Energy & Environmental Science



REVIEW

View Article Online



Cite this: Energy Environ. Sci., 2025, 18, 6934

Challenges in membrane electrode assemblies at elevated temperatures for proton exchange membrane fuel cells: a review

Caizheng Yue, Dab Weibo Zheng, *ab Qiuya Wang, Zhendong Wang, Bing Li, Dab Cunman Zhang oab and Pingwen Ming*ab

Elevated temperatures within membrane electrode assemblies (MEAs) have gained considerable attention as the development of proton exchange membrane fuel cells (PEMFCs) has progressed. For PEMFCs to achieve widespread commercialization, ensuring the stable operation of MEAs at elevated temperatures is essential, but elevated temperatures create several challenges that can negatively impact a PEMFC's performance and lifespan. These challenges include limited mass transfer, charge conduction, and electrochemical reactions, as well as difficulty managing water and gas flows within a system. Elevated temperatures also induce the accelerated degradation of materials and structures, thereby affecting the long-term durability of various MEA components. This review provides an in-depth analysis of these challenges, describing the causes and classification of elevated temperature states within a MEA and the critical changes in transport processes and load response characteristics that they cause. It also discusses the effects of elevated temperatures on the material properties and structural integrity of MEA components (catalyst layer, proton exchange membrane, and gas diffusion layer), and the resulting performance loss mechanism. Recent progress in overcoming challenges caused by the elevated temperatures is also summarised. Finally, based on the identified challenges and current research, this review emphasises the significant role of heat in suboptimal MEA performance and lifespan and offers valuable insights into future directions for optimizing MEA performance and longevity at elevated temperatures. This provides quidance for the further development of PEMFCs for applications at elevated temperatures.

Received 25th February 2025, Accepted 19th May 2025

DOI: 10.1039/d5ee01108f

rsc.li/ees

Broader context

The global environmental challenges associated with the widespread use of fossil fuels underscore the urgent need for sustainable energy solutions. Hydrogen energy, particularly through the proton exchange membrane fuel cell (PEMFC), presents a clean and efficient alternative. The operation of the membrane electrode assemblies (MEAs) at elevated temperatures has garnered increasing attention as PEMFC technology progresses. However, the performance and longevity of PEMFCs under such conditions remain significantly limited, primarily due to the inability of current operating strategies and cell designs to effectively address the new challenges faced by MEA performance at elevated temperatures. A comprehensive understanding of the effects of elevated temperatures on the internal transport processes and degradation mechanisms of MEA components is an essential prerequisite for improving PEMFC operation under such conditions. In this review, we thoroughly analyze the causes, classification, and impacts of the elevated temperature state in the MEA, highlighting the associated challenges in MEA transport processes and component durability and summarizing the recent progress in overcoming elevated temperature limitations. We also propose valuable insights into future directions for improving performance at elevated temperatures, which are expected to be implemented in next-generation PEMFCs.

1. Introduction

The environmental challenges associated with the widespread combustion of fossil fuels are global in scope and require international collaboration. In response, the development of sustainable green energy sources has become critical, with hydrogen energy emerging as a clean and efficient alternative. Proton exchange membrane fuel cells (PEMFCs) are promising power sources that directly convert chemical energy stored in hydrogen molecules into electrical energy through electrochemical reactions. Their advantages, including zero emissions, higher energy conversion efficiencies, and rapid responses, have driven their applications in fields such as vehicles, submarines, and drones.1-3

^a School of Automotive Studies, Tongji University, Shanghai, 201804, China. E-mail: zhengweibo@tongji.edu.cn, pwming@tongji.edu.cn

^b Clean Energy Automotive Engineering Center, Tongji University, Shanghai, 201804,

^c School of Materials Science and Engineering, East China University of Science and Technology, Shanghai, 200237, China

As the power density of fuel cells increases, the heat generation rate of the fuel cell stacks also rises significantly, creating challenges for heat dissipation within the operating temperature range from 60 °C to 80 °C.^{4,5} Increasing the operating temperature has been proposed as a solution to this issue, particularly for high-power PEMFC stacks subjected to long-term operation under heavy-duty conditions. Both theoretical analyses and experimental studies have indicated that operating PEMFCs above 100 °C offers several advantages, such as improved reaction kinetics, simplified water management (only water vapour when exceeding 100 °C), and greater CO tolerance.⁶ Projections by Japan's New Energy and Industrial Technology Development Organization (NEDO) have suggested that fuel cell operating temperatures will rise to 105 °C by 2030 and 120 °C by 2040. Therefore, higher operating temperatures have become a favourable option in the development of PEMFCs. Efforts to enable stable and efficient PEMFC operation at elevated temperatures include the development of modified perfluorosulfonic acid (PFSA) membranes with improved thermal stability and water retention capacities, as well as the use of graphitised carbon nanotube supports that are thermally stable at elevated temperatures.^{8,9} However, the performance and lifetimes of PEMFCs are still significantly limited at elevated operating temperatures, which present challenges to the stable and efficient operation of the membrane electrode assemblies (MEAs), the core components of PEMFCs.

