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Ce–Mn bimetallic oxide-doped SPEEK/SPPO blend composite membranes to induce high oxidative tolerance and proton conductivity for hydrogen fuel cells†

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Utilizing the potential of transition metals to quench free radicals in hydrogen fuel cells, we prepared cerium–manganese-based bimetallic oxide (CeMnO_x) nanostructures by alkali-aided deposition precipitation to harness as fillers in hydrocarbon-based polymer electrolyte membranes (PEMs). To improve the stability of the PEMs, a blend of sulfonated poly ether ether ketone (SPEEK) and sulfonated poly(2,6-dimethyl-1,4-phenylene oxide) (SPPO) with different weight ratios was employed as a membrane backbone. The interfacial adhesion and coupling configuration of three-dimensional CeMnO_x with sulfonic moieties of blend polyelectrolytes enhances the bound water content in the PEMs. It also constructs an extensive hydrogen bonding network with proton transport channels, uplifting the proton conductivity (K^m) by promoting vehicular and Grotthuss-type transport. Reversible redox cycling of Ce³⁺–Ce⁴⁺ and Mn²⁺–Mn³⁺ endorses the quenching of harsh radicals and augments the oxidative stability of the membranes. Composite PEMs exhibit only 4–9% mass loss and 2–5% K^m loss after exposure to Fenton's solution. SSM-73 membrane showed a peak power density of 431.2 mW cm⁻² with a maximum current density of 1272.6 mA cm⁻² at 75 °C in 100% RH and showed a high OCV retention of 88% over 50 h. The fuel cell performance of SSM-73 was 50% higher than that of its corresponding blend membrane, *i.e.*, SS-73, which showed a peak power density of 287 mW cm⁻² and a maximum current density of 872.3 mA cm⁻².

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1. Introduction

In spite of advancements in the modern world, fossil fuels remain the primary source of global energy consumption.^{1,2} As a result, a conundrum has emerged, necessitating the development of environmentally benign alternative energy sources. Proton exchange membrane fuel cells (PEMFCs) have attracted substantial attention in the energy sector due to their high energy conversion efficiency, eco-friendly operation, zero emissions, fast refueling, and scalability.^{3,4} PEMFCs are membrane-based electrochemical devices that convert chemical energy into electrical energy. Although redox flow batteries (RFBs) function the same way, there is continuous supply of fuel to fuel cells in PEMFCs, enabling them to deliver uninterrupted electrical energy, unlike charge–discharge in RFBs.⁵

Polymer electrolyte membranes (PEMs) composed of a polymeric backbone integrated with charged functional groups are the most crucial components in fuel cells as they allow for preferential transfer of ions, enable circuit completion, and prevent mixing of reactants inside the fuel cell device.⁶ Nafion, a perfluorosulfonic acid polymeric membrane, is widely recognized as the state-of-the-art PEM and has achieved commercial success due to its extraordinarily high hydrated conductivity and mechanical and chemical stability.⁷ However, at elevated temperatures and low humidity, Nafion suffers from shortcomings such as poor proton conductivity and degraded mechanical stability.^{7,8} Moreover, the complex synthetic process and high cost of the Nafion membrane challenge scientists to develop alternative, cost-effective, and more robust PEMs.

Aromatic non-fluorinated PEMs, basically sulfonated derivatives of poly(arylene ethers), have attracted the attention of many researchers as potential PEMs for fuel cells due to their ease of sulfonation, controllable degree of sulfonation and high film-forming ability.⁹ The use of sulfonated poly ether ether ketone (SPEEK),^{10,11} sulfonated polysulfone (SPSf),¹² sulfonated polyethersulfone (SPES),¹³ sulfonated poly(2,6-dimethyl-1,4-

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phenylene oxide) (SPPO),¹⁴ sulfonated polystyrene and its derivatives,¹⁵ and sulfonated polybenzimidazole¹⁶ as potential PEMs for PEMFCs has been reported so far. Among these PEMs, SPEEK and SPPO have drawn much attention due to their easy synthetic process, commercial availability, and low cost of poly ether ether ketone (PEEK) and poly(2,6-dimethyl-1,4-phenylene oxide) (PPO). Nevertheless, these sulfonated PEMs have weak chain mobility and average power density.¹⁷ A higher degree of sulfonation (DoS) is necessary to enhance ionic conductivity. However, the increased number of sulfonic acid groups elevates the ion exchange capacity (IEC), resulting in higher water uptake, which leads to dimensional instability.¹⁸ One more eminent hurdle of these hydrocarbon-based PEMs is their poor oxidative stability. These hydrocarbon-based PEMs are intrinsically prone to radical-induced membrane deterioration, leading to high H₂ crossover and low performance in PEMFCs.¹⁹ H₂ crossover induces an exothermic reaction between H₂ and O₂, possibly resulting in pinholes in the PEMs. Harshly, the chemical reactions at the cathode sides of the cell also craft ·OH, ·H, and ·OOH radicals, accountable for the chemical invasion on the membrane and catalyst layers.^{20,29} Hence, it is desirable to construct such PEMs that can overcome the above-mentioned limitations to achieve remarkable PEMFC performance.

Plentiful approaches such as blending SPEEK or SPPO with other polymers,^{21,22} crosslinking of SPEEK,²³ and grafting of PEEK/SPEEK²⁴ to enhance the chemical and mechanical stabilities have been reported. However, these techniques are inefficient in providing desirable current and power density. Among several methodologies, embedding inorganic fillers in the polymer matrix has proven to be a facile and impactful technique to enhance the membrane properties of PEMFCs.²⁵ Furthermore, if the integrating composite material is proton-conductive, it will trigger the overall conductivity of the membrane. It will produce additional nano pathways for proton conduction and also enhance the synergistic interaction between the polymeric backbone and the filler.²⁶ In the search for suitable fillers, transition metal-based doping is highly preferable due to their anti-oxidative attributes, which will mitigate the oxidative decomposition of the membranes.²⁷ There are reports using Ce, Zr, Mn, Fe, and Ti-based ions, oxides, MOFs, MXenes, or bimetallic oxides as intercalating agents into PEMs, resulting in improved chemical stability.^{7,28–36} Crown ethers, organic anti-oxidants, silica supports, and hetero-polyacids embedded in transition metal ions have also been investigated as fillers for the improved oxidative stability of PEMs.^{37–39} However, there are some shortcomings in doping fillers into a polymer matrix, such as introducing phase separation and agglomeration in the membrane.²⁶ Under certain conditions, it may also obstruct the sub-nano proton conduction channels, resulting in lower proton conductivity and PEMFC performance. Hence, it is essential to tune the filler loading to achieve accelerated conductivity, open circuit voltage (OCV), and power density.

Among the radicals generated during PEMFC operation, ·OH is specifically more harmful to non-fluorinated aromatic hydrocarbon-based membranes. ·OH attacks the aromatic

rings at a rate constant of 10⁹ to 10¹⁰ M⁻¹ s⁻¹, which is much higher than the kinetics with fluorinated membranes (approx. 10⁶ M⁻¹ s⁻¹).⁴⁰ Studies revealed that cerium (Ce) is a prominent free radical scavenger among transition metals and can also be treated as a repairing agent in sulfonated hydrocarbon-based polymeric membranes.¹⁹ The unique anti-oxidant and catalytic attributes of Ce nanoparticles are mainly due to the simultaneous existence of Ce³⁺ and Ce⁴⁺ ions and their reversible redox cycling between both oxidation states. Direct doping of CeO₂ in the polymer matrix will not result in better PEMFC performance due to the major existence of Ce⁴⁺ in CeO₂, while the main anti-oxidative property of Ce is due to its Ce³⁺ state.⁴¹ Hence, Ce nanoparticles with some structural modifications are preferable for doping in the PEMs. Similar to Ce, Mn also exhibits redox catalytic properties as well as anti-oxidant behavior. Both Ce³⁺ and Mn²⁺ ions can be treated as radical scavengers as they quench almost 89% and 46% of ·OH radicals, respectively when the doping is only about 1% with respect to the sulfonated hydrocarbon membrane.^{36,37} However, the hydrogen pre-coverage of Mn^{m+} ions is superior to that of Ceⁿ⁺, resulting in higher proton conduction. To achieve the optimal properties of these two transition metals, Sharma *et al.* synthesized and incorporated a CeMnO_x bimetallic oxide into a SPEEK polymer electrolyte to fabricate a stable PEM for enhanced performance and durability in PEMFCs and achieved a peak power density of 88 mW cm⁻² at 90 °C in 100% RH.³⁶ After several cycles of operation in PEMFCs, the anti-oxidative nature of Ce/Mn may suffer a little degradation at high RH.³⁷ Hence, it is only the SPEEK membrane that performs the electrochemical reaction under harsh conditions. Although SPEEK exhibits excellent physicochemical and electrochemical properties, it has some bottlenecks in terms of stability. SPPO exhibits slightly better mechanical, tensile, and oxidative stabilities than those of SPEEK. Hence, the SPEEK/SPPO blend polymer electrolyte may be chosen to achieve the most favorable combination of electrochemical properties and stability to construct PEMs. This combination may help resist the radical attack without compromising on the proton conductivity to generate high current and power densities in PEMFC applications.

Considering all the above-mentioned facts, we propose CeMnO_x bimetallic oxide-incorporated SPEEK/SPPO blend membranes to fabricate chemically stable PEMs for PEMFCs. The degree of sulfonation of SPPO was kept lower than that of SPEEK, to provide high oxidative tolerance in the blend membranes. A series of PEMs have been fabricated with different weight ratios of SPPO to SPEEK (70/30 w/w and 50/50 w/w, keeping a fixed CeMnO_x loading of 2% w/w). The impact of the CeMnO_x on the physicochemical and electrochemical properties has been thoroughly investigated. Our novelty lies in the blending chemistry and an optimized loading of CeMnO_x to produce optically transparent CeMnO_x@SPEEK/SPPO hybrid PEMs for enhanced performance and durability in hydrogen fuel cells. The present work comes with a wider scope and can beat the monopoly of perfluorinated membranes in the modern world of fuel cell technologies.

2. Experimental

2.1. Materials

PEEK (Solvay, KetaSpire® 820 P), PPO (Merck), sulfuric acid (H₂SO₄, Qualigens, 98%), chlorosulfonic acid (CSA, Research Lab, 97%), chloroform (CHCl₃, SRL Chemicals, 99%), dimethylacetamide (DMAc, Research Lab, 99.5%), dimethyl sulfoxide (DMSO, 99%, Finar), deuterated dimethyl sulfoxide (DMSO-d₆, 99.9%, TCI), ceric ammonium nitrate (CAN, Molychem, 98%), manganese dioxide (MnO₂, SRL Chemicals, 99%), sodium hydroxide (NaOH, Fisher Scientific, 97%), hydrogen peroxide (H₂O₂, Research Lab, 30%), nitric acid (HNO₃, Research Lab, 69%), ferrous chloride (FeCl₂·6H₂O, Loba Chemie, 97%), hydrochloric acid (HCl, Qualigens, 37%), sodium chloride (NaCl, Research Lab, 99%), and a Nafion 117-containing solution (Merck, 5% v/v) were purchased and used. Fuel cell accessories, including gas diffusion layer (GDL), platinum catalyst (DURA-lyst, 40% Pt on Vulcan XC72), and gaskets were obtained from Sainergy Fuel Cell India Pvt Ltd. Ultrapure water was used in all experimental techniques.

2.2. Synthetic procedures for materials and membranes

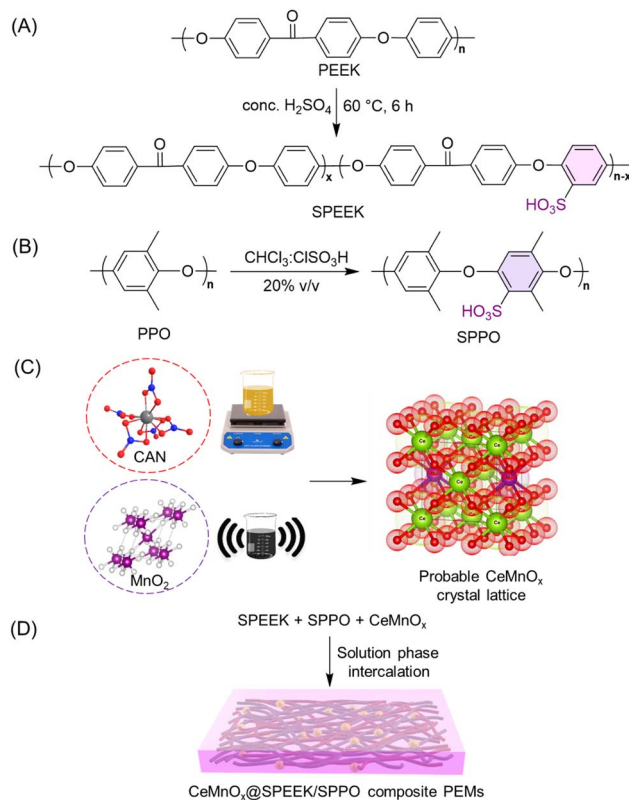
2.2.1. Synthesis of SPEEK fibers. PEEK was sulfonated using the simple chemistry of direct electrophilic aromatic substitution reaction, reported earlier.^{36,42} Briefly, 10 grams of PEEK was dissolved in 150 mL of concentrated H₂SO₄ (6.67% w/v) in a 500 mL beaker covered with a glass plate until a clean, transparent, and viscous reddish-orange solution was formed. Then the solution was stirred at 60 °C for 6 h. After that, the reaction mixture was removed from the stirrer and precipitated in cold water to obtain SPEEK fibers. These fibers were subsequently washed several times with water, followed by a final wash with methanol to achieve a neutral pH. Finally, the fibrous material was vacuum-dried at 60 °C for 12 h. Scheme 1A demonstrates the synthesis procedure of SPEEK fibers. The DoS was evaluated using two methods: (i) ¹H-nuclear magnetic resonance (NMR) spectroscopy of SPEEK fibers and (ii) ion exchange capacity (IEC) of the virgin SPEEK membranes.^{43,44} Eqn (1) and (2) represent the mathematical calculations used for evaluating DoS by ¹H NMR and IEC, respectively:

$$\frac{n}{12 - 2n} = \frac{I_c}{\sum I_{a,b,d,d',e,e'}} \quad (1)$$

$$\text{IEC}(\text{meq g}^{-1}) = \frac{\text{DoS} \times 1000}{M_{\text{PEEK}}(\text{g mol}^{-1}) + M_{\text{SO}_3\text{H}}(\text{g mol}^{-1}) \times \text{DoS}} \quad (2)$$

where I_x is the intensity of the signal ascribed to x proton and the DoS was calculated as $n \times 100$. M_{PEEK} and $M_{\text{SO}_3\text{H}}$ represent the molecular weight of PEEK and SO₃H single units, respectively.

