\text{S cm}^{-1}$ under ambient conditions, but are safer than gel-polymer electrolytes that exhibit high conductivity of ca. $10^{-1}$ $\text{S cm}^{-1}$ under ambient conditions. The aforesaid polymer-based electrolytes have the advantages of a wide potential window of ca. 4 V and hence can provide high energy-density. Gel-polymer electrolytes are generally prepared using organic solvents that are environmentally malignant. Hence, replacement of organic solvents with water in gel-polymer electrolytes is desirable, which also minimizes the device cost substantially. The water containing gel-polymer electrolytes, called hydrogel-polymer electrolytes, are, however, limited by a low operating potential-window of only about 1.23 V. This article reviews salient features of electrochemical capacitors employing hydrogel-polymer electrolytes.

1. Introduction

Electrochemical capacitors (ECs) are energy storage devices with high power densities as evident from the Ragone plot shown in Fig. 1. ECs are complementary to secondary batteries and are projected as future power sources for high-power applications such as electric vehicles, utility load-leveling, heavy-load starting assists for diesel locomotives, military and medical applications as also low-power applications such as camera-flash equipment, lasers, pulsed-light generators, and as back-up power for computer memory.1–3 On the basis of mechanisms involved in the charge-storage process, an EC is classified as an electrical double-layer capacitor (EDLC) or a pseudocapacitor. An EDLC typically comprises an electrolyte sandwiched between two electrodes. High energy-density of EDLCs, as compared to dielectric capacitors, arises due to the large surface-area of the electrode materials such as activated carbons,4–6 aerogel or xerogel carbons,7–9 carbon nanotubes,4,5,10,11 and exfoliated graphite.12 Owing to the physical nature of charge storage and discharge, EDLCs have several advantages over secondary batteries, viz. faster charge-discharge, longer cycle-life (>105 cycles), and higher power-density.1 Pseudocapacitors, also referred to as redox capacitors, involve redox reactions in the charge storage and discharge processes.

Broader context

Electrochemical capacitors (ECs) are electrochemical energy storage devices with high power densities. ECs possess properties that are complementary to secondary batteries and hence can be hybridized to yield devices, which can be employed as power sources for high-power applications such as electric vehicles, utility load-leveling, heavy-load starting assists for diesel locomotives, military and medical applications as also for low-power applications such as camera-flash equipment, lasers and as back-up power for computer memory. Electrochemical capacitors employ two electrodes and an aqueous or a non-aqueous electrolyte, either in liquid or solid form. Solid electrolytes provide certain advantages such as freedom from leakage of any liquid component, compactness and reliability. Solid electrolytes employed in ECs are mostly polymer-based and are classified as solid-polymer electrolytes, gel-polymer electrolytes and composite-polymer electrolytes. Solid-polymer electrolytes exhibit lower ionic conductivity values than gel-polymer electrolytes. Gel-polymer electrolytes are generally prepared using organic solvents that are environmentally malignant. Hence, replacement of organic solvents with water in gel-polymer electrolytes is desirable, which also minimizes the device cost substantially. The water containing gel-polymer electrolytes are called hydrogel-polymer electrolytes.
The ECs employ either aqueous or non-aqueous electrolytes and it may be in liquid or solid form, the latter providing advantages of compactness, reliability, freedom from leakage of any liquid component and a large operating potential-window. The vapour pressure of the solid electrolytes is generally low and hence they are amenable for use at higher than ambient temperatures. They are also relatively easy to handle. Polymeric solid electrolytes are flexible, and can be easily fabricated as thin and large area membranes facilitating low internal-resistance. They also enhance the operational reliability of the ECs. Polymer electrolytes are the most widely studied solid electrolytes. Complexation of functional-groups of certain polymers with cations results in the formation of polymer–cation complexes commonly referred to as solid-polymer electrolytes (SPEs). Fenton et al. and Wright were the first to report the ionic conductivity measurements on poly(ethylene oxide) (PEO)–alkali metal salt complexes. However, it was Armand et al., in 1978, who pointed out the potential use of these materials in batteries. In order to achieve a well-complexed polymer–metal salt system, the choice of polymer as well as the metal salt is of utmost importance. The choice of a host polymer depends principally on factors such as (i) presence of a sequential functional group with high electron donation capability to form a strong coordinate bond with a metal ion, (ii) a low steric hindrance to bond rotations so that the polymer chain mobilities called segmental motions is facilitated, and (iii) an appropriate distance between the donor atom and the cation. The complexation of cations with the functional groups of the polymer and the stability of the resulting polymer–cation complex are explained on the basis of the Hard–Soft Acid–Base (HSAB) principle. Based on the HSAB principle, the polymers are considered as arrays of hard or soft bases that form coordination bonds with cations. The stability of polymer electrolytes is primarily dictated by the polymer–cation solvation energy with only a little contribution from the lattice energy of the salt.