The output performance of PEMFCs, including their steadystate response characteristics (such as polarization and power density curves) and dynamic response characteristics (such as voltage overshoot and recovery time), is significantly limited at elevated temperatures. This is primarily due to the inability of existing materials or operational strategies to accommodate the substantial alterations in transport processes within the MEA caused by increased temperatures. For instance, the lower water content of ionomers within the proton exchange membranes (PEMs) and the catalyst layer (CL) at temperatures near or above the boiling point of water significantly reduces the proton conductivity. 10-14 Increasing the inlet humidity can theoretically mitigate excessive dehydration of ionomers by maintaining a sufficient environmental humidity. 15 However, excess water vapour entering the PEMFC lowers the partial pressure of reactant gases, thereby reducing the reaction rate at active sites and increasing the polarization.¹⁶ Consequently, at elevated temperatures, the coupling between reactant gases and water vapour is intensified, which may render the current water and gas management strategies for maintaining PEMFC performance at low temperatures ineffective. This issue stems from the limited systematic understanding of the transport processes within the MEA at elevated temperatures. Moreover, elevated temperatures can influence the transport processes by modifying critical structural features, such as the three-phase boundary (TPB) and charge transport pathways, primarily due to changes in the physical properties of the ionomer. 17-19 In elevated temperature environments, the dynamic load response of the PEMFC becomes more complex and difficult to control due to significant variations in the water content, reactants, and charged species. 20,21 Thus, it is imperative to systematically analyse critical changes in the transport processes within the MEA due to elevated temperatures to ensure steady-state and dynamic load response characteristics of PEMFCs at elevated temperatures.

Due to the accelerated degradation of MEA components at elevated temperatures, MEA components must have a high thermal stability to ensure the long-term stable operation of PEMFCs in such environments. Ruiu et al. 22 demonstrated that short-term temperature cycling between 90 °C and 120 °C caused more severe degradation of the MEA components than long-term stable operation under automotive-relevant conditions at 80 °C. This led to a fivefold increase in the irreversible performance loss rate of the PEMFC. Factors such as a high electrode potential, humidity fluctuations, and attack by free radicals are commonly considered to be the primary drivers of MEA degradation, while the role of elevated temperatures has often been overlooked. 4,23,24 Elevated temperatures induce structural changes in the materials of MEA components and accelerate other degradation processes within the MEA, both of which result in a more severe, complex, and highly coupled degradation mechanism of the MEA at elevated temperatures. This makes it essential to have a clear understanding of the mechanisms by which elevated temperature degrades MEA components, along with identifying the key degradation factors and characteristics. This knowledge will provide a critical theoretical foundation for developing and selecting materials for MEAs with enhanced stability for extending the service life of PEMFCs that operate at elevated temperatures. However, a systematic understanding of the elevated temperature-induced degradation mechanism in the MEA is lacking.

Hence, this review summarises the challenges associated with MEAs at elevated temperatures in PEMFCs. First, the factors contributing to elevated temperatures and the classification of elevated temperature states in the MEA are discussed. Subsequently, critical alterations in the transport processes, covering reaction gas and water transport, charged species conduction, and load response characteristics, including steady-state and dynamic responses at elevated temperatures, are analysed. Then, the durability challenges of materials and structures in the MEA, focusing on CL, PEM, and gas diffusion layer (GDL), as well as their impacts on PEMFC performance loss are summarised. Moreover, recent progress in overcoming the challenges due to elevated temperatures is summarised, followed by recommendations for future research.