2.2.2. Synthesis of the SPPO powder. The sulfonation of PPO was carried out by methods reported previously.^{14,45} PPO powders were dissolved in 100 mL of CHCl₃ (10% w/v) in a 250 mL round-bottom flask (RB). After complete dissolution, the SPPO/CHCl₃ solution was cooled down below 5 °C in an ice



Scheme 1 Synthesis procedure of (A) SPEEK, (B) SPPO, (C) CeMnO_x, and (D) composite PEMs.

bath. Then 35 mL of separately prepared CHCl₃/CSA (20% v/v) solution was added dropwise using a dropping funnel with vigorous stirring. After complete addition, the sulfonation reaction was allowed to proceed for a further 6 h. Subsequently, the resulting reddish-brown reaction mixture was precipitated in ice-cooled water, followed by multiple washing with water until a neutral pH was achieved. Finally, the material was dried in a vacuum oven at 60 °C for 12 h. Scheme 1B illustrates the chemical synthesis route of SPPO. The extent of functionalization was measured in terms of DoS, calculated from the ¹H NMR spectra of SPPO fibers and IEC of the SPPO membranes using eqn (3) and (4), respectively:^{46,47}

$$\text{DoS} = \frac{I_m}{0.5I_p + I_m} \quad (3)$$

$$\text{DoS} = \frac{120\text{IEC}}{1000 - 80\text{IEC}} \quad (4)$$

where I_m and I_p stand for the intensity of the aromatic protons from the substituted and unsubstituted units in SPPO, respectively; 120 and 80 are the molecular weights of the PPO and –SO₃H single units.

2.2.3. Synthesis of the CeMnO_x bimetallic oxide. The CeMnO_x bimetallic oxide was fabricated following an alkali-aided deposition precipitation procedure (Scheme 1C).³⁶ In brief, 2 g CAN was completely dissolved in 50 mL 0.1 M HNO₃ in a beaker. Simultaneously, in another beaker, 0.32 g of MnO₂

was dispersed in 50 mL ultrapure water using an ultrasonicator. Both the solutions were allowed to dissolve/disperse for a span of 6 h at 60 °C. Then, a yellow CAN/HNO₃ solution was added dropwise to the MnO₂ dispersion and kept under stirring overnight at 75 °C. Then, 1 M NaOH solution was added to the reaction mixture to achieve a pH range of 11–12. The addition of alkali caused precipitation in the reaction mixture. This precipitate solution was further stirred for 24 h, followed by 48 h of digestion. Finally, ultrapure water was added to the reaction mixture and centrifuged at 7000 rpm (5–6 times) to obtain a neutral pH (pH = 7). The final product was dried in a vacuum oven to obtain CeMnO_x bimetallic oxide nanoparticles.

2.2.4. Fabrication of composite blend PEMs. The PEMs were constructed by solution-phase blending of SPEEK and SPPO by varying their weight ratios. To begin, 0.7 g of SPEEK and 0.3 g of SPPO were dissolved in 10 mL and 5 mL of DMAc respectively, in separate beakers at 60 °C. After perfect dissolution, both solutions were mixed and stirred for 2 h at 60 °C. Simultaneously, 20 mg of CeMnO_x (2% w/w with respect to total polymer content) was dispersed in 2 mL of DMAc using an ultrasonicator. Finally, the dispersed CeMnO_x solution was added to the blend mixture with vigorous stirring to avoid any physical agglomeration of composite materials and kept for 2 h. Then, the CeMnO_x@SPPEK/SPPO casting solution was poured into a glass plate (10 cm × 17 cm) and kept under an IR lamp at 60 °C for 16 h for drying. After drying, the membrane was peeled out and kept in ultrapure water, followed by 0.1 M HCl for 6 h for functional group activation and then stored in ultrapure water. The resulting PEM was abbreviated as SSM-73 (which indicates a SPEEK/SPPO blend of 70 : 30 w/w with a filler loading of 2%, with the first 'S' representing SPEEK, the second 'S' representing SPPO, and M representing the filler material). A blend membrane without the incorporation of a CeMnO_x material was also fabricated to compare the PEMFC performance abbreviated as SS-73 (with first 'S' representing SPEEK, second 'S' representing SPPO, 7 corresponding to 70% weight content, and 3 corresponding to 30% weight content). Similarly, the other two PEMs were constructed by changing the SPEEK/SPPO content to 50 : 50 w/w and named SSM-55 (with filler) and SS-55 (without filler). A blank SPEEK membrane was also prepared for comparison purposes. Moreover, PEMs with different SPEEK/SPPO ratios (w/w) were also fabricated for comparison purposes (refer to Table S1†).

2.3. Characterizations

2.3.1. Structure, morphology, and phase. The structure of the sulfonated polymers was characterized by ¹H NMR spectroscopy. The functional groups present in sulfonated polymers and the prepared PEMs were evaluated by attenuated total reflection infrared (ATR-IR) spectroscopy analyses. The phases present in the CeMnO_x bimetallic oxides were characterized by wide-angle X-ray diffraction spectroscopy (WAXS). The vibrational modes of the composite materials were evaluated by Raman spectroscopy. The elements and chemical states present in CeMnO_x were analyzed by X-ray photoelectron spectroscopy (XPS). The surface morphology of the prepared PEMs and

CeMnO_x structure was characterized using a field emission scanning electron microscope (FE-SEM). Elemental mapping of CeMnO_x was performed using energy-dispersive X-ray (EDX) analysis. The phase separation and topology of the PEMs were studied using an atomic force microscope (AFM) and transmission electron microscope (TEM). Besides the phase of PEMs, TEM illuminates the lattice fringes present in the crystalline CeMnO_x bimetallic oxides. The co-continuous charge distribution was studied by dipping the sample membrane pieces in a methylene blue dye solution. The detailed procedure for all the analyses is given in the ESI.†

2.3.2. Stability. The overall stability of the PEMs was evaluated by analyzing their thermal, mechanical, tensile, and oxidative stabilities. The thermal stability of the PEMs was studied by thermogravimetric analysis (TGA). The glass-transition temperature (*T_g*) of the PEMs was studied by differential scanning calorimetry (DSC). The mechanical stability was evaluated using a dynamic mechanical analyzer (DMA). The tensile properties of the PEMs were investigated by their stress-strain analysis using a universal testing machine (UTM). Finally, the oxidative stability of the PEMs was investigated using Fenton's reagent. The detailed procedures for all the techniques are given in ESI.†

2.3.3. Physico/electrochemical properties. The water uptake (WU, %), swelling ratio (SR, %), hydration number (λ_{hyd}), fixed charge density (χ^m), and IEC of the PEMs were evaluated by the methods previously reported by us.⁴⁸ The ionic conductivity (proton conductivity, κ^m) of the PEMs was studied by electrochemical impedance spectroscopy (EIS) by constructing the Nyquist plot. The activation energy (*E_a*) for the membranes was calculated by evaluating their Arrhenius plots. The detailed explanations of all the methods and the employed equations are given in ESI.†

2.4. Membrane electrode assembly and PEMFC testing

The membrane electrode assembly (MEA) with a fuel cell stack of 16 cm² was fabricated using commercial Pt/C (40% w/w) as both the cathode and anode catalysts. The Pt catalyst was loaded on a commercially supplied gas diffusion layer (GDL, ML coated). In brief, 50 mg of Pt catalyst was finely dispersed in IPA using an ultrasonicator for 1 h. Subsequently, 30% Nafion 117-containing solution (5% v/v) was added to it and sonicated for further 30 min. The resulting catalyst slurry was coated onto the GDL (brush coating) to obtain a catalyst loading of 0.5 mg cm⁻² on both the GDLs. Now, a membrane piece of dimension 7.5 cm × 7.5 cm was sandwiched between the prepared electrodes by hot-pressing at 90 °C for 3 min at 30 kN cm⁻². Then, the resulting MEA was placed with Teflon gaskets and assembled in a fuel cell stack. The PEMFC performance of the PEMs was evaluated using a fuel cell test station (FCT) supplied by Fuel Cell Technologies, Inc., USA. H₂ and O₂ gases were allowed to pass through the anode and cathode at a flow rate of 60 and 150 mL min⁻¹, respectively, through a bubble humidifier. The PEMFC performance was evaluated at a cell temperature of 75 °C at 100% RH. The fuel cell testing was performed several times to get equilibrated data.

2.5. Durability test

The oxidative resistance and durability of the MEAs of the blend membranes (SS-73 and SS-55) and the composite PEMs (SSM-73 and SSM-75) were further investigated by the accelerated stress test (AST) with a slightly modified DoE protocol. The OCV of the single-cell PEMFC was evaluated as a function of time for 50 h. The cell temperature was maintained at 75 °C with a low humidity of 30% RH throughout the test. The anode compartment was fed with pure H₂ gas at 100 mL min⁻¹, and the cathode side was fed with O₂ gas at 150 mL min⁻¹. The OCV of the cell was measured using the same FCT. Moreover, the conductivity loss of the prepared PEMs after the durability test was also evaluated by EIS using CH Instruments, Inc. (CHI608E) in the fuel cell setup with the working electrode on the cathode side and the reference and counter electrodes on the anode side under similar conditions.

3. Results and discussion

3.1. Structure of SPEEK, SPPO, and CeMnO_x

The direct electrophilic substitution of hydrophilic sulfonic acid groups on the PEEK polymeric backbone was done at the *ortho* position to the aryl ether linkage rather than to the aryl keto linkage (Scheme 1). This is due to the presence of a higher electronic density at the aryl ether linkage, which is highly prone to electrophilic substitution.⁴³ The structure of SPEEK was determined by its ¹H NMR spectra using DMSO-d₆ as the solvent (Fig. 1). The single signal present at $\delta = 7.5$ ppm is due to the aromatic proton present next to the C-SO₃H linkage. This confirms the successful sulfonation of the PEEK backbone. Besides, multiplet signals at $\delta = 7.15$ and 7.25 ppm are ascertained to other hydrogen atoms present in the SPEEK backbone. The DoS was evaluated using the ¹H NMR spectra and calculated to be 64.65%, which can be correlated with the DoS obtained using IEC, which is 58%. Similarly, the synthesis of SPPO also follows a similar substitution mechanism. The addition of sulfonic acid moieties in the PPO backbone was confirmed by ¹H NMR spectroscopy using DMSO-d₆ as the solvent (Fig. 1, only

compound signals have been shown to avoid the merging of solvent and compound signals). The SPPO structure is mainly characterized by two prominent signals at $\delta = 6.43$ ppm (aromatic ring protons) and $\delta = 6.02$ ppm (protons parallel to sulfonic acid groups) and utilized to evaluate DoS, which comes out to be 24.13%. Moreover, a range of split signals from $\delta = 1.9$ –2.5 ppm can be ascribed to the benzylic protons in SPPO. These split signals are due to the change in the electronic environment near the benzylic protons by due to the effect of sulfonic acid groups in the SPPO polymeric backbone.⁴⁵

Further, the assessment of functional groups in SPEEK, SPPO, and blend PEMs was done by ATR-IR analysis (Fig. 2A and B). From the curves, the broad band near 3000–3600 cm⁻¹ is ascribed to the O–H stretching vibrations of the sulfonic acid groups present in SPEEK, SPPO, and blend PEMs. This peak also represents the O–H vibrations due to the moisture content absorbed by the hydrophilic –SO₃H segments.^{43,45} Higher moisture trapping due to the addition of a hydrophilic CeMnO_x material in the PEMs produces a broader transmission band in SSM-73 and SSM-55. Moreover, a band at ≈ 1647 cm⁻¹ is observed in SPEEK and the blend PEMs is ascertained to the existence of aromatic carbonyl groups in the SPEEK backbone, which is absent in SPPO. Additionally, all membranes exhibit bands at ≈ 1306 cm⁻¹, ≈ 1079 cm⁻¹, ≈ 1024 cm⁻¹, and ≈ 706 cm⁻¹ due to the C–H, O=S=O, S=O stretching and O=S=O bending vibrations, respectively.³⁶ In the filler-loaded PEMs, *i.e.*, SSM-73 and SSM-55, some weak bands are added at the fingerprint IR regions due to the transmission exhibited by the chemical bonds of CeMnO_x nanostructures. Some bands at 515 cm⁻¹, 523 cm⁻¹, 531 cm⁻¹, 550 cm⁻¹, and 600 cm⁻¹ can be attributed to the vibrational frequency of Ce–O and Mn–O bonds of the CeMnO_x crystal lattice.^{49,50} This confirms the successful incorporation of the CeMnO_x nanostructures in the PEMs.