Ionic mobility in SPEs has been reported to be due to segmental motion and the ion transport is mainly restricted to the amorphous phase of the polymer electrolytes. Mixing a polymer with an alkali metal salt dissolved in an organic solvent results in the formation of a gel-polymer electrolyte (GPE). GPEs exhibit liquid-like ionic conductivity while maintaining the dimensional stability of a solid system. Various delivery processes. Energy-storage mechanisms in pseudocapacitors involve fast Faradaic reactions such as intercalation, under-potential deposition or redox processes occurring at or near a solid–electrode surface at an appropriate potential. Redox processes often occur in metal oxides, conducting polymers and hybrid organic–inorganic nanocomposite materials making them attractive electrode materials for pseudocapacitors.
polymers, such as poly(propylene oxide), poly(ethylene imine), thio-alkane, polystyrene, poly(vinyl alcohol), poly(vinylidene fluoride) (PVdf), poly(vinylidene carbonate), poly(acrylonitrile), poly(vinyl chloride), poly(vinyl sulfone), poly(-phenylene terephthalamide), and poly(vinylpyrrolidone), have been reported to form GPEs with conductivities ranging between $10^{-4}$ and $10^{-3}$ S cm$^{-1}$ under ambient conditions. The low molecular-weight organic solvents with high dielectric constants employed in GPEs are commonly referred to as plasticizers. Solvents such as ethylene carbonate, propylene carbonate, tetrahydrofuran, dimethyl formamide, diethyl carbonate, diethyl phthalate, methyl ethyl carbonate, γ-butyrolactone, dimethyl carbonate, glycol sulfite, and alkyl phthalates, have been commonly employed as ‘plasticizers’ in GPEs. The glass-transition temperature of GPEs varies with the extent of plasticization which depends on the proportion of polymer and plasticizer in the GPEs. GPEs are generally prepared by heating a mixture of appropriate quantities of a polymer, a plasticizer and a salt to a temperature above the glass-transition temperature of the polymer followed by casting the resultant hot viscous solution onto a glass plate and cooling to ambient temperatures. When water is used as a plasticizer, the gelled polymer electrolyte is referred to as ‘hydrogel-polymer electrolyte’.

Hydrogels are three-dimensional polymeric networks that trap water in the polymer matrices mainly by surface tension. Based on the nature of the pendant groups in the precursor, hydrogels are classified as neutral or ionic, while the classification as affine or phantom networks is based on their structural and mechanical features. Alternatively, depending on the nature of the polymer, hydrogels are classified as homopolymeric or copolymeric hydrogels. Hydrogels are further classified as amorphous, semicrystalline, hydrogen-bonded structures, supramolecular structures and hydrocolloidal aggregates based on the physical structure of the network. The three-dimensional network formation of hydrogels and their insolubility in the precursor solution are due to the presence of chemical cross-linkages or physical entanglements. Physical hydrogels differ from chemical hydrogels in the type of cross-linkages, randomness of the network formation, and the effects of these parameters on the rigidity and elastic moduli of the resultant networks. Unlike the covalent cross-linking points in chemical hydrogels formed by the reaction between the polymer and a cross-linking reagent such as glutaraldehyde, physical hydrogels are formed through association of several laterally associated polymer helices in extended junction zones, wherein the hydrogel network is stabilized by physical entanglements, electrostatic attractive forces and hydrogen bonding. The schematic diagram of a chemical hydrogel with point cross-links and a physical hydrogel with multiple junction zones are shown in Fig. 2(a) and (b), respectively. Physical hydrogels are thermally reversible and can be viewed as viscoelastic solids. Physiologically responsive hydrogels often show a swelling behavior in response to a changing external environment. Some of the factors responsible for the swelling behavior are pH, ionic strength, temperature and electromagnetic radiation.

Poly(vinyl alcohol) (PVA), discovered by Haehnel and Herrmann in 1924, is one of the most widely investigated polymers. It is cheap, non-toxic and chemically stable. The –OH groups of PVA react with –CHO groups of certain aldehydes to form acetal or hemiacetal linkages under acidic conditions. The resultant polymeric entity with acetal or hemiacetal linkages is water insoluble and gel-like in nature. It can be cast to form thin large surface-area membranes suitable for use as electrolyte-selector in ECs. Hydrogels comprising synthetic polymers, such as PVA have high structural integrity and good mechanical properties. These hydrogels have a large water content absorbed in the polymer matrix that helps in fine-tuning their ionic conductivity. Such hydrogels, however, possess a high degree of hydrophobicity due to the presence of long carbon-chains with fewer hydrophilic pendant groups. By contrast, hydrogels consisting of natural polymers, such as gelatin, possess a high degree of hydrophilicity, due to the presence of a large number of hydrophilic pendant groups, which helps in high retention of water in the polymer matrix leading to an enhanced ionic conductivity.

Research in the field of biomaterials has led to advances in hydrogel science and technology supporting a wide spectrum of applications in biomaterials, agriculture, tissue engineering, aquaculture, infant care and nanotechnology. KCl-doped physical hydrogels have long been employed in salt bridges for electrochemical experiments. Unlike the hydrogel electrolytes comprising synthetic polymers, biopolymer-based hydrogel electrolytes with natural cross-linkers, such as genipin find applications in biomedical devices such as defibrillators. The effect of indifferent electrolytes on the rate of gelation of gelatin by dextranialdehyde has been reported in the literature. Hydrogel comprising gluteraldehyde cross-linked poly(vinyl alcohol)/poly(acrylic acid) blend has been evaluated as a coating on graphite electrodes for use in electrochemical sensors. Hydrogels have also found applications in enzyme-based electrochemistry.