2. Description of elevated temperature states in the MEA

2.1 Causes of elevated temperature states

To better understand how elevated temperatures occur in MEAs, it is necessary to first outline the fundamental operating principle of PEMFCs. As PEMFC technology has advanced, the design of typical fuel cells has evolved into a multilayer structure composed of multiple functional components. As shown in Fig. 1(a), a single PEMFC generally consists of a MEA, an anode bipolar plate (ABP), and a cathode bipolar plate (CBP), with

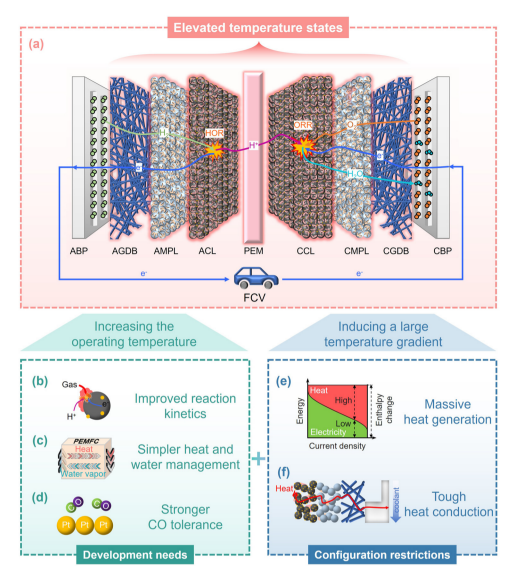


Fig. 1 Illustration of the reason for the MEA maintaining the elevated temperature states. (a) The fundamental operating principle of PEMFCs. (b) Enhanced reaction kinetics. (c) Simplified water and heat management. (d) Improved tolerance of the catalyst to CO impurities. (e) Massive heat generation within the MEA. (f) Low heat dissipation capability of the MEA components.

both plates containing gas channels and coolant channels. This review discusses an MEA without a sealed frame and mainly includes the anode gas diffusion backing, anode microporous layer, anode catalyst layer (ACL), proton exchange membrane (PEM), cathode catalyst layer (CCL), cathode microporous layer, and cathode gas diffusion backing. Electricity generation is driven by half-electrode reactions, specifically the hydrogen oxidation reaction (HOR) at the ACL and the oxygen reduction reaction (ORR) at the CCL.²³ To facilitate these reactions, various reactants must be transported to the reaction sites. Hydrogen (H₂) and oxygen (O₂) are first supplied to the fuel cell and distributed throughout the MEA through gas channels. H₂ passes through the anode gas diffusion backing and anode microporous layer to reach the ACL, where the HOR occurs and releases electrons and protons. The electrons travel through an external circuit to generate electricity, while the protons are conducted by the ionomer on the catalyst, through the PEM,

and finally to the CCL, where the ORR occurs and produces water (H₂O) through a four-electron transfer mechanism.²⁵

Electrochemical reactions, charge-transfer processes, and the phase change of water products release heat.²³ Due to the variety, uneven distribution, and different magnitudes of heat sources within a PEMFC, it is typically necessary to control the operating temperature using fluids such as water or ethylene glycol to maintain a desired temperature and ensure reliable electrochemical reactions. The MEA component is not a perfect thermal conductor, resulting in a temperature gradient between the MEA and the external operating environment.²³ This causes the internal temperature of the MEA to be higher than the external operating temperature. The temperature of the MEA is influenced by two factors: the external operating temperature, where a higher operating temperature will raise the overall PEMFC temperature, and the temperature gradient between the MEA and its external surroundings, where the configuration

Review

of the MEA causes heat accumulation, further elevating its internal temperature. Sections 2.1.1 and 2.1.2 will discuss how these two factors jointly contribute to the elevated temperatures within the MEA.

2.1.1. Development needs. To meet the evolving demands of PEMFCs, higher operating temperatures have become a favoured strategy due to enhanced reaction kinetics, simplified water and heat management, and the alleviation of the detrimental effects of CO impurities on Pt catalysts, as shown in Fig. 1(a)-(c).

Sluggish reaction kinetics create activation overpotentials during the half-electrode reactions, particularly the ORR, which is a major contributor to output voltage losses in PEMFCs. Elevated temperatures can significantly accelerate the reduction of adsorbed oxygen. 26-28 Yano et al. 29 demonstrated that the apparent rate constant of the ORR on Pt in 0.1 M HClO₄ increased exponentially between 30 °C and 110 °C, in accordance with the Arrhenius equation. There was a 3.6-fold increase in the reaction rate when the temperature rose from 80 °C to 120 °C, which improved the electrode reaction kinetics and enhanced the output performance, as shown in Fig. 1(b).