The textural structure and phases of CeMnO_x were studied by WAXS analysis (Fig. 2C). The WAXS curve of the prepared material covered diffraction lines pertaining to CeO_x (cerianite, PDF 43-1002, JCPDS: 01-081-1984) phases at diffraction angles of $2\theta = 28.6^\circ$ (1 1 1), 33.2° (2 0 0), 47.6° (2 2 0), and 56.86° (3 1 1). Similarly, the diffraction pattern is assigned to the crystal planes of MnO_x (ramsdellite, PDF 44-142 and pyrolusite, PDF 24-735, JCPDS: 01-072-1984) phases at $2\theta = 37.78^\circ$, 41.2° , 43° , and 77.46° (3 0 1). It is worth noticing that the WAXS curve of the CeMnO_x nanostructure exhibits sharp diffraction peaks for the CeO_x phase ($2\theta = 28.6^\circ$, 33.1° , 47.5° , 56.6°), which signifies the existence of well-crystallized and ordered cerium oxides within it. Meanwhile, the diffraction peaks corresponding to MnO_x ($2\theta = 37.3^\circ$, 77°) are weaker than those of CeO_x, which indicates the abundance of Mn²⁺ in the highly dispersed solution with lower crystallinity.⁵¹ Hence, the overall crystallinity in the CeMnO_x nanostructures tends to decrease compared to CeO₂, promoting more defective crystallite sites. These defects manifest the increment of Ce³⁺ per total Ce ions (Ce_t), which is highly favorable for scavenging the harsh ·OH and ·OOH radicals.⁵² The crystallite size (D_{crystal}) of Ce phases was obtained using the Debye–Scherrer equation (eqn (S7), ESI[†]). The D_{crystal} value for the (1 1 1) plane of Ce (in CeMnO_x) was calculated to be

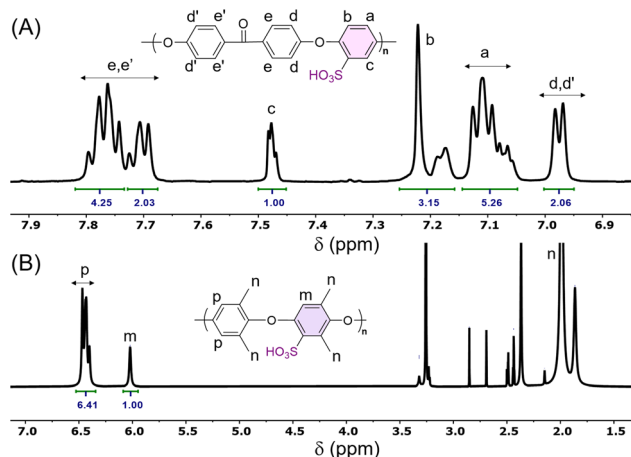


Fig. 1 ¹H NMR spectra of (A) SPEEK and (B) SPPO.

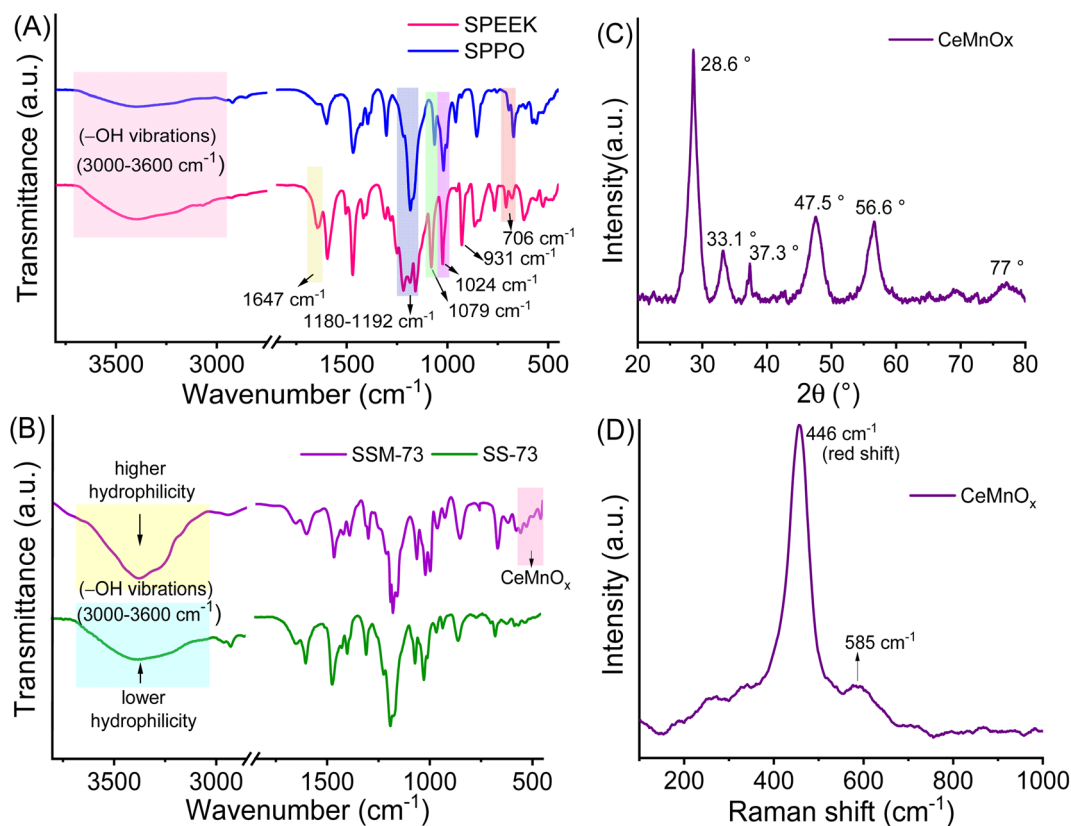


Fig. 2 ATR-IR spectra of (A) SPEEK and SPPO and (B) SSM-73 and SS-73 respectively. (C) and (D) XRD pattern and Raman spectra of CeMnO_x.

5.67 nm, whereas for CeO₂, it is 3.5 nm, indicating the enhancement of the surface area and volume of the crystal lattice in the CeMnO_x nanostructure. The vibrational modes of CeMnO_x were illustrated by its Raman spectrum analysis (Fig. 2D). The spectrum is primarily dominated by the vibrational band at 446 cm⁻¹ due to the fluorite structured degenerated F_{2g} Raman active mode (Ce–O–Ce symmetrical stretching), followed by a weak satellite band at 585 cm⁻¹ due to the defect-induced mode (D mode) of cerium oxides.^{14,53} The successful Mn doping in the crystal lattice leads to a red shift of F_{2g} Ce–O–Ce symmetrical stretching to 446 cm⁻¹ from 460 cm⁻¹, which is observed for CeO₂ nanoparticles.³⁰

The chemical valence states and bonds of Ceⁿ⁺ and Mn^{m+} in CeMnO_x were evaluated by XPS (Fig. S1, ESI† and Fig. 3). Fig. 3A displays the XPS profiles of Ceⁿ⁺ in their 3d_{5/2} and 3d_{3/2} spin states. The abundance of Ce³⁺ is ascertained to four deconvoluted peaks located at 882.1 eV (3d_{5/2}), 887.6 eV (3d_{5/2}), 900.6 eV (3d_{3/2}), and 905.2 eV (3d_{3/2}). Similarly, there are six peaks for Ce⁴⁺ 3d states, which are ascribed to the peaks present at 884.9 eV (3d_{5/2}), 889.8 eV (3d_{5/2}), 898.1 eV (3d_{5/2}), 902.8 eV (3d_{3/2}), 907.1 eV (3d_{3/2}), and 916.4 eV (3d_{3/2}). The Ce³⁺ content in the prepared CeMnO_x was evaluated using [Ce³⁺]/[Ce_T] ratio (from the XPS curves), which comes out to be 42%. It is worth noting that the alkali-assisted preparation strategy of the CeMnO_x nanostructure enhanced the abundance of Ce³⁺ ions, as evident from the XPS profiles. Fig. 3B represents the 2p_{1/2} and 2p_{3/2} profile of Mn^{m+} moieties. The presence of Mn²⁺, Mn³⁺, and Mn⁴⁺

states in the CeMnO_x samples is evident from the deconvoluted peaks assigned precisely to binding energies of 639.8 eV, 641.1 eV, and 642.5 eV, respectively. A weak band at 645.8 can be ascribed to Mn^{m+} oxides (Mn^{δ+}–O^{δ-}) in the nanostructure framework.⁵¹ Additionally, three types of O 1s chemical states are observed in the XPS spectrum. The lattice oxygen state (O_{latt}) is located at 529.1 eV. The adsorbed oxygen state (O_{ad}), such as O₂⁻ or O⁻ is present at 530.6 eV. These O_{ad} values are also responsible for the defects creating oxygen vacancies.⁵⁴ Finally, hydrophilic adsorbed oxygen (O_{hyd}) such as water or hydroxyl moieties (H₂O or HO⁻) is assigned at a binding energy of 531.2 eV (Fig. 3C). Generally, the abundance of Ce³⁺ states in a nanostructure promotes oxygen vacancies.⁵⁵ This implies that the prepared CeMnO_x nanostructures contain desirable oxygen vacancies and promote ion conduction.

The morphology of the prepared PEMs was studied by their cross-sectional FE-SEM analysis (Fig. S2, ESI† and Fig. 4A and B). The SEM images illustrated featureless morphology with a smooth surface without cracks or pinholes. Hence, the PEMs are dense and can reduce the fuel crossover in PEMFCs. Fig. 4A, B and S3, ESI,† manifest the FE-SEM images of the prepared nanostructures. The shapes of CeMnO_x nanostructures look like a fusion of octahedron-cubic structures. The EDX elemental mapping further presents the distribution of Ce, Mn, and O segments in the composite material (Fig. 4C–F and S4, ESI†). The [Ce]/[Mn] ratio was calculated to be 14/1, indicating that the desired quantity of Mn is perfectly coupled with Ce to impart

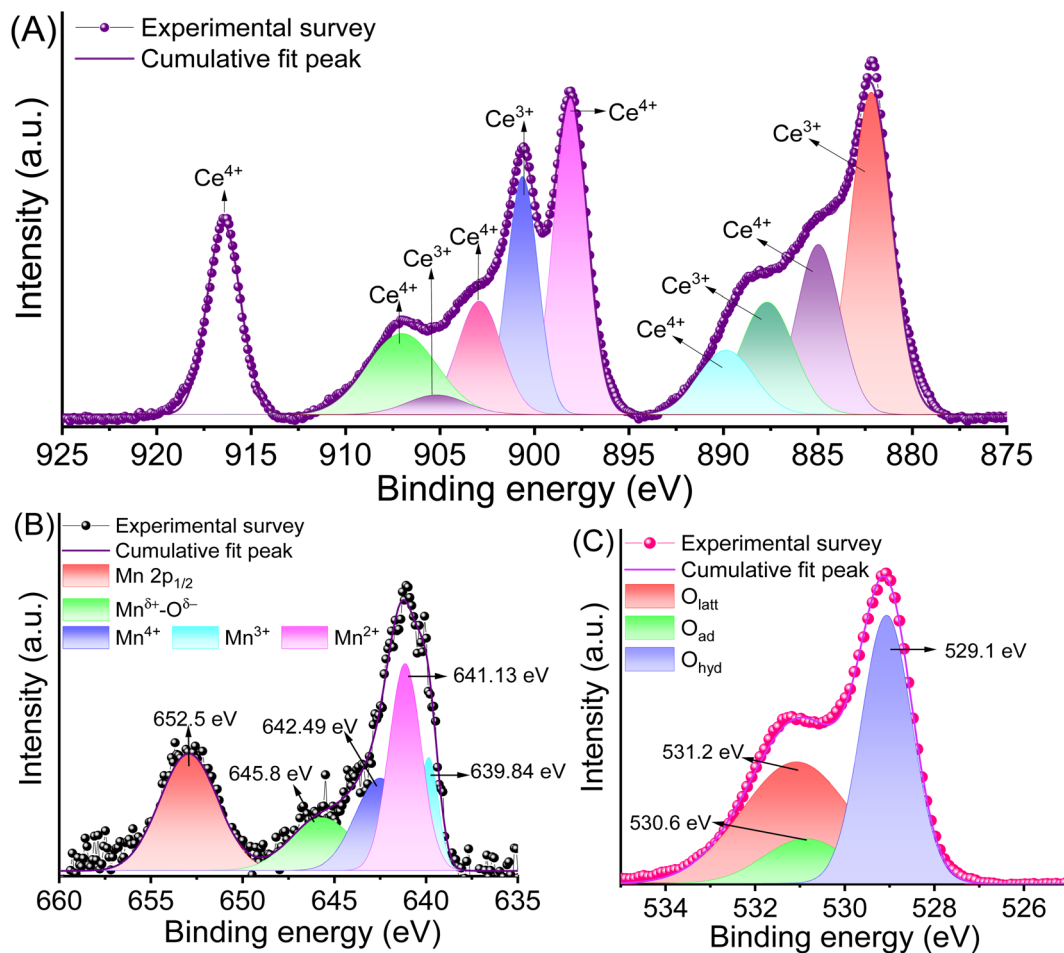


Fig. 3 Deconvoluted XPS curves of CeMnO_x representing (A) Ce 3d, (B) Mn 2p, and (C) O 1s.