The ECs employing aqueous polymer membranes such as nafion and sulfonated poly(ether ether ketone) (SPEEK) have been employed in EDLCs and pseudocapacitors. Ironically, the literature on ECs with hydrogel-polymer electrolytes is scanty. In this article, we review some hydrogel-electrolytes employed in ECs.
2. Organic hydrogel electrolytes

2.1. Poly(ethylene oxide) hydrogel electrolytes

An alkaline hydrogel electrolyte based on poly(ethylene oxide) (PEO), namely PEO–KOH–H₂O, has been employed as GPE-cum-separator in Ni-based batteries by various researchers. The disadvantages of this class of electrolytes include high crystallinity and low melting-point of PEO, limited operating temperature-range, and a low hydroxide-ion transference number coupled with poor interfacial characteristics. Vassal et al. have reported a family of alkaline hydrogels based on copolymers of poly(epichlorohydrin) and poly(ethylene oxide) with KOH as dopant and studied their use in nickel/metal hydride secondary battery and zinc/air primary batteries. The amorphous nature of poly(epichlorohydrin) combined with the solvating behavior of poly(ethylene oxide) is exploited in this study. The same hydrogel polymer electrolyte has been employed by Lewandowski et al. in an all solid-state EC that used activated carbon as the electrode material. They have reported the performance of PEO–KOH–H₂O and compared with 6 M KOH as electrolyte for ECs. Cyclic voltammograms (CV) reveal an ill-defined box-like shape and this behaviour is attributed to the low conductivity of the electrolyte. In the case of the PEO–KOH–H₂O electrolyte, the ionic transport is essentially due to the solution entrapped in the polymer matrix contributing to lower value of conductivity than that of a free-liquid electrolyte. The complex-impedance plot shows a semicircle at high frequencies with a transition to a linear part at low frequencies corresponding to a capacitive behavior. The resistance of the alkaline polymer-hydrogel-based EC as obtained from the impedance spectrum is found to be 0.77 Ω and the RC-time constant is observed to be 1.4 s. Specific capacitance values for the ECs employing the PEO–KOH–H₂O and 6 M KOH aqueous solution electrolytes are reported to be 93 and 90 F g⁻¹, respectively.

2.2. Potassium poly(acrylate) hydrogel electrolytes

Potassium poly(acrylate), with chemical formula –(CH₂–CHCOOR)ₙ–, dissolves in water with a concomitant decrease in pH of the resultant solution. Estimation based on the dissociation equilibrium constant of acetic acid (pKₐ = 4.76) or propanoic acid (pKₐ = 4.89) suggests that >95 % of H⁺ are ion-exchanged by K⁺ at the R-sites in the above chemical formula. PAAK-based hydrogels such as PAAK–KOH–H₂O and PAAK–KCl–H₂O could be classified as physical hydrogels as no chemical reaction is involved in their formation. Formation of PAAK-based physical hydrogels presumably takes place by immediate network formation making them more stable than the corresponding PEO–KOH–H₂O hydrogels. The degree of network formation depends upon the pH and ionic strength of the medium and also the type of counter ion and functional charge-density of the polymer. Ivakura et al. have prepared PAAK–KOH–H₂O hydrogel electrolyte and subsequently employed them in Ni/MH batteries. Ivakura et al. and Wada et al. have reported ECs employing alkali cross-linked potassium poly(acrylate) hydrogel, namely PAAK–KOH–H₂O, as electrolyte and compared the performance with an EC employing 10 M KOH aqueous electrolyte; the performances of the ECs are found to be similar. The ionic conductivity of the PAAK–KOH–H₂O hydrogel electrolyte is reported to be comparable to that of KOH aqueous solution electrolyte over wide ranges of KOH concentrations and temperatures while the PEO–KOH–H₂O hydrogel electrolyte exhibits low conductivity. The voltammograms for the ECs employing PAAK–KOH–H₂O and aqueous 10 M KOH electrolytes recorded at a scan rate of 10 mV s⁻¹ are shown in Fig. 3. Specific capacitances for the ECs, obtained from the CVs, are recorded at 25 °C and 144 F g⁻¹ with PAAK–KOH–H₂O and 10 M KOH aqueous electrolytes, respectively. The charge–discharge plots for both the ECs are observed to be nearly linear with little ohmic drop. This is ascribed to the high ionic-conductivity of the electrolytes and good electrical contact of the electrolytes with the carbon electrodes. The galvanostatic charge–discharge data for the ECs with PAAK–KOH–H₂O and 10 M KOH aqueous electrolytes recorded at a current density of 1 mA cm⁻² are shown in Fig. 4; the respective discharge capacitance values from these data are determined to be 146 and 142 F g⁻¹ for the ECs. The discharge capacitance of the EC with polymer hydrogel electrolyte is 82% of the initial value after 20 000 cycles, while it is 72% in the case of the EC with KOH aqueous solution electrolyte. This clearly indicates that EC assembled by using PAAK–KOH–H₂O hydrogel electrolyte exhibits better performance than the EC employing 10 M KOH aqueous solution electrolyte, suggesting
that the hydrogel polymer electrolyte prepared from PAAK with higher water holding capacity is more effective in maintaining good electrode/electrolyte interface than the aqueous alkaline electrolyte. To investigate the electrochemical behavior at the electrode/electrolyte interface in detail, ac impedance measurements have been carried out. The Nyquist plots for the ECs with cross-linked PAAK–KOH–H₂O and 10 M KOH aqueous solution electrolytes are shown in Fig. 5. The equivalent series resistance values are estimated to be 0.14 and 0.12 Ω for PAAK–KOH–H₂O and 10 M KOH aqueous solution electrolytes, respectively. Nohara et al.⁸⁹ have reported the self-discharge characteristics of an EC using PAAK–KOH–H₂O electrolyte. The EC has been found to exhibit lower voltage-decay on open circuit and lower leakage-current than the EC with KOH aqueous solution electrolyte. The same authors⁹⁰ have reported a hybrid EC assembled using activated carbon (AC) as negative electrode, Ni(OH)₂ as positive electrode and PAAK–KOH–H₂O as the electrolyte. The hybrid EC has been shown to operate over a wide potential range exhibiting ca. 2.4 times higher capacitance than the EC with two symmetrical AC electrodes.