As shown in Fig. 1(c), when operating at temperatures ≤ 80 °C, PEMFCs often struggle with inadequate heat removal using radiator technologies in transport vehicles, especially in heavyduty fuel cell vehicles. This issue necessitates specialised cooling systems, increasing overall costs, but the heat transfer rate improves due to a larger temperature gradient between the fuel cell and the external environment as the operating temperature of the PEMFC increases. This improvement allows the use of existing cooling systems in transport vehicles, enhancing massspecific energy densities, weight efficiency, and overall energy performance. 30-32 Moreover, when operating at ≤80 °C under atmospheric pressure, PEMFCs operate with a dual-phase water system (liquid water and water vapour) to maintain proper humidification within the PEM and prevent flooding in the CL. This presents challenges for water management 33-35 that are not present at higher temperatures (>100 °C), where only a single gaseous phase exists. This avoids the flooding issue, resulting in a more straightforward water management system.

Trace amounts of CO in hydrogen feed gas can significantly reduce a PEMFC's performance due to its strong adsorption onto Pt.31 At higher temperatures, however, the adsorption of CO on Pt is reduced, which improves the CO tolerance, 8,36 as depicted in Fig. 1(d). For instance, the CO tolerance increases from 10–20 ppm at 80 $^{\circ}$ C to 1000 ppm at 130 $^{\circ}$ C and can even reach 30 000 ppm at 200 °C. 37 This eliminates multiple stages of fuel processing and gas cleaning, thereby enabling the use of more cost-effective fuel.

2.1.2. Configuration restrictions. Due to the configuration restrictions of the MEA, the heat generated within the MEA cannot be efficiently dissipated and so it accumulates, resulting in a further rise in the overall temperature of the MEA. Additionally, localised hotspots can form within the MEA under harsh operating conditions. Elevated temperatures are primarily influenced by two factors: the large amount of heat generated within the MEA, and the low heat dissipation capability of the MEA components.

Regarding heat generation, for a PEMFC with an efficiency of 50%, approximately 45% of energy is released as heat.^{79,80} Heat generation has a more significant effect on fuel cells operating at high current densities, as depicted in Fig. 1(e). This substantial heat generation mainly arises from the combined contributions of multiple heat sources, including reversible reaction heat (Q_{rev}) , irreversible reaction heat (Q_{irrev}) , and Joule heat $(Q_i)^{23}$ Fig. 2(a) depicts the influence of the key factors for massive heat generation, where the colour intensity of the arrow is directly proportional to the magnitude of the applied factor. Lighter colours indicate a lower magnitude, whereas darker colours indicate a greater magnitude. Existing commercial Pt/C catalysts do not achieve outstanding power generation efficiency (i.e., catalyst activity),81 leading to the release of a large amount of reaction heat. Furthermore, increasing the current density releases more reaction heat and Q_i due to the higher chemical reaction intensity^{76,82,83} and charged species conduction,⁷⁶ respectively. Decreasing the catalyst loading can increase Qirrey due to significantly higher gas transport resistance. 84-86 Furthermore, the harsh working conditions of the PEMFC itself and reaction flow such as slow electron and proton conduction capacities, poor intercomponent contact, severe gas shortage, 50,78 and insufficient humidification may also generate additional heat.

The magnitude of thermal conductivity or thermal conduction resistance usually increases the heat conduction capacity. Statistical analysis of the thermal conductivities for PEMFC components participating in heat dissipation in the through-plane direction was conducted using 37 relevant studies from 2008 to now, and the results are illustrated in Fig. 2(b). The median thermal conductivities of the CL, microporous layer (MPL), gas diffusion backing (GDB), and bipolar plate (BP) were respectively 0.21, 0.10, 0.42, and 16.65 W m⁻¹ K⁻¹. Except for the BP, the thermal conductivities of other components were so low that they were equivalent to those of some thermal insulation materials. This shows that it is difficult to remove heat generated inside the MEA, as demonstrated in Fig. 1(f). Statistical results of 12 relevant studies from 2013 to 2024 show that the average temperature difference between the PEM and CL was 10.41 K, while that between the GDL and coolant was 9.31 K, as illustrated in Fig. 2(c). Therefore, based on the current MEA configuration, the actual temperature in the MEA will be higher, especially under harsh working conditions.

2.2. Classification of elevated temperature states

NEDO has established 120 °C as the maximum operating temperature target for sulfonic acid-based PEMFCs, implying that elevated temperature operation can range from 80 °C to 120 °C. Considering an approximate 10 °C temperature difference between the MEA and coolant, as shown in Fig. 2(c), the actual MEA temperature may reach 90-130 °C. Moreover, during fuel cell operation at 80 °C, local hotspots caused by membrane pinholes may raise local temperatures to around 140 °C, creating temperature gradients of up to 60 °C,87 but this result has only been obtained using qualitative analysis of thermochromic pigments. To more comprehensively investigate the effects of elevated temperatures, this review investigated the transport