catalytic activity in PEMFCs. Moreover, the surface FE-SEM images of SSM-73 and SS-73 are shown in Fig. S3,[†] along with their EDX elemental mapping illustrating the distribution of elements within the membranes. The presence of Ce, Mn, and a higher amount of O in SSM-73 indicates the successful incorporation of CeMnO_x into the membrane. Furthermore, upon analyzing the EDX data from different sections of the same membrane sample, it is observed that the weight and atomic content of C, S, O, Ce, and Mn almost remained similar, which confirms the uniform distribution of the fillers in the membrane. The topography and phase separation behavior in the PEMs were investigated by AFM analysis (Fig. 4G, H, S5 and S6, ESI[†]). The AFM images show no microphase separation and support nanophase-separated morphology. This suggests the successful incorporation of CeMnO_x into the SPEEK/SPPO polyelectrolyte without any agglomeration. However, a filler loading higher than 2% resulted in agglomeration in the PEMs. The roughness in the PEMs is within the nanoscale range, suggesting a smooth surface of the fabricated PEMs. The dark domains in the AFM image can be assigned to the hydrophilic phases within the PEMs, whereas the bright domains represent the hydrophobic regions. The TEM analysis also reveals the crystal lattice fringes of CeMnO_x , which are associated with the

(1 1 1), (1 0 0), and (1 1 0) planes due to Ce (Fig. 4I and J). This can also be correlated with the XRD pattern of the material, indicating the successful synthesis of CeMnO_x with crystalline nature. The uniform distribution of the fillers and the nanophase-separated morphology in the PEMs is also supported by their TEM analysis. Fig. S7, ESI[†] shows the TEM images of the composite PEMs, demonstrating the distribution of the fillers in the polymer matrix. The presence of lattice fringes, as well as the bright and dark domains representing hydrophobic and hydrophilic moieties in the polymer, indicates the distribution of fillers throughout the membrane. The uniform negative charge distribution on the prepared PEMs was confirmed by the dye test. The blue color from the dye was uniformly absorbed by the hydrophilic moieties, which suggests a co-continuous charge distribution throughout the PEMs and virgin SPEEK membrane, implying a uniform phase in PEMs similar to SPEEK (Fig. 4K and S8, ESI[†]).

3.2. Stability of the composite PEMs

The overall stability of the PEMs was analyzed in terms of their thermal, mechanical, tensile, and chemical stability. The T_g value of the PEMs was evaluated by their DSC analysis (Fig. 5A and B). The useful temperature range for PEMFC testing and the hot

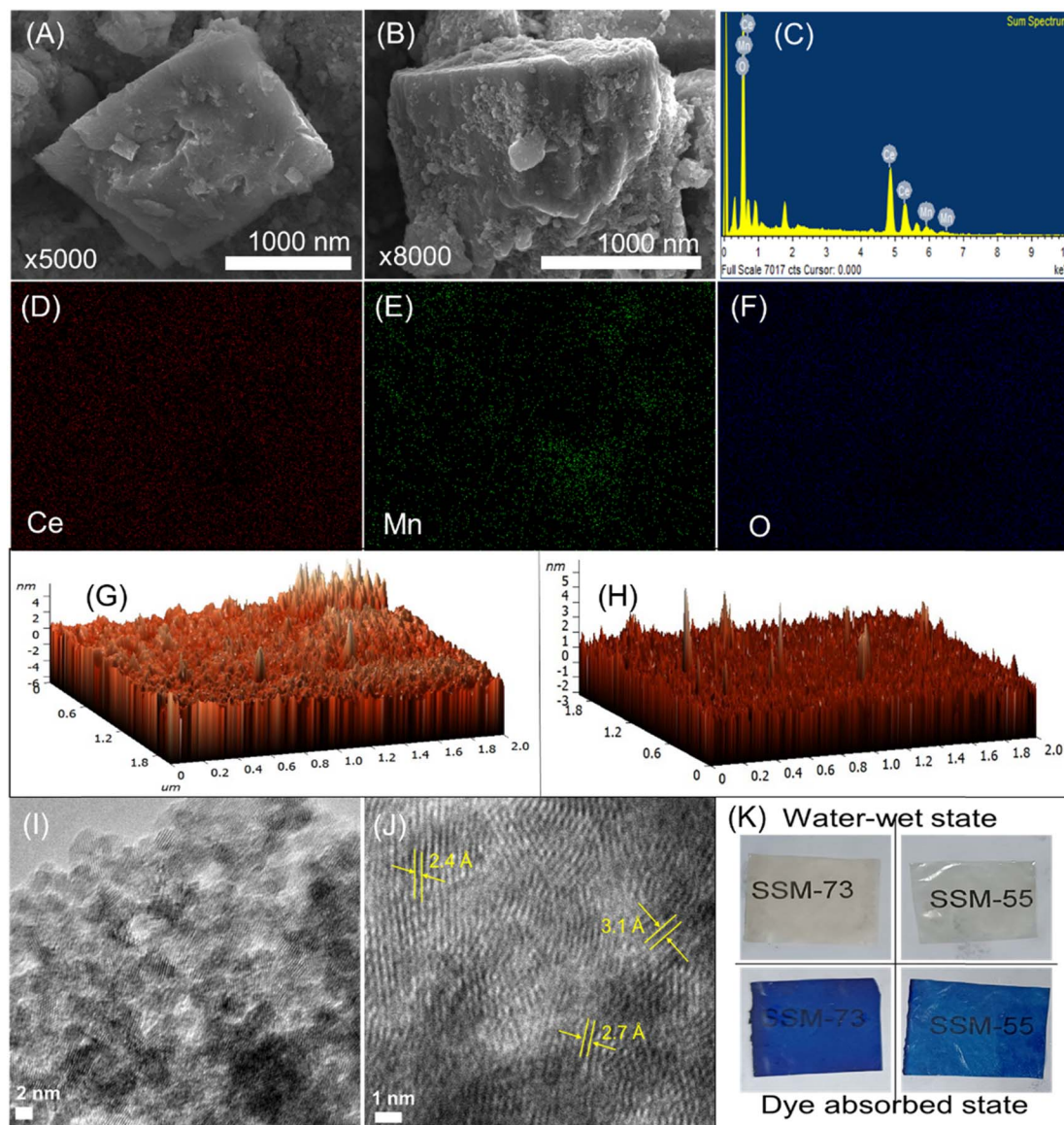


Fig. 4 (A and B) FE-SEM images, (C) EDX spectra, and (D–F) elemental mapping of CeMnO_x representing Ce, Mn, and O, respectively. (G and H) Topographical AFM images of SS-73 and SSM-73 respectively. (I and J) TEM images showing lattice fringes in CeMnO_x . (K) Physical images after dye absorption tests of SSM-73 and SSM-55 respectively.

pressing for MEA fabrication can be deduced concerning the T_g values of the membranes. All PEMs showed single T_g , which confirms an outstanding miscibility between SPEEK, SPPO, and the filler.⁵⁴ The T_g value of the PEMs follows the order of SS-73 (104.6 °C) < SS-55 (105.4 °C) < SSM-73 (113.2 °C) < SSM-55 (113.7 °C). The higher the SPPO content, the higher the T_g value. From the T_g values, it is clear that the incorporation of CeMnO_x nanostructures enhanced the T_g value of the membranes. It is known that the increase in hydrophilicity and charge in the membranes results in a decrease in the T_g value. However, in this case, the filler chemically interacts with the polymeric backbone (*via* H-bonding and bridge interactions) and impedes early glassy transition.

The thermal stability of the PEMs was further evaluated by TGA (Fig. 5C). All the PEMs exhibited three stages of

degradation. The first decomposition, around 90–130 °C, can be attributed to the loss of moisture and bound water content (due to hydrophilic sulfonic groups and filler) present in the samples. Enhanced thermal stability of the composite PEMs (SSM-55 and SSM-73) compared to that of the blend membranes (SS-55 and SS-73) is evident from the second-stage thermo-responsive degradation at 270–410 °C, attributed to the disruption of the sulfonic acid groups. It is worth noting that SSM-55 and SSM-73 show minimal weight loss in the range of 270–410 °C compared to SS-55 and SS-73 due to the intercalation of CeMnO_x in the composite PEMs. This can be explained in terms of H-bond network formation between the filler and the sulfonic acid groups in the membrane matrix, resulting in enhanced stability of the composite PEMs. The last stage of degradation at >440 °C is due to the decomposition of the

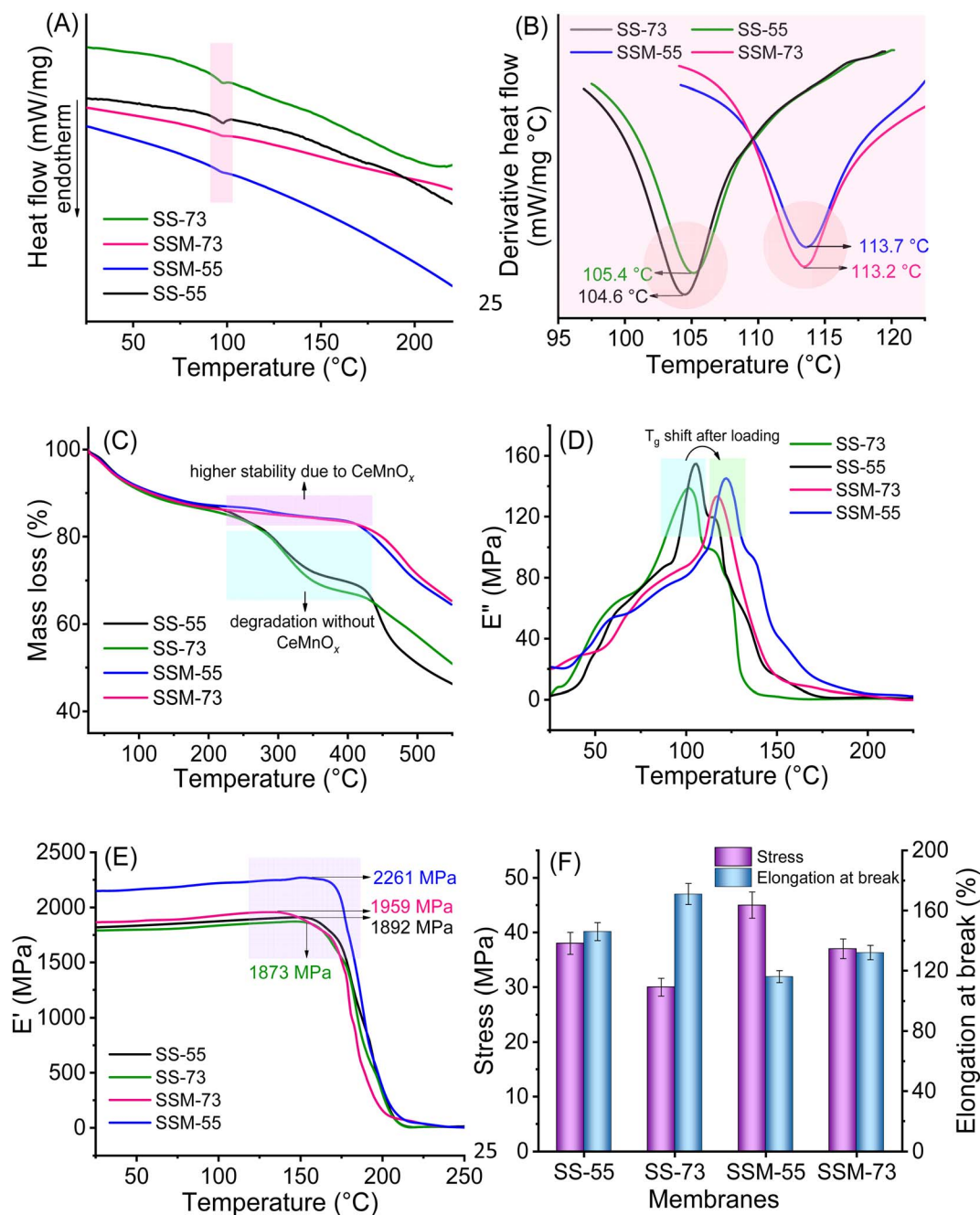


Fig. 5 (A) DSC curves, (B) derivative DSC curves, (C) TGA thermogram, (D) loss modulus vs. temperature curves, (E) storage modulus vs. temperature curves, and (F) UTM stress–strain analysis of the respective PEMs.

hydrocarbon-based polymeric backbone. Comparing the blend PEMs, SS-55 exhibits a higher thermal stability than that of SS-73 due to the presence of a higher SPPO content in the former. The mechanical stability of the PEMs was investigated using a DMA. The T_g values obtained from the DSC curves can be correlated with the loss modulus vs. temperature curves (Fig. 5D). The maxima of the loss modulus curves indicate the T_g value of the prepared membranes. Fig. 5E represents the storage modulus vs. temperature curves for the prepared PEMs. The SSM-55 membrane possesses the highest mechanical

stability with a storage modulus of 2261 MPa followed by SSM-73 (1959 MPa), SS-55 (1892 MPa), and SS-73 (1873 MPa). CeMnO_x imparts high mechanical strength in the membranes by holding the polymeric chains *via* chemical interactions.