A PAAK-based neutral hydrogel electrolyte has been studied by Lee and Wu.⁸² A pseudocapacitor with MnO₂·nH₂O as the electrode and PAAK–KCl–H₂O as the electrolyte has been assembled and its performance compared with a similar EC employing 1 M KCl aqueous solution as electrolyte. The PAAK–KCl–H₂O hydrogel electrolyte along with a partially immersed MnO₂·nH₂O-based EC in a container in an inverted position is shown in Fig. 6. The CVs for the MnO₂·nH₂O-based ECs with 1 M KCl aqueous solution and PAAK–KCl–H₂O hydrogel electrolytes are shown in Fig. 7(i) and Fig. 7(ii), respectively. Specific capacitance values of 168 and 116 F g⁻¹ have been observed for the ECs with PAAK–KCl–H₂O hydrogel and 1 M KCl aqueous solution electrolytes, respectively. The higher capacitance value for the hydrogel-based EC as compared to the EC employing aqueous KCl solution has been ascribed to the compatibility in the hydrophobicities of PVdF binder and PAAK. The hydrophobic nature of the polymeric component of the hydrogel electrolyte improves the compatibility between the PVdF binder component of the electrode matrix and the hydrogel electrolyte, resulting in good penetrability of the latter within the pores of electrodes. Consequently, the overall cell capacitance increases due to a decrease in the unwetted electrode area. Nyquist plots for the MnO₂·nH₂O-based ECs with PAAK–KCl–H₂O hydrogel and 1 M KCl aqueous solution electrolytes are shown in Fig. 8. These plots exhibit characteristics of a porous pseudocapacitor. The similarity in charge-transfer resistances for the two electrolytes-based ECs suggests that the electrochemical processes at the oxide–electrolyte interface are comparable in the two types of electrolytes. Hence, PAAK-based electrolytes seem to be good candidates to be used in conjunction with other electrode materials such as RuO₂·xH₂O, carbon, MnO₂ for pseudocapacitors as well as hybrid capacitors.

2.3. Poly(vinyl alcohol) hydrogel electrolytes

An EC employing an activated carbon fiber cloth electrode and a hydrogel electrolyte comprising aqueous PVA cross-linked
with glutaraldehyde has been reported by Wada et al.\textsuperscript{91} and its performance compared with a similar EC employing 1 M H$_2$SO$_4$ aqueous solution electrolyte. The ionic conductivity of the cross-linked PVA hydrogel electrolyte (PHE) has been found to be comparable to that of 1 M H$_2$SO$_4$ aqueous solution. The galvanostatic charge–discharge data for the ECs with cross-linked PHE and 1 M H$_2$SO$_4$ aqueous solution electrolyte are shown in Fig. 9. Discharge capacitance values of 110 and 115 F g$^{-1}$ have been found for the ECs with cross-linked PVA hydrogel electrolyte (PHE) and 1 M H$_2$SO$_4$ aqueous solution electrolyte, respectively. To monitor the self-discharge characteristics, the ECs are initially charged to 0.8 V and the time course of the cell voltage under an open circuit condition is recorded. The open circuit voltage–time decay data for the ECs with cross-linked PHE and 1 M H$_2$SO$_4$ aqueous solution electrolyte are shown in Fig. 10. As seen from Fig. 10, the open-circuit voltage–time decay for the cross-linked PHE-based EC is lower than that for the EC with 1 M H$_2$SO$_4$ aqueous solution electrolyte. This implies that the PHE-based EC exhibits lower self-discharge than the EC employing 1 M H$_2$SO$_4$ aqueous solution as electrolyte.