The tensile property of the PEMs was evaluated by their stress–strain analysis using UTM in the water-wet state (Fig. 5F). SS-73 shows the lowest tensile stress of 30 MPa, as it was fabricated using only the blend polyelectrolyte with 70% (w/w) of SPEEK. Although SS-55 was also prepared using the blend solution, a higher SPPO content in SS-55 provides extra tensile

strength with a stress value of 38 MPa. Composite PEMs, *i.e.*, SSM-55 and SSM-73, exhibited higher tensile stress (45 MPa and 37 MPa, respectively) than the blend membranes. This indicates that the addition of CeMnO_x in the polymer matrix provides chain support by creating a three-dimensional network structure and H-bonding. The engineering strain of the PEMs was evaluated using their elongation at break (%) value. In this case, SS-73 exhibited the highest elongation of 171% due to the presence of a maximum amount of SPEEK in the membrane, which is followed by SS-55, SSM-73, and SSM-55 with elongation values of 146%, 132%, and 116%, respectively. Contrary to the tensile stress values, the composite PEMs show a lower tensile strain than the blend membranes. Incorporating CeMnO_x into the blend membranes enhances the rigidity of the membrane, resulting in low flexibility, reduced chain mobility, and lower elongation values.⁵⁶ A higher filler loading will increase the tensile stress and lower the elongation, making the membrane brittle. Hence, a fixed filler loading of 2% (w/w) provides a perfect stress–strain balance for PEMFC applications. Finally, the chemical stability of the prepared PEMs was investigated *via* exposure with Fenton's reagent (see later).

The stability trend (thermal, mechanical, and tensile) of all the prepared PEMs indicates that SSM-55 exhibits the highest overall stability compared to others, whereas SS-73 and SS-55 without any filler show lower stability. This can be explained in terms of chemical interactions present in the membrane matrix. In the sulfonated hydrocarbon-based SPEEK/SPPO blend membranes, besides strong H-bonding between –SO₃H moieties, multiple other chemical interactions such as permanent dipole–dipole interaction (keto–keto and sulfone–sulfone), weak London forces acting between aromatic polymer chains, and π – π interactions are present.⁵⁷ All these forces together minimize the strong H-bonding interactions in the PEMs. When CeMnO_x is added as a filler to the membrane matrix, it comes in between the polymer chains forming a three-dimensional structure that impedes dipole–dipole and London interactions between the aromatic chains. Moreover, the hydrophilic oxygen-rich surface of CeMnO_x itself comprises H-bonding and bridge interactions with the polymer backbone, which provides extra strength to the membranes.¹⁴ Hence,

composite PEMs are highly preferable for electrochemical energy devices. The stability analysis of the prepared PEMs also reveals that blending of SPPO is essential to enhance the oxidative stability of the SPEEK membrane even though radical scavengers are already in use. The sulfonation of PEEK is more facile, resulting in a greater degree of sulfonation than that of SPPO. This increased sulfonation leads to higher water uptake and swelling ratios, potentially causing dimensional and mechanical degradation of the PEMs. The blending of SPPO not only enhances oxidative stability but also provides additional mechanical and dimensional stability to the membrane.

3.3. Physical and electrochemical analyses

3.3.1. WU, SR, and IEC. Table 1 displays all the physico/electrochemical parameters of the prepared PEMs. The WU of the membrane is a crucial parameter to use in electrochemical separation applications.^{58,59} The composite PEMs undoubtedly show higher WU than the blend membranes and follow the order of SSM-73 (41.2%) > SSM-55 (39.4%) > SS-73 (37.2%) > SS-55 (36.3%). Comparing CeMnO_x with the blend membranes, the latter has more water-retaining capacity than CeMnO_x due to the presence of sulfonic groups. However, the intercalation of the same in the blend membranes uplifts the WU of the composite PEMs. In brief, the WU values showed an uplift of 10–12% in composite PEMs than the blend PEMs. The upgraded WU in SSM-73 and SSM-55 can be attributed to the coupling configuration of hydrophilic CeMnO_x and –SO₃H segments, which enhances the overall hydrophilic territory, catering surplus space to retain and sorb water. Similarly, the SR of the PEMs follows the same order due to the above-discussed attributes. The WU behavior of the PEMs in the ascended temperature ranges was also derived. All the PEMs exhibited a similar trend: the higher the temperature, the higher the WU. This effect can be explained by two factors; (i) the solvation effect and (ii) activated chain mobility at higher temperatures. However, the incorporation of the filler facilitates dimensional stability with higher WU at high temperatures.

IEC is one of the pivotal properties of a PEM, which is directly proportional to the degree of sulfonation, indicating the

Table 1 Physicochemical and electrochemical parameters of the prepared PEMs^a

Parameters	Membranes			
	SS-73	SS-55	SSM-73	SSM-55
Thickness (μm)	62.0	58.0	61.0	59.0
WU (%)	37.2 \pm 0.8	36.3 \pm 0.6	41.2 \pm 1.2	39.4 \pm 1.0
SR (%)	29.2 \pm 0.7	26.8 \pm 0.7	30.5 \pm 0.9	28.5 \pm 0.8
IEC (meq g ⁻¹)	1.70 \pm 0.03	1.68 \pm 0.02	1.36 \pm 0.02	1.32 \pm 0.02
χ^m (meq cm ⁻³)	0.45	0.47	0.33	0.36
λ_{hyd}	12.1	12.0	16.8	16.6
Surface resistance (Ω)	0.63	0.79	0.52	0.56
K^m (mS cm ⁻¹)	12.7	9.4	15.0	13.5
K^{m*} (mS cm ⁻¹)	23.2	19.2	26.7	24.3
E_a (kJ mol ⁻¹)	11.39	12.54	9.47	10.89

^a K^{m*} represents K^m at 80 °C.

$-\text{SO}_3\text{H}$ content in the membrane. The proton conduction mechanism is significantly affected by their IEC values, especially the Grotthuss-type migration as it reflects the hopping sites embedded in the membrane. The IEC values of the PEMs follow the order of SSM-55 (1.32 meq g^{-1}) < SSM-73 (1.36 meq g^{-1}) < SS-55 (1.67 meq g^{-1}) < SS-73 (1.70 meq g^{-1}). Unlike WU, the IEC of the composite PEMs is lower than the blend membranes. This can be demonstrated by two factors; (i) charge dilution: $-\text{SO}_3\text{H}$ partially interacts with CeMnO_x and is not fully available for interaction with foreign cations, and (ii) electrostatic repulsion: high valence states of Ce^{z+} and Mn^{m+} repel the cations electrostatically and impede chemical interactions with $-\text{SO}_3\text{H}$ moieties.¹⁴

3.3.2. K^m , E_a , and ion conduction mechanism. K^m is the most critical electrochemical property of a PEM, as it is responsible for proton conduction which commands operational potential and power density of PEMFCs.⁶⁰ K^m of the PEMs also shows a similar trend to that of WU. SSM-73 exhibits the highest K^m value of 15.0 mS cm^{-1} , followed by SSM-55, SS-73, and SS-55 with 13.5, 12.7, and 9.4 mS cm^{-1} at RT under hydrated conditions (Fig. 6A and B). The proton conduction in the blend membranes is due to the proton conducting pathways created by sulfonic acid groups, whereas, in the case of composite membranes, the conduction is profited due to the addition of conducting CeMnO_x fillers in the polymer electrolyte (Scheme 2). The incorporation of CeMnO_x nanostructures

in SSM-73 showed a hike of 18% in their K^m value than that of SS-73. Hence, the newly constructed coupling interfacial configuration of CeMnO_x and blend polymer is accountable for higher proton conductivity.⁷ The increased bound water content due to the addition of fillers generates extra carrier sites for proton diffusion in the form of hydronium ions *via* the vehicular proton transport mechanism (Scheme 2).¹⁴ Additionally, 3D conductive CeMnO_x constructs extensive H-bond networks with sulfonic acid groups, reducing the proton hopping gap and enabling protonation-deprotonation at hydrophilic sites for the Grotthuss-type proton transport mechanism (Scheme 2). Furthermore, the filler creates sub-nano ionic channels, which contribute to efficient proton conduction. These channels are created due to the induction of a prominent dipole because of the oxygen vacancies (lattice defects) and the electronegativity difference between the metallic sites.³⁶ The K^m values of the membranes were derived from their respective Nyquist plots. Fig. 6B shows the magnified Nyquist plot in higher frequencies to elucidate the exact membrane resistance. SSM-73 showed the lowest resistance of 0.52Ω (when sandwiched between electrodes with an area of 0.78 cm^2) compared to other prepared PEMs. The higher the conductivity, the lower the resistance. A filler loading more than 2% (w/w) shows agglomeration in the membranes, which reduces its interfacial adhesion with sulfonic groups, resulting in lower K^m .

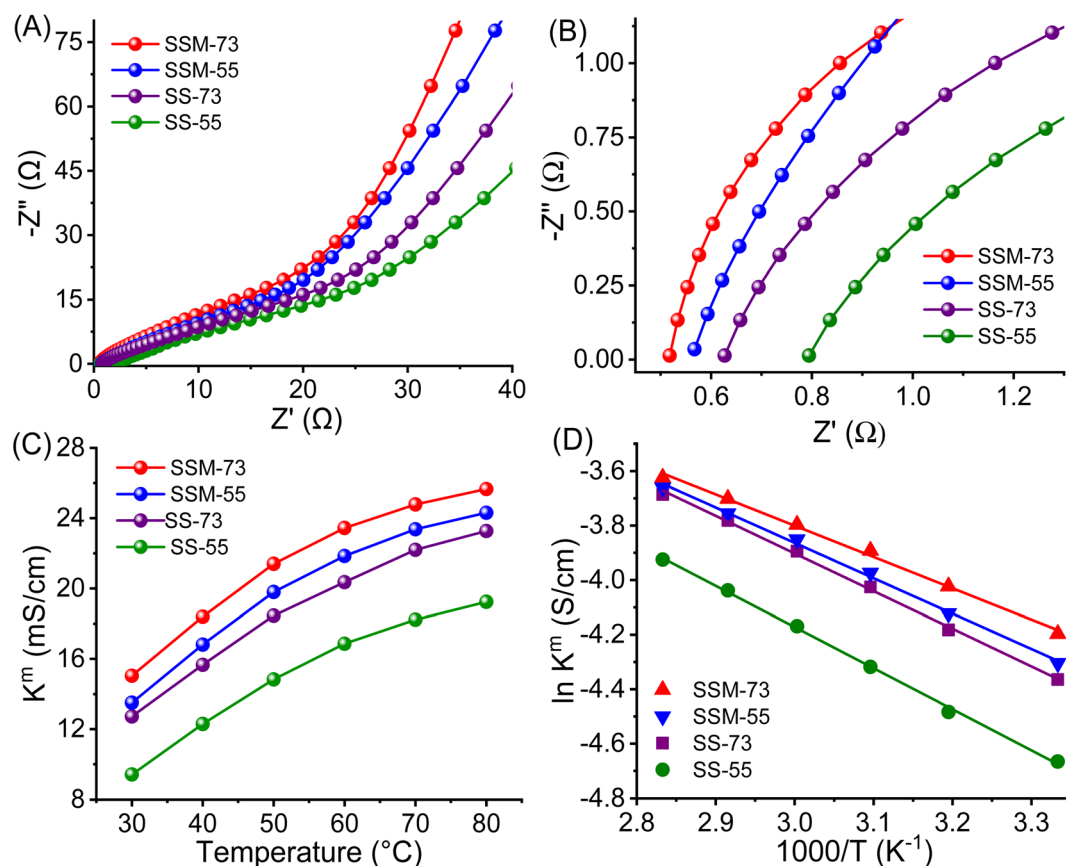
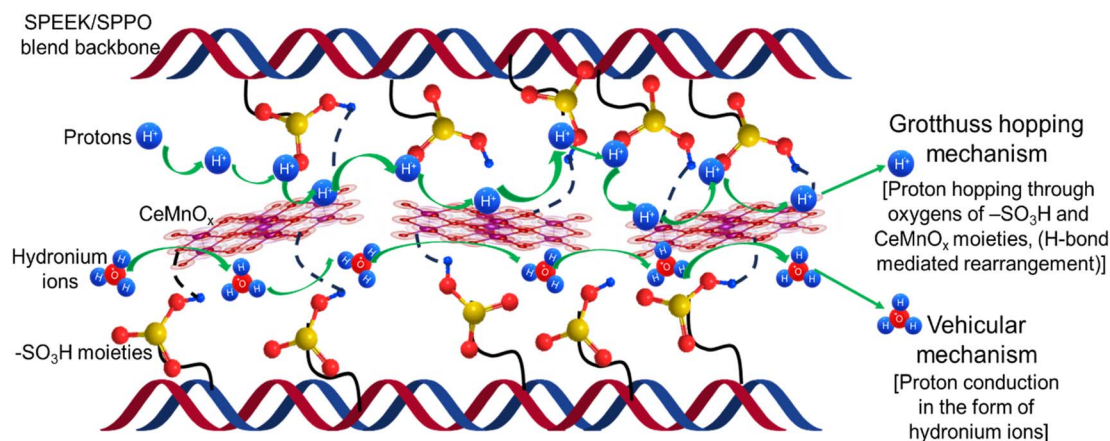


Fig. 6 (A) Nyquist plot, (B) magnified Nyquist plot, (C) K^m vs. temperature curves, and (D) Arrhenius plot of the prepared PEMs.



Scheme 2 Schematic illustration of the proton conduction mechanism through the composite PEMs.

The proton transport properties of the PEMs were further assessed by their temperature-dependent conductivities and portrayed in Fig. 6C. Similar to WU, the K^m value of all the prepared PEMs increased gradually with temperature. SSM-73 showed a K^m value of 26.7 mS cm^{-1} at 80°C , which is 78% higher than its K^m value at RT (15.0 mS cm^{-1}). The flexible chain mobility and activated water content at higher temperatures increase the entropy of the system, promoting proton transport kinetics. Further, the E_a value of the PEMs has been elucidated from the Arrhenius plot (Fig. 6D and eqn (S9)†). It is well known that Nafion exhibits an E_a value of $\approx 10 \text{ kJ mol}^{-1}$. From the prepared PEMs, SSM-73 displayed an E_a value of 9.47 kJ mol^{-1} , whereas other PEMs showed higher E_a values than that of Nafion. The E_a values for SSM-55, SS-73, and SS-55 are $10.89 \text{ kJ mol}^{-1}$, $11.39 \text{ kJ mol}^{-1}$, and $12.54 \text{ kJ mol}^{-1}$, respectively. Integrating with the proton conducting parameter, SSM-73 exhibits the least proton transport barrier, resulting in the lowest E_a values among all.

3.3.3. Oxidative stability and mechanism. The resistivity of the prepared PEMs to the chemical oxidative degradation has been measured in terms of conductivity and mass loss after

exploiting membrane samples in harsh Fenton's solution ($3\% \text{ H}_2\text{O}_2 + 3 \text{ ppm Fe}^{2+}$) at 60°C for 6 h keeping addition of fresh Fenton's solution after 2 h each. The loss in K^m and mass with respect to time is represented in Fig. 7. Fig. S9, ESI,† displays the digital images of the prepared membranes at different time intervals, confirming the oxidative stability of the PEMs. H_2O_2 in Fenton's reagent creates a similar chemical environment as PEMFCs, where $\cdot\text{OH}$ radicals are formed due to fuel crossover (either from the cathode to the anode side or *vice versa*).¹⁴ Hence, Fenton's reagent can be utilized to test the defense mechanism of the PEMs towards $\cdot\text{OH}$ and $\cdot\text{OOH}$ radicals. The chemical stability of the prepared PEMs was found to be proportional to the SPPO and CeMnO_x content in the blend polyelectrolyte. The prepared PEMs exhibited a mass loss of 2%, 5%, 14%, and 18% ascertained to SSM-55, SSM-73, SS-55, and SS-73, respectively after Fenton's test (Fig. 7). Similarly, the PEMs showed a minimal loss of K^m value of 4%, 9%, 13%, and 21% for SSM-55, SSM-73, SS-55, and SS-73, respectively (Fig. 7). From the loss values, it is evident that the incorporation of CeMnO_x in the membranes provides surplus chemical stability to the PEMs (mechanism discussed later). It is clear from Fig. 7 that both the mass loss and K^m loss

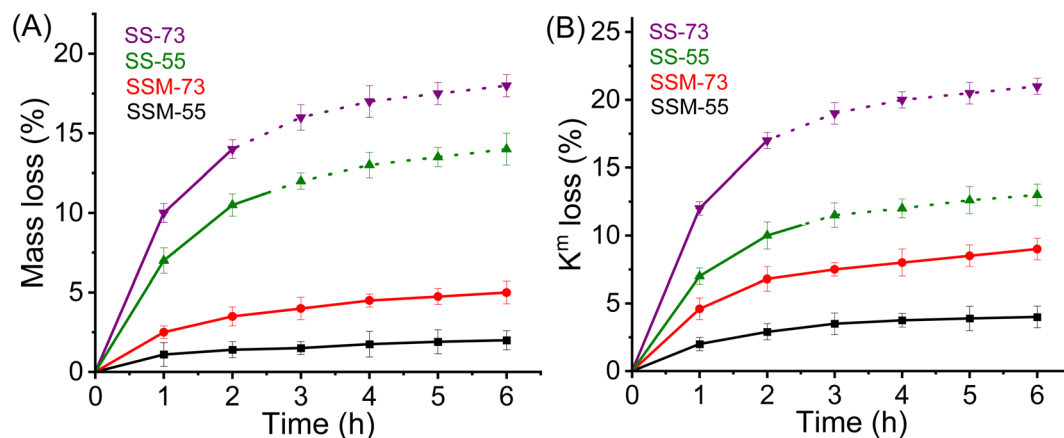


Fig. 7 Oxidative stability analysis in terms of (A) mass loss vs. time and (B) conductivity loss vs. time after Fenton's test (dotted regions indicate the average oxidative stability measured with the recast membrane samples).

have a sharp slope up to 3 h and then achieve a steady-state phase, which illustrates the ability of the PEMs to resist chemical degradation over time. It is observed that SS-73 and SS-55 exhibited some visible degradations after 2 and 2.5 h, respectively. Hence, the K^m loss was measured by its recast SS-73 and SS-55 membranes, and is shown as a dotted line in Fig. 7.

The prominent factor governing oxidative stability is the simultaneous existence of Ce^{3+}/Ce^{4+} ions and Mn^{2+}/Mn^{3+} ions in the composite PEMs. These metal ions act as radical scavengers and protect the membrane backbone from radical attack. The main trait which is responsible for the protection from $\cdot OH$ radicals is the Ce^{3+} ion. Ce^{3+} reacts with $\cdot OH$ to produce Ce^{4+} and $\cdot OOH$ radicals. Similarly, Mn^{2+} reacts with $\cdot OH$ and forms Mn^{3+} . The rate constant for the reaction of $\cdot OH$ with Ce^{3+} is quite higher ($3 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$) than with Mn^{2+} ($0.4 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$).⁶¹ Ce^{4+} and Mn^{3+} further oxidize $\cdot OOH$ and H_2O_2 to regenerate Ce^{3+} and Mn^{2+} ions respectively, along with O_2 . This redox cycling goes on over the fuel cell testing period and restricts the radicals from harming the membrane. Hence, the respective quenching chemistry of Ce^{3+} and Mn^{2+} states present in the $CeMnO_x$ nanostructures serves as efficient protectors from the chemical degradation of the membranes. Fig. S10 and S11 (ESI)[†] show the MEA condition after multiple single-cell PEMFC testing and the durability test. It is clear from the images that the prepared MEAs are devoid of any pinholes or cracks and stable after the durability test.

Moreover, the effect of $CeMnO_x$ bimetallic oxides over their individual oxides, *i.e.*, CeO_2 and MnO_2 as fillers was explored using the SSM-73 membrane with 2% filler loading each. The oxidative stability in terms of mass and K^m loss of the PEMs using all these fillers is displayed in Fig. S12, ESI.[†] It is worth noting that the oxidative stability of SSM-73 using a $CeMnO_x$ filler is higher than that of CeO_2 and MnO_2 . The higher oxidative stability of $CeMnO_x$ can be ascertained to its higher Ce^{3+} content. The insertion of Mn^{m+} ions into the crystal lattice of CeO_2 distorts its lattice with an increased Ce^{3+} content. However, in the CeO_2 nanoparticles, the Ce^{4+} content is higher than the Ce^{3+} content. As Ce^{3+} plays a superior role in sustaining oxidative tolerance over Ce^{4+} , $CeMnO_x$ exhibited a higher oxidative stability than that of CeO_2 . Moreover, SSM-73 using a MnO_2 filler exhibited the least oxidative stability compared to CeO_2 and $CeMnO_x$. This may be due to the lower efficiency of Mn^{2+}/Mn^{3+} as a free radical scavenger than that of Ce^{3+}/Ce^{4+} . Furthermore, WU and K^m of SSM-73 using these fillers were also compared, and are shown in Table S2.[†] $CeMnO_x$ outplays other materials as fillers, as it imparts higher K^m in SSM-73. This is because the doping of Mn^{m+} in the CeO_2 lattice creates more oxygen vacancies, which helps in proton conduction. The corresponding Nyquist plots using $CeMnO_x$, CeO_2 , and MnO_2 as fillers are depicted in Fig. S13, ESI.[†] Considering the above-mentioned facts, it is clear that the $CeMnO_x$ bimetallic oxide is more effective as the filler than its individual oxides. Furthermore, to assess the acid stability of composite PEMs employing $CeMnO_x$ and CeO_2 as fillers, two pieces of SSM-73 membranes with 2% filler loading were immersed in 3 M, 6 M, and 9 M HCl for 7 days. The acid stability was measured in terms of mass loss of the respective PEM samples. SSM-73 with

$CeMnO_x$ and CeO_2 filler exhibited a negligible mass loss of 0.2–0.3% in 3 M HCl, indicating that the composite PEMs are highly stable in 3 M HCl. However, SSM-73 with $CeMnO_x$ exhibited a mass loss of 6.2% and 17.7% in 6 M and 9 M HCl, respectively, whereas it showed a mass loss of 6.7% and 18.8% using CeO_2 as the filler in 6 M and 9 M HCl, respectively. It is noteworthy that the acid stabilities of SSM-73 with $CeMnO_x$ and CeO_2 fillers is close to each other. From the above-mentioned experiment, it can be inferred that the Mn^{m+} doping in the CeO_2 lattice to prepare the $CeMnO_x$ bimetallic oxide does not exert a significant influence on the acid stability of the PEMs. However, the oxidative stability of the PEMs showed a remarkable superiority using $CeMnO_x$ as fillers over CeO_2 or MnO_2 .

The physicochemical, electrochemical, and oxidative stability analysis of the other prepared blend PEMs with different SPEEK/SPPO ratios is shown in Table S3, ESI.[†] Although SS-91 and SS-82 exhibited higher K^m values than that of SS-73, their high WU and low oxidative stability make them dimensionally unstable and prone to radical attack, respectively. Consequently, pristine and composite membranes based on SS-91 and SS-82 were not considered for further investigations, *i.e.*, hydrogen fuel cell testing and OCV degradation test. It is worth noting that the K^m value decreases as the SPPO content increases in the blend PEMs (due to lower DoS of SPPO than that of SPEEK). Therefore, there is no need to construct blend PEMs with a higher SPPO content than the SPEEK content as it will lower the K^m value and fuel cell performance of the PEMs. Hence, blend PEMs were fabricated only up to a SPEEK/SPPO ratio of 50/50 (w/w). Considering the above-mentioned properties (Table S3[†]), only SS-73 and SS-55 were only considered for further investigation.

3.4. Single-cell PEMFC studies

The single-cell H_2 - O_2 fuel cell performance of all the prepared PEMs, *i.e.*, SSM-73, SSM-55, SS-73, and SS-55, was investigated by their polarization curves represented in Fig. 8A. In terms of thermal, mechanical, and chemical stabilities, SSM-55 is a better candidate, whereas in terms of proton conducting attributes, IEC, and E_a SSM-73 should be a better candidate. All the prepared PEMs showed an OCV of 0.91–0.95 V, which assures them as a potential candidate for fuel cells. The single-cell stack packed with all the prepared MEAs (16 cm² each) showed a cell resistance in the range of 62–68 m Ω , reducing the total ohmic loss in the system. The cell resistance follows the order of SSM-73 (62 m Ω) < SSM-55 (64 m Ω) < SS-73 (67 m Ω) < SS-55 (68 m Ω). This suggests that the prepared membranes are highly conducting and exhibit lower resistance as a fuel cell components. As the resistance of other components, such as electrodes and bipolar plates remains constant, the resistance of the PEM becomes the sole determinant of the total cell resistance or ohmic loss in the system. The lower membrane resistance results in reduced ohmic losses in the fuel cell system. After testing the respective MEAs in the FCT, SSM-73 showed a peak power density of 431.2 mW cm⁻² and a maximum current density of 1272.6 mA cm⁻² at 75 °C (100% RH), which is the highest among all the other prepared PEMs.