Studies on acidic, alkaline and neutral poly(vinyl alcohol)/poly(acrylic acid) blend hydrogel electrolytes (BHEs) and their application to ECs have been reported by Choudhury et al.\textsuperscript{54} Ionic conductivity data as a function of polymer composition for the PVA/PAA BHE with: (a) HClO$_4$, (b) NaOH, and (c) NaCl dopant are shown in Fig. 11. The ionic conductivity of the acidic PVA/PAA BHEs decreases with increase in the proportion of PAA in the blend while that of the alkaline and neutral PVA/PAA BHEs exhibit a reverse effect. This contrasting behavior has been explained based on the assumption that proton conduction in the acidic PVA/PAA BHE occurs by the Grothuss-type mechanism whereas the ionic conductivity in alkaline as well as neutral PVA/PAA BHEs is brought about predominantly by segmental motion.

The cyclic voltammograms for the ECs with black pearl carbon (BPC) electrodes and PVA/PAA BHEs comprising: (a) PVA : PAA = 5 : 0, (b) PVA : PAA = 4 : 1, and (c) PVA : PAA = 3 : 2 by volume, in (i) acidic, (ii) alkaline, and (iii) neutral forms
are shown in Fig. 12. With increasing proportion of PAA in the blend, the specific capacitance values for the acidic BHE-based ECs decrease while those for the alkaline and neutral BHE-based ECs increase. The variation in specific capacitance values with polymer composition for the ECs employing acidic, alkaline and neutral BHEs follow the same trend as that observed in the variation of ionic conductivity with polymer composition for the three types of BHEs.

To investigate the effect of variation of HClO₄ dopant concentration on specific capacitance of the ECs, CVs have been recorded for ECs comprising BPC electrodes and PHEs with 0.1, 0.5, 1.0, 1.5 and 2.0 N HClO₄ dopants and the data are shown as (a), (b), (c), (d) and (e) in Fig. 13, respectively. The specific capacitance value at 0 V increases from 13.4 to 95.6 F g⁻¹ as the concentration of HClO₄ in the PHE is increased from 0.1 to 2.0 N. The increase in specific capacitance with increasing dopant concentration is explained on the basis of increased accessibility of the electrode surface by increased amount of dopant ions during potential cycling. In order to probe the effect of variation in HClO₄ concentration on phase angle, electrochemical impedance data have been recorded for ECs comprising BPC electrodes and PHEs with 0.1, 0.5, 1.0, 1.5 and 2.0 N HClO₄ dopants and the data are shown as (a), (b), (c), (d) and (e) in Fig. 14, respectively. The phase angle at 1 Hz increases from 76° to 79° as the concentration of HClO₄ in the PHE is increased from 0.1 to 2.0 N. The increase in phase angle with increasing dopant concentration suggests an improvement in the capacitive behavior of the ECs. To study the effect of variation in HClO₄ concentration on discharge capacitance and coulombic efficiency, galvanostatic chronopotentiometric data for the ECs with PHEs containing 0.1, 0.5, 1.0, 1.5 and 2.0 N HClO₄ have been recorded and the data are shown as (a), (b), (c), (d) and (e), in Fig. 15, respectively. An increase in both charge and discharge times is observed with increasing HClO₄ concentration in the PHE. The unequal time-scales associated with the first few cycles of the charge and discharge processes for ECs need to be probed further. The discharge capacitance increases from 8.7 to 96.8 F g⁻¹ as the concentration of HClO₄ in the PHE is increased.
from 0.1 to 2.0 N. As the concentration of HClO₄ in the PHE is increased from 0.1 to 1.5 N, the coulombic efficiency of the ECs is found to increase from 85.9 to 87.5 %. With further increase in concentration of HClO₄ in the PHE to 2.0 N, the coulombic efficiency decreases to 76.5 %. The data pertaining to the electrochemical studies depicted in Fig. 13–15 are summarized in Table 1.

The PVA/PAA based materials are also amenable for fabricating pseudocapacitors using transition metal oxides. Capacitors employing RuOₓ·xH₂O/C electrodes and acidic PVA/PAA BHEs with varying polymer composition show voltammograms [(a) PVA : PAA = 5 : 0, (b) PVA : PAA = 4 : 1, and (c) PVA : PAA = 3 : 2 by volume, Fig. 16] that are nearly rectangular in shape. A decrease in specific capacitance with increasing proportion of PAA in the blend is observed. The variation in specific capacitance of the RuOₓ·xH₂O/C electrode-based ECs employing acidic BHEs follows the same trend as that observed for the ionic conductivity of the acidic BHEs as a function of polymer composition in the blend. A maximum specific capacitance value of ca. 1000 F g⁻¹ is obtained for the pristine acidic PHE. Electrochemical impedance data for a typical EC with RuOₓ·xH₂O/C electrodes and acidic pristine PVA hydrogel electrolyte recorded in the frequency range between 5 mHz and 10 kHz with a perturbation of 5 mV rms show a charge-transfer resistance of ca. 32 Ω and a phase angle of ca. 82°.

### Table 1

<table>
<thead>
<tr>
<th>HClO₄ concentration/M</th>
<th>Electrochemical data obtained from CVs at 0 V</th>
<th>CVs by integrating total enclosed area</th>
<th>Bode plot at 1 Hz</th>
<th>Galvanostatic chronopotentiometry at 0.5 mA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Specific capacitance/F g⁻¹</td>
<td>Average specific capacitance/F g⁻¹</td>
<td>Phase angle⁰</td>
<td>Specific capacitance/F g⁻¹</td>
</tr>
<tr>
<td>0.1</td>
<td>13.4</td>
<td>12.8</td>
<td>76.3</td>
<td>8.7</td>
</tr>
<tr>
<td>0.5</td>
<td>19.3</td>
<td>18.0</td>
<td>77.9</td>
<td>11.1</td>
</tr>
<tr>
<td>1.0</td>
<td>27.1</td>
<td>26.5</td>
<td>77.6</td>
<td>25.2</td>
</tr>
<tr>
<td>1.5</td>
<td>38.7</td>
<td>38.5</td>
<td>82.0</td>
<td>59.3</td>
</tr>
<tr>
<td>2.0</td>
<td>95.6</td>
<td>91.2</td>
<td>78.5</td>
<td>96.8</td>
</tr>
</tbody>
</table>

2.4. Gelatin hydrogel electrolytes

Gelatin is a polypeptide consisting mostly of proline, hydroxyproline, and glycine. It is extracted from denatured collagen. It dissolves in boiling water to form a pale yellow, semi-transparent, viscous solution. On cooling below 308 K, a concentrated solution of gelatin forms a physical hydrogel. The utility of such a hydrogel is limited due to its low network rigidity. The mechanical strength of gelatin hydrogel is improved by chemical cross-linking with aldehydes. Choudhury et al. have reported
the use of a hydrogel electrolyte comprising aqueous gelatin cross-linked with glutaraldehyde in ECs. The effect of variation in concentration of NaCl dopant on specific capacitance has been followed using cyclic voltammetry. Fig. 18 shows that the specific capacitance increases from 37.6 to 81.2 F g\(^{-1}\) as the concentration of NaCl in the GHEs is increased from 0 to 3 N. The increase in capacitance with increasing dopant concentration has been explained on the basis of increased accessibility of BPC by dopant ions during charge-discharge cycling. The low concentration of ions present in pristine GHE is not sufficient to access all the active sites of BPC that possess high surface-area. With increasing dopant concentration, there is a corresponding increase in the number of ions available for electric-double-layer formation at the electrode/electrolyte interface, leading to a better utilization of BPC electrode material in the ECs with doped GHEs as compared to pristine GHE-based ECs. The enhanced utilization of electrode material in the ECs with doped GHEs is responsible for enhanced specific capacitance of the latter. The effect of variation in concentration of NaCl on phase angle of ECs comprising BPC electrodes and GHEs as studied by electrochemical impedance is shown in Fig. 19. The phase angle of ECs at 1 Hz increases from 49 to 75° as the concentration of NaCl in the GHE is increased from 0 to 3 N. The charge–discharge data as recorded by galvanostatic chronopotentiometry for GHEs containing varying concentrations of NaCl are shown in Fig. 20. The discharge capacitance increases from 12.4 to 70 F g\(^{-1}\) as the concentration of NaCl in the GHE is increased from 0 to 3 N. The data pertaining to the electrochemical studies depicted in Fig. 18–20 are summarized in Table 2.

The discharge capacitance, coulombic efficiency and cell resistance data for the ECs comprising pristine as well as 3 N NaCl-doped GHEs are studied as a function of charge–discharge cycles and the data\(^9\) are shown in Fig. 21. As shown in Fig. 21, the EC with 3 N NaCl-doped GHE has a higher discharge-capacitance of 10.4 F g\(^{-1}\) as against 5.2 F g\(^{-1}\) observed for pristine GHE-based EC. As shown in Fig. 22, the coulombic efficiencies for the ECs with both pristine and 3 N NaCl-doped GHEs are as low as ca. 50% for the first charge–discharge cycle, which gradually increase to ca. 85% for the one hundredth cycle. The EC with 3 N NaCl-doped GHE has nominally higher coulombic efficiency as compared to the EC with pristine GHE. As shown in Table 3, the coulombic efficiency increases from 52 to 100% as the current density increases from 0.1 to 1.0 mA cm\(^{-2}\). As the current density increases, the time required to attain the cut-off voltage of 1 V decreases affecting the formation of the electric double-layer leading to an increase in coulombic efficiency of the ECs.\(^{94}\) As shown in Fig. 23, the cell resistance for the EC with 3 N NaCl doped GHE is ca. 1.2 Ω as compared to ca. 36 Ω for the EC with pristine GHE before recording charge–discharge cycles. The lower cell resistance for the doped GHE-based EC as compared to the pristine GHE-based EC is mainly due to the higher ionic conductivity for the doped GHE in relation to the pristine GHE. The discharge capacitance and coulombic efficiency decrease while cell resistance increases with increasing charge–discharge cycle number. The EC with 3 N NaCl doped GHE exhibits a longer life of ca. 4.7 h over 900 charge–discharge cycles as against the EC with pristine GHE that has an endurance of only ca. 4.3 h over 1700 cycles.

**Fig. 18** Cyclic voltammograms recorded at a scan rate of 5 mV s\(^{-1}\) for the ECs comprising BPC electrodes and GHEs containing (a) 0, (b) 1, (c) 2, and (d) 3 N NaCl dopants (from ref. 93).

**Fig. 19** Electrochemical impedance data recorded in the frequency range between 1 Hz and 10 kHz at an ac amplitude of 5 mV rms for the ECs comprising BPC electrodes and GHEs with (a) 0, (b) 1, (c) 2, and (d) 3 N NaCl dopants (from ref. 93).