This is followed by the SSM-55 membrane with a maximum power density of 370.5 mW cm^{-2} and a current density of $1078.1 \text{ mA cm}^{-2}$. As the CeMnO_x loading is the same in SSM-73 and SSM-55 (2%, w/w), a higher conductivity (26.7 mS cm^{-1} at high temperature) and an optimum balance between membrane hydrophilicity and mass transport in the former is responsible for its better PEMFC performance. It is true that the conductivity of the membrane primarily facilitates proton transport and may not directly influence the performance in the low-voltage region, it indirectly impacts the overall cell performance. Higher proton conductivity ensures efficient proton transport within the membrane, which can help maintain a stable proton gradient across the MEA interface. However, the relationship between membrane hydrophilicity and gas transport resistance is indeed complex. It is true that higher hydrophilicity can enhance the water retention of the membrane and may lead to the accumulation of water and flooding issues in the cell, which will increase the gas transport resistance. However, the architecture of the prepared composite PEMs is so designed that it imposes a balance between the hydrophilicity and the mass transport within the fuel cell. SSM-73 with 2% filler loading showed improved water management property, which helps maintain proper hydration levels within the membrane and fuel cell system. This, in turn, inhibits the flooding issue, facilitating proton conduction and ensuring efficient mass transport of reactants (primarily oxygen), thereby leading to improved performance of SSM-73 at low-voltage regions. The SS-73 membrane displayed a peak power density of 286.9 mW cm^{-2} (34% lesser than SSM-73) and a maximum current density of 872.3 mA cm^{-2} (40% lesser than SSM-73) under the same operational conditions. However, the pristine SPEEK membrane without incorporation of filler exhibited a peak power density of 230.7 mW cm^{-2} and a maximum current density of 692.3 mA cm^{-2} at 75°C (100% RH) (Fig. S14, ESI†). Moreover, the impact of filler loading at three different levels, namely 1%, 2%, and 3%, on the fuel cell performance has been investigated. The fuel cell performance was less favorable in cases where the filler loading was either 1% or 3%. SSM-73 with 1% filler loading demonstrated a peak power density of 330 mW cm^{-2} and a maximum current density of 959

mA cm^{-2} , whereas SSM-73 with 3% filler loading showed a peak power density of 364 mW cm^{-2} with a maximum current density of 1056 mA cm^{-2} under similar conditions. The performance with 1% filler loading was inferior to that with 2% loading, mainly due to the relatively limited influence of fillers at the lower loading level. However, when the filler content was increased to 3%, it resulted in agglomeration on the membrane, which prevented it from effectively contributing to fuel cell performance.¹⁴ The Nyquist plot of SSM-73 with filler loadings of 1%, 2%, and 3% is shown in Fig. S15, ESI†. The surface resistance of SSM-73 with 2% filler loading exhibits the lowest resistance of 0.52Ω followed by 3% filler loading (0.55Ω) and 1% filler loading (0.57Ω). This can be clearly correlated with the fuel cell performance of SSM-73 using different filler loadings (w/w).

Besides Fenton's test, the chemical stability of the prepared MEAs was evaluated by AST. Fig. 8B displays the OCV variation of the cell as a function of time at an operating temperature of 75°C at 30% RH. Comparing SS-73 and SS-55, it is revealed that the OCV of SS-73 is higher than that of SS-55 initially. The higher OCV of SS-73 than that of SS-55 was maintained over a period of 28 h, after which the OCV decay was more pronounced in SS-73 than in SS-55. After 50 h, the OCV decay for SS-55 was 36.5%, which is lower than that of SS-73 (43.7%). SS-55 resists the free radical attack more than SS-73 due to its higher SPPO content, which imparts better oxidative stability in the former. The addition of CeMnO_x in the membrane matrix dramatically enhances the OCV retention of SSM-73 and SSM-55. CeMnO_x as the antioxidant plays the role of free radical scavengers and mitigates the radical attack on the polymeric backbone. SSM-55 exhibited the least OCV decay of 12.2% followed by SSM-73, which showed an OCV decay of 13.4%. The durability test of the prepared PEMs can be correlated with the oxidative stability analysis using Fenton's reagent. Furthermore, the enhancement of oxidative stability using SPPO as a blend material was further validated by the durability test of these membranes. A new SPEEK membrane without any SPPO content having 2% CeMnO_x loading was prepared and exposed to its durability test. SSM-73 (blend composite PEM) exhibited an OCV degradation of 13.4%, whereas the SPEEK composite

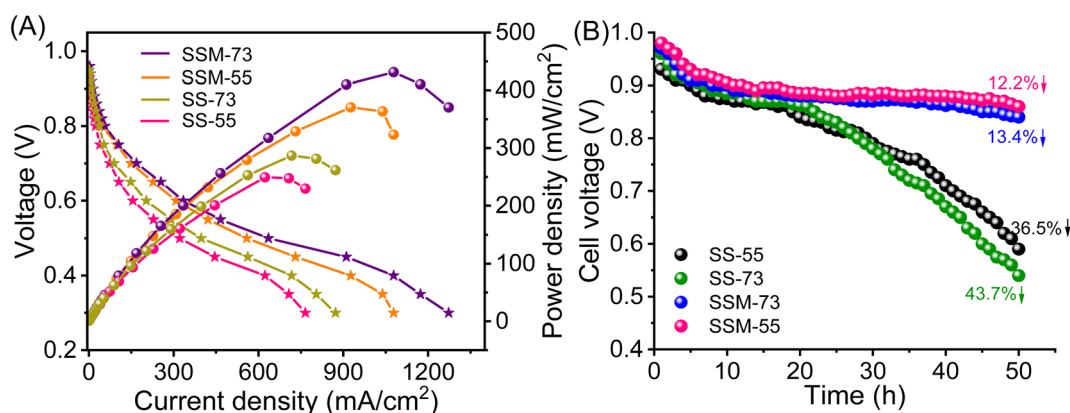


Fig. 8 (A) Polarization curves of the prepared PEMs at 75°C and 100% RH. (B) Durability test in terms of OCV measurements with respect to time.

membrane displayed an OCV degradation of 16.2% after 50 h of accelerated stress test. Moreover, the K^m retention of the PEMs after the accelerated stress test was also measured to evaluate the durability in terms of conductivity (Fig. S16, ESI†). The K^m value of the PEMs after the durability test follows the order of SSM-73 (23.5 mS cm^{-1}) > SSM-55 (22.3 mS cm^{-1}) > SS-73 (19.7 mS cm^{-1}) > SS-55 (16.5 mS cm^{-1}). However, it is worth noting that the K^m decay percentage of SSM-73 is slightly higher at 6.5%, than that of SSM-55, which had a K^m decay of 6%. This discrepancy is attributed to the higher oxidative stability of SSM-55, as confirmed by the results from Fenton's test. Moreover, the hydrogen crossover experiments were conducted by linear sweep voltammetry (LSV), and are represented in Fig. S17, ESI.† The H_2 crossover was measured by its hydrogen crossover current with hydrogen flowing at the anode (100 mL min^{-1}) and nitrogen at the cathode (300 mL min^{-1}), 75°C cell temperature and 100% RH. The scan rate was kept constant at 50 mV s^{-1} ,

ranging from 0 to 0.8 V. The maximum current in the LSV curves indicates a higher H_2 crossover through the respective MEA. This is because a higher current suggests more hydrogen gas is crossing over from the anode to the cathode side through the membrane and splits into proton and electron also at the cathode side. The H_2 crossover experiments suggest that the composite PEMs exhibited lesser H_2 crossover than their parent blend membranes. The real-time acid stability of the PEMs was accessed using the cross-sectional FE-SEM images of the prepared MEAs before and after the fuel cell performance and durability (accelerated stress test) analysis. The cross-sectional FE-SEM images of the MEAs are presented in Fig. 9. It is clear from these images that the membranes sandwiched between the two gas diffusion electrode layers are devoid of any cracks or pinholes, suggesting robust acid stability of the PEMs under real fuel cell testing conditions. Furthermore, after the AST, the product water eluent was systematically collected and subjected

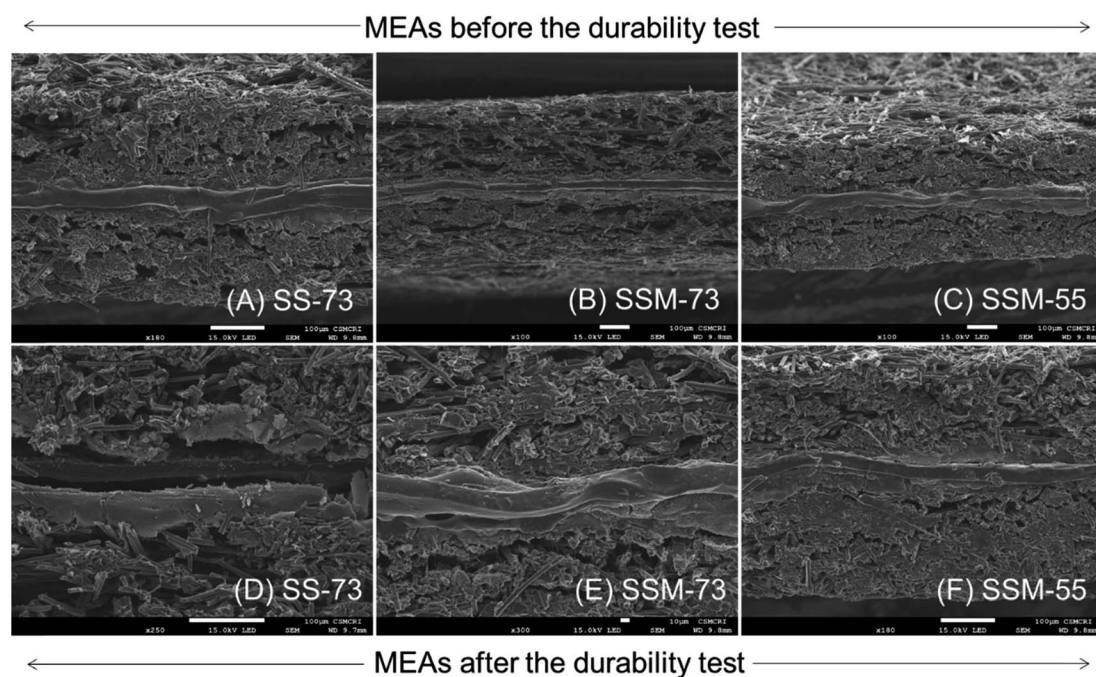


Fig. 9 Cross-sectional FE-SEM images of the prepared MEAs before and after the durability test: (A and D) SS-73, (B and E) SSM-73, (C and F) SSM-55.

Table 2 Comparative study of the PEMFC performance of the prepared PEMs with the similar reported PEMs

Sl no.	Membrane	Peak power density (mW cm^{-2})	Maximum current density (mA cm^{-2})	Ref.
1	SPEEK/SGO	378	830	63
2	SPEEK/HPW@MIL 101	383	784	64
3	C-SP90/SP90NF	485	1750	65
4	SPEEK/W-SBA-15	405	785	66
5	SPEEK-CNT-FA-6	672	1625	67
6	SPAEEK/f-SiO ₂	445	700	68
7	M-SP/DMNPs@HPW-2	433	1180	69
8	mPTA-Si/SPEEK	180.87	690	70
9	SSM-73	431.2	1272.6	This work
10	SSM-55	370.5	1078.1	This work

to UV-visible spectroscopy to examine any component leakage from the composite PEMs.⁶² It is clear from Fig. S18, ESI,† that CeMnO_x-dispersed water showed an absorbance peak at 313 nm, whereas there is no significant absorbance for the product water obtained during the AST, indicating no traces of CeMnO_x leaching out from the composite PEMs. However, there may be a possibility of CeMnO_x leaching, which remains in the catalyst layer when it interacts with ionic species such as Nafion ionomers, which may not be detected in the eluent. Table 2 shows the comparison of the fuel cell performance of the reported membranes with our prepared PEMs, which indicates the high potential of the prepared PEMs for better performance and durability in PEMFCs.

4. Conclusions

In summary, three-dimensional CeMnO_x bimetallic oxide nanostructures have been synthesized and employed as fillers to fabricate composite polymer electrolyte membranes for H₂-O₂ fuel cells. The blending chemistry of SPEEK and SPPO was utilised to design highly stable CeMnO_x@SPEEK/SPPO hydrocarbon-based membranes. The filler loading was optimized and fixed at 2% (w/w) in composite membranes. In this work, four typical membranes, namely two blend membranes (with different blending ratio) and two composite blend membranes, were fabricated. All the membranes showed nanophase separated morphology, indicating the excellent miscibility of the blend polyelectrolyte and the filler. Apparent reinforcement in stability and proton conductivity was manifested in the composite membranes due to the consistent distribution of CeMnO_x nanostructures over the polymer matrix. The outshining properties of the composite membranes are due to (i) a higher volume of CeMnO_x, which increases the bound water content and (ii) the formation of H-bond networks, coupling configurations, and bridge interactions of fillers with sulfonic acid segments in the membrane, which enhances stability and creates sub-nano pathways for proton conduction. CeMnO_x endorses both vehicular and Grotthuss-type pathways, resulting in fast proton conduction (26.77 mS cm⁻¹ for SSM-73 at 80 °C) and low activation energy barrier (9.47 kJ mol⁻¹ for SSM-73) in the composite PEMs. Moreover, the existence of Ce³⁺ and Mn²⁺ in the filler helps in the radical scavenging to produce outstanding oxidative stability with a negligible mass loss of 5–8% and a minimal K^m loss of 2–5% in the composite membranes. The composite PEMs (SSM-73 and SSM-55) showed >86% of OCV retention after 50 h of accelerated stress test, confirming their enhanced durability compared to SS-55 and SS-73. The representative SSM-73 membrane exhibited a peak power density of 431.2 mW cm⁻² and the highest current density of 1272.6 mA cm⁻² at 75 °C and 100% RH, which proves the potential of the composite membrane as a meritorious candidate for hydrogen fuel cells.