**Fig. 20** Galvanostatic chronopotentiometric data recorded at a current density of 0.1 mA cm\(^{-2}\) for the ECs comprising BPC electrodes and GHEs containing (a) 0, (b) 1, (c) 2, and (d) 3 N NaCl dopants (from ref. 93).
Ionic conduction in pristine GHE takes place by the Gro¨tthus-type mechanism of proton transport whereas that in NaCl-doped GHEs occurs predominantly by free diffusion of hydrated Na⁺ and Cl⁻ ions. As described in the literature,

transport of protons in aqueous media by a Gro¨tthus-type mechanism takes place along chains of hydrogen bonded water molecules. In this mechanism, protons are transported by alternation of covalent and hydrogen bonds with little physical movement. This mechanism explains the facile proton transport and accounts for its highest conductivity among all the ions.

Na⁺ and Cl⁻ ions in aqueous media such as doped GHEs are associated with large hydration-shells. During charge–discharge cycling of ECs, Na⁺ and Cl⁻ ions with their large shells of hydrated water molecules shuttle back and forth between the electrodes. The ionic shuttling in ECs takes place at high rates owing to their high power-density. Because of the fast potential-driven migration of hydrated Na⁺ and Cl⁻ ions back and forth between the electrodes during charge–discharge cycling, the NaCl-doped GHEs degrade faster than pristine GHEs with a consequent sharp increase in cell resistance for the NaCl-doped GHE-based ECs as shown in Fig. 23. The decrease in both the discharge capacitance and coulombic efficiency values with concomitant increase in cell resistance for the ECs during charge–discharge cycling is due to the gradual evaporation of water from the GHEs. This can be substantiated from the study of the EC performance as a function of time during which the EC is kept under open-circuit condition in a closed chamber maintained at 100% relative humidity. There is only a small decrease in discharge capacitance during the first 50 h of operation subsequent to which the performance is almost

### Table 2

<table>
<thead>
<tr>
<th>NaCl concentration/N</th>
<th>CVs at 0 V</th>
<th>CVs by integrating total enclosed area</th>
<th>Bode plot at 1 Hz</th>
<th>Galvanostatic chronopotentiometry at 0.1 mA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Specific capacitance/F g⁻¹</td>
<td>Average specific capacitance/F g⁻¹</td>
<td>Phase angle/°</td>
<td>Specific capacitance/F g⁻¹</td>
</tr>
<tr>
<td>0</td>
<td>37.6</td>
<td>35.6</td>
<td>49</td>
<td>12.4</td>
</tr>
<tr>
<td>1</td>
<td>58</td>
<td>58.8</td>
<td>65</td>
<td>31.2</td>
</tr>
<tr>
<td>2</td>
<td>70.4</td>
<td>69.6</td>
<td>71</td>
<td>56.8</td>
</tr>
<tr>
<td>3</td>
<td>81.2</td>
<td>82.4</td>
<td>75</td>
<td>70</td>
</tr>
</tbody>
</table>

**Fig. 21** Plots of specific capacitance, evaluated from galvanstatic chronopotentiometric data recorded at a current density of 0.1 mA cm⁻², as a function of charge–discharge cycles for the ECs comprising BPC electrodes and GHEs containing (a) 0 and (b) 3 N NaCl (from ref. 93).

**Fig. 22** Plots of coulombic efficiency, evaluated from galvanstatic chronopotentiometric data recorded at a current density of 0.1 mA cm⁻², as a function of charge–discharge cycles for the ECs with GHEs containing (a) 0 and (b) 3 N NaCl (from ref. 93).

**Fig. 23** Plots of cell resistance, evaluated from electrochemical impedance data recorded in the frequency range between 100 mHz and 10 kHz with a perturbation of 5 mV rms, as a function of charge–discharge cycles for the ECs with GHEs containing (a) 0 and (b) 3 N NaCl (from ref. 93).
invariable. The capacitive performance data for the pristine GHE-based EC for 10 days of continuous operation are summarized in Table 3. It is noteworthy that unlike the synthetic-polymer-based hydrogel electrolytes, biopolymer-based hydrogel electrolytes with natural cross-linkers, such as genipin,\textsuperscript{39} have a unique advantage for applications in biomedical devices such as defibrillators.\textsuperscript{9} They are also presently shown to exhibit excellent performance in electrochemical energy systems.