Author contributions

Uma Chatterjee: conceptualization, methodology, supervision, funding acquisition, project administration, and writing – review and editing. Sk Miraz Hossain: writing – review and editing, methodology, investigation, and formal analysis.

Pratyush Patnaik: writing – original draft, investigation, and formal analysis. Suman Sarkar: data curation, validation, and visualization. Ritika Sharma: visualization and formal analysis.

Conflicts of interest

There are no conflicts to declare.

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References

- Z. J. Baum, L. L. Diaz, T. Konovalova and Q. A. Zhou, *ACS Omega*, 2022, 7, 32908–32935.
- Y. Wang, D. F. R. Diaz, K. S. Chen, Z. Wang and X. C. Adroher, *Mater. Today*, 2020, 32, 178–203.
- P. Patnaik, R. Mondal, S. Sarkar, A. Choudhury and U. Chatterjee, *Int. J. Hydrogen Energy*, 2022, 47, 41920–41931.
- I. Staffell, D. Scamman, A. V. Abad, P. Balcombe, P. E. Dodds, P. Ekins, N. Shah and K. R. Ward, *Energy Environ. Sci.*, 2019, 12, 463–491.
- R.-A. Felseghi, E. Carcadea, M. S. Raboaca, C. N. Trufin and C. Filote, *Energies*, 2019, 12, 4593.
- D. W. Shin, M. D. Guiver and Y. M. Lee, *Chem. Rev.*, 2017, 117, 4759–4805.
- Y. Liu, J. Zhang, X. Zhang, Y. Li and J. Wang, *ACS Appl. Mater. Interfaces*, 2016, 8, 20352–20363.
- X. Ge, F. Zhang, L. Wu, Z. Yang and T. Xu, *Macromolecules*, 2022, 55, 3773–3787.
- M. Pirali-Hamedani and S. Mehdipour-Ataei, *Polym. Adv. Technol.*, 2017, 28, 1495–1503.
- N. A. M. Harun, N. Shaari and N. F. H. N. Zaiman, *Int. J. Energy Res.*, 2021, 45, 19671–19708.
- P. Hosseinabadi, K. Hooshyari, M. Javanbakht and M. Enhessari, *New J. Chem.*, 2019, 43, 16232–16245.
- J. Li, H. Wu, L. Cao, X. He, B. Shi, Y. Li, M. Xu and Z. Jiang, *ACS Appl. Nano Mater.*, 2019, 2, 4734–4743.
- S. Gahlot, P. P. Sharma, V. Kulshrestha and P. K. Jha, *ACS Appl. Mater. Interfaces*, 2014, 6, 5595–5601.
- S. M. Hossain, P. Patnaik, R. Sharma, S. Sarkar and U. Chatterjee, *ACS Appl. Mater. Interfaces*, 2024, 16, 7097–7111.
- J. Shin, B. Chang, J. Kim, S. Lee and D. Suh, *J. Membr. Sci.*, 2005, 251, 247–254.
- S. Subhadarshini, J. S. Sravan, O. Sarkar, S. V. Mohan, T. K. Roy and T. Jana, *ACS Appl. Energy Mater.*, 2023, 6, 1422–1438.
- S. Takamuku, A. Wohlfarth, A. Manhart, P. Räder and P. Jannasch, *Polym. Chem.*, 2015, 6, 1267–1274.
- J. Pan, L. Zhu, J. Han and M. A. Hickner, *Chem. Mater.*, 2015, 27, 6689–6698.

- 19 T. de Wild, T. Nemeth, T. M. Nolte, T. J. Schmidt, T. Nauser and L. Gubler, *J. Electrochem. Soc.*, 2021, **168**, 054514.
- 20 J. Wu, X. Z. Yuan, J. J. Martin, H. Wang, J. Zhang, J. Shen, S. Wu and W. Merida, *J. Power Sources*, 2008, **184**, 104–119.
- 21 M. I. Khan, A. Shanableh, S. Shahida, M. H. Lashari, S. Manzoor and J. Fernandez, *Membranes*, 2022, **12**, 263.
- 22 A. R. Kim, M. B. Poudel, J. Y. Chu, M. Vinothkannan, R. S. Kumar, N. Logeshwaran, B.-H. Park, M.-K. Han and D. J. Yoo, *Composites, Part B*, 2023, **254**, 110558.
- 23 Y. Yagizatlı, A. Sahin and I. Ar, *Int. J. Hydrogen Energy*, 2022, **47**, 40445–40461.
- 24 H. Li, G. Zhang, W. Ma, C. Zhao, Y. Zhang, M. Han, J. Zhu, Z. Liu, J. Wu and H. Na, *Int. J. Hydrogen Energy*, 2010, **35**, 11172–11179.
- 25 W. Wu, Y. Li, P. Chen, J. Liu, J. Wang and H. Zhang, *ACS Appl. Mater. Interfaces*, 2016, **8**, 588–599.
- 26 A. Choudhury, P. Patnaik, R. Mondal, S. Sarkar and U. Chatterjee, *Chem. Eng. J.*, 2023, **451**, 138901.
- 27 X. Liu, Y. Li, M. Li, N. Xie, J. Zhang, Y. Qin, Y. Yin and M. D. Guiver, *J. Membr. Sci.*, 2021, **629**, 119282.
- 28 S. Mukhopadhyay, A. Das, T. Jana and S. K. Das, *ACS Appl. Energy Mater.*, 2020, **3**, 7964–7977.
- 29 A. M. Baker, J. H. Dumont, R. Mukundan, S. G. Advani, A. K. Prasad, D. Spornjak and R. L. Borup, *ECS Trans.*, 2017, **80**, 643–650.
- 30 R. Zhiyan, L. Qingbing, H. Youxiu, D. Rui, L. Jia, L. Jia and L. Jianguo, *RSC Adv.*, 2021, **11**, 32012–32021.
- 31 D. Zhao, B. L. Yi, H. M. Zhang and H. M. Yu, *J. Membr. Sci.*, 2010, **346**, 143–151.
- 32 L. Gubler and W. H. Koppenol, *J. Electrochem. Soc.*, 2011, **159**, B211–B218.
- 33 Z. Wang, J. Ren, Y. Sun, L. Wang, Y. Fan, J. Zheng, H. Qian, S. Li, J. Xu and S. Zhang, *J. Membr. Sci.*, 2022, **645**, 120193.
- 34 M. Vinothkannan, B. Son and S. Shanmugam, *J. Mater. Chem. A*, 2022, **10**, 8975–8988.
- 35 A. R. Kim, M. Vinothkannan, K. H. Lee, J. Y. Chu, B. Park, M. Han and D. J. Yoo, *Int. J. Energy Res.*, 2022, **46**, 4835–4851.
- 36 J. Sharma, P. Upadhyay, S. Mishra and V. Kulshrestha, *Int. J. Hydrogen Energy*, 2023, **48**, 10941–10954.
- 37 V. D. C. Tinh and D. Kim, *J. Membr. Sci.*, 2020, **613**, 118517.
- 38 C. D'Urso, C. Oldani, V. Baglio, L. Merlo and A. S. Aricò, *J. Power Sources*, 2016, **301**, 317–325.
- 39 S.-H. Shin, A. Kodir, D. Shin, S.-H. Park and B. Bae, *Electrochim. Acta*, 2019, **298**, 901–909.
- 40 G. V. Buxton, C. L. Greenstock, W. P. Helman and A. B. Ross, *J. Phys. Chem. Ref. Data*, 1988, **17**, 513–886.
- 41 P. Eriksson, A. A. Tal, A. Skallberg, C. Brommesson, Z. Hu, R. D. Boyd, W. Olovsson, N. Fairley, I. A. Abrikosov, X. Zhang and K. Uvdal, *Sci. Rep.*, 2018, **8**, 6999.
- 42 V. Yadav, A. Rajput, N. H. Rathod and V. Kulshrestha, *Int. J. Hydrogen Energy*, 2020, **45**, 17017–17028.
- 43 V. Sharma, P. Upadhyay, N. H. Rathod, J. Sharma, S. Mishra, S. K. Raj, V. Kishore and V. Kulshrestha, *Int. J. Hydrogen Energy*, 2023, **48**, 37784–37795.
- 44 X. Li, C. Liu, H. Lu, C. Zhao, Z. Wang, W. Xing and H. Na, *J. Membr. Sci.*, 2005, **255**, 149–155.
- 45 S. Mishra, J. Sharma, N. H. Rathod and V. Kulshrestha, *ACS Appl. Energy Mater.*, 2022, **5**, 4850–4860.
- 46 B. Kruczek, in *Polyphenylene Oxide and Modified Polyphenylene Oxide Membranes*, Springer US, Boston, MA, 2001, pp. 61–104.
- 47 R. Guan, C. Gong, D. Lu, H. Zou and W. Lu, *J. Appl. Polym. Sci.*, 2005, **98**, 1244–1250.
- 48 R. Mondal, S. Pal, P. Patnaik, D. V. Bhalani, S. K. Gupta, U. Chatterjee and S. K. Jewrajka, *Desalination*, 2022, **531**, 115682.
- 49 J. Su, H. Song and C. Wang, *Adv. Funct. Mater.*, 2019, **29**, 1907154.
- 50 M. E. Culica, A. L. Chibac-Scutaru, V. Melinte and S. Coseri, *Materials*, 2020, **13**, 2955.
- 51 H. Zhao, F. Dong, W. Han and Z. Tang, *Ind. Eng. Chem. Res.*, 2019, **58**, 18055–18064.
- 52 Y. Lee, G. He, A. J. Akey, R. Si, M. Flytzani-Stephanopoulos and I. P. Herman, *J. Am. Chem. Soc.*, 2011, **133**, 12952–12955.
- 53 H. He, X. Lin, S. Li, Z. Wu, J. Gao, J. Wu, W. Wen, D. Ye and M. Fu, *Appl. Catal., B*, 2018, **223**, 134–142.
- 54 G. Xiao, S. Xin, H. Wang, R. Zhang, Q. Wei and Y. Lin, *Ind. Eng. Chem. Res.*, 2019, **58**, 5388–5396.
- 55 S. Cai, D. Zhang, L. Zhang, L. Huang, H. Li, R. Gao, L. Shi and J. Zhang, *Catal. Sci. Technol.*, 2014, **4**, 93–101.
- 56 S. Pal, A. Choudhury, P. Patnaik, S. Sarkar and U. Chatterjee, *Int. J. Hydrogen Energy*, 2022, **47**, 41806–41819.
- 57 E. Sgreccia, J.-F. Chailan, M. Khadhraoui, M. L. Di Vona and P. Knauth, *J. Power Sources*, 2010, **195**, 7770–7775.
- 58 V. Bhadja, U. Chatterjee and S. K. Jewrajka, *RSC Adv.*, 2015, **5**, 40026–40035.
- 59 P. Patnaik, S. M. Hossain, S. Pal, S. Sarkar, R. Sharma and U. Chatterjee, *J. Membr. Sci.*, 2023, **688**, 122105.
- 60 S. Pal, R. Mondal and U. Chatterjee, *Renewable Energy*, 2021, **170**, 974–984.
- 61 F. D. Coms, H. Liu and J. E. Owejan, *ECS Trans.*, 2008, **16**, 1735–1747.
- 62 A. K. Undrajavarapu, R. Sakthivel, A. Sethi, B. M. Dass, S. D. Bhat and V. M. Dhavale, *New J. Chem.*, 2023, **47**, 14960–14971.
- 63 R. Kumar, M. Mamlouk and K. Scott, *RSC Adv.*, 2014, **4**, 617–623.
- 64 B. Zhang, Y. Cao, Z. Li, H. Wu, Y. Yin, L. Cao, X. He and Z. Jiang, *Electrochim. Acta*, 2017, **240**, 186–194.
- 65 D. Han, J. Zhang, C. Wang, P. Hu and Y. Liu, *Int. J. Hydrogen Energy*, 2024, **52**, 11–24.
- 66 V. Elumalai, C. K. K. Sravanthi and D. Sangeetha, *Appl. Nanosci.*, 2019, **9**, 1163–1172.
- 67 G. Sivasubramanian, K. Hariharasubramanian, P. Deivanayagam and J. Ramaswamy, *Polym. J.*, 2017, **49**, 703–709.
- 68 K. H. Lee, J. Y. Chu, A. R. Kim and D. J. Yoo, *Int. J. Energy Res.*, 2019, **43**, 5333–5345.
- 69 D. Han, J. Sun, J. Ge, H. Guo, J. Zhang, C. Wang, P. Hu and Y. Liu, *Int. J. Hydrogen Energy*, 2024, **57**, 301–314.
- 70 M. H. Mohamed, H. Ilbeygi, J. Jaafar, M. Aziz, M. H. D. Othman and M. A. Rahman, *Int. J. Hydrogen Energy*, 2022, **47**, 10736–10746.