### 3. Inorganic hydrogel electrolytes

Inorganic hydrogels prepared by the sol–gel method are one of the most promising host materials to design high proton-conducting composites. Unlike organic polymer-based hydrogels, silica hydrogels incorporating perchloric acid and heteropoly acid dopants have been found to possess ionic conductivity values as high as $10^{-2}$ S cm$^{-1}$ under ambient conditions as they contain a large number of micropores and mesopores filled with liquid which facilitate fast proton-transport. Perchloric acid-doped silica hydrogels containing PVA as a binder have been prepared by a sol–gel method and employed\textsuperscript{95} in all solid-state ECs along with activated carbon as the electrode. The dependence of room temperature ionic conductivity for the silica hydrogel electrolyte on HClO$_4$ concentration is shown in Fig. 24. The conductivity of the silica hydrogel electrolyte increases with increasing HClO$_4$ concentration attaining the maximum value of $5 \times 10^{-2}$ S cm$^{-1}$ at room temperature for a HClO$_4$/TEOS molar ratio of 1. The variation of IR-drop with varying discharge current for an all solid-state EC fabricated employing silica-hydrogel electrolyte with an HClO$_4$/TEOS molar ratio of 0.5 is shown in Fig. 25. The IR-drop has been found to be almost directly proportional to the discharge current. Variation in specific capacitance for the ECs comprising PVA-containing silica hydrogel electrolytes as a function of HClO$_4$ concentration is depicted in Fig. 26. The room-temperature specific capacitance of the EC drastically increases with an increase in HClO$_4$ concentration and tends to attain a value of 44 F g$^{-1}$ for a HClO$_4$/TEOS molar ratio of 0.5. The large capacitance of the all solid-state ECs is attributed to the high proton conductivity of the hydrogel electrolyte and the homogeneous hybridization of the activated carbon and the hydrogels.

### 4. Summary

Hydrogel-electrolyte-based all solid-state ECs are promising and environment-friendly energy-storage devices. Among the various alkaline polymer-hydrogel-electrolyte-based ECs, cross-linked PAAK–KOH–H$_2$O based-EC with activated carbon fiber cloth electrodes exhibits a specific capacitance value as high as

<table>
<thead>
<tr>
<th>Time/day</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average specific capacitance/F g$^{-1}$</td>
<td>43.6</td>
<td>40.8</td>
<td>37.6</td>
<td>36.4</td>
<td>36.0</td>
<td>36.8</td>
<td>36.8</td>
<td>37.6</td>
<td>37.6</td>
<td>37.6</td>
<td>38.0</td>
</tr>
</tbody>
</table>

---

**Table 3** Average specific capacitance values for pristine GHE-based ECs evaluated by keeping it in a closed chamber maintained at 100% relative humidity (ref. 93)

---

**Fig. 24** Variation in ionic conductivity of PVA-containing silica hydrogel electrolytes as a function of HClO$_4$ dopant concentration (from ref. 95).

**Fig. 25** Variation in IR-drop with discharge current for an EC employing PVA-containing silica hydrogel electrolyte with a typical HClO$_4$/TEOS molar ratio of 0.5 and activated carbon powder hybridized with silica hydrogel as polarizable electrode (from ref. 95).

**Fig. 26** Variation in specific capacitance for ECs comprising PVA-containing silica hydrogel electrolytes with varying HClO$_4$ dopant concentrations and activated carbon powder hybridized with silica hydrogel as polarizable electrode (from ref. 95).
150 F g⁻¹. A pseudocapacitor employing a MnO₂-nH₂O electrode and a neutral hydrogel-polymer-electrolyte, namely PAAK–KCl–H₂O, yields higher specific capacitance values of ca. 168 F g⁻¹. ECs with chemical hydrogel electrolytes, namely cross-linked PVA and gelatin-based hydrogel electrolytes, are both mechanically and chemically very stable as compared to ECs with conventional non-cross-linked hydrogel-polymer-electrolytes and hence are very promising. ECs employing acidic cross-linked PVA hydrogel electrolyte and activated carbon fiber cloth electrodes remain limited to specific capacitance values of only about 110 F g⁻¹. However, optimization of various physical parameters is desired to achieve optimum specific capacitance values with these ECs. Interestingly, cross-linked PVA hydrogel electrolytes can be prepared with acidic, alkaline and neutral dopants. BHEs consisting of PVA/PAA BHEs show interesting behavior in relation to polymer composition. With increasing proportion of PAA in the PVA/PAA BHEs, both ionic conductivity and specific capacitance values for the acidic BHE-based ECs decrease while those for the alkaline and neutral BHE-based ECs increase. A pseudocapacitor employing RuOₓ-xH₂O/C electrode and acidic pristine PVA hydrogel electrolyte exhibits a specific capacitance value as high as 1000 F g⁻¹. Biopolymer-based hydrogel electrolytes such as cross-linked gelatin hydrogel electrolytes are highly hydrophilic in nature and possess a better water-retention capacity as compared to synthetic polymers such as PVA-based hydrogel electrolytes. Silica hydrogel-based inorganic hydrogel electrolytes are attractive materials for application in ECs. Silica-based hydrogel electrolytes have been found to exhibit ionic conductivity values as high as 10⁻² S cm⁻¹ and specific capacitance values as high as 44 F g⁻¹ at room temperature. The use of physical hydrogels, such as gelatin, agar, starch, collagen, and gelatin gum, and also chemical hydrogels, namely chitosan, polyether, and polyamide, as electrolytes in ECs needs to be explored. Other hydrogels based on organic compounds such as bile acids are to be explored for electrochemical applications, particularly capacitors. In this article, we have limited the discussion only to the use of polymeric materials as electrolytes for ECs. It is noteworthy that the use of polymeric materials as electrodes in ECs is well documented in the literature.

References

4 A. Burke, J. Power Sources, 2000, 91, 37.
33 A. Bide and K. Harirahan, J. Power Sources, 2006, 159, 1450.