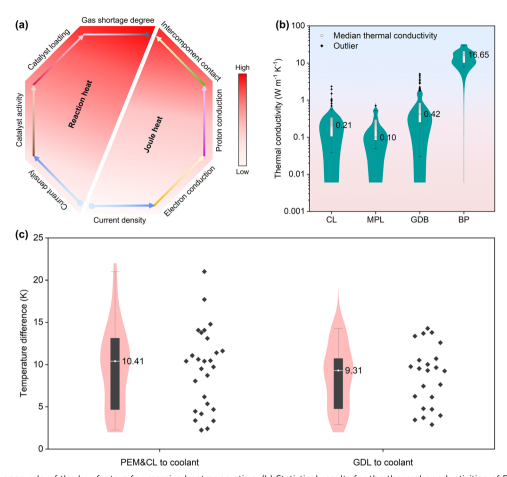


Fig. 2 (a) The influence rule of the key factors for massive heat generation. (b) Statistical results for the thermal conductivities of PEMFC components including the CL, $^{23,38-52}$ microporous layer (MPL), $^{23,38,40,42-47,49,51-54}$ gas diffusion backing (GDB), $^{23,38,40,42-47,49,51,52,55-72}$ and bipolar plate (BP). $^{23,38,40,42-47,49,51,52,73}$ (c) Statistical results for the temperature differences between the PEM and the CL, GDL and coolant. $^{42,46,49,50,52,54,56,74-78}$

and degradation mechanisms in the MEA over the broader temperature range of 90-180 °C.

Within the elevated temperature range, both the water phase and ionomer in the MEA undergo significant phase transitions, as depicted in Fig. 3. When the temperature exceeds the boiling point of water (~110 °C at 143.27 kPa), liquid water converts to vapour, which drastically changes the internal hydration environment of the MEA. This shift affects material utilization and transport-reaction processes. In the vapour phase, the absence of liquid water films on Pt catalyst surfaces reduces the electrochemical surface area (ECSA).88 Ionomer hydration/dehydration causes deformation, leading to a redistribution of the carbon support and disruption of electron network pathways.¹⁷ Schroeder's paradox highlights that an ionomer absorbs less water when it contacts saturated vapour than liquid water, 89 leading to diminished proton conductivity. Moreover, changes in the water environment also influence the degradation of the MEA components. As the temperature rises beyond the ionomer's glass transition temperature (~ 130 °C), its chain mobility increases, which accelerates ionomer instability. 90 Based on these behaviours and current research processes, the MEA's elevated temperature range discussed in this review is classified into three regimes: (1) sub-boiling at moderate temperatures (90–110 °C), defined as the (M) regime; (2) super-boiling at sub-glass transition temperatures (110–130 $^{\circ}$ C), defined as the (H) regime, and (3) super-glass transition at extreme temperatures (130-180 °C), defined as the (E) regime. Correspondingly, operating temperatures are grouped into two ranges: 80-100 °C and 100-120 °C, corresponding to regimes (M) and (H), respectively. Regime (E)corresponds to fault conditions, such as local hotspots. The respective changes in MEA transport characteristics and degradation behaviours across the three different elevated temperature regimes are discussed in the following section.

Critical changes in transport processes and load responses at elevated temperatures

Half-electrode reactions are how PEMFCs power electrical appliances and require a sufficiently fast transport process to guarantee that the PEMFC has the desired steady and dynamic load response capacity. The major transport and electrochemical processes in a MEA are listed in Table 1. The transport processes refer to the movements of reactive species within the different media of a PEMFC. There are two transport processes

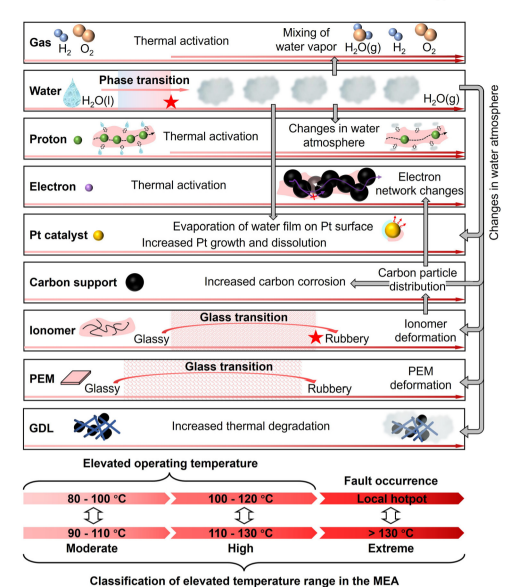


Fig. 3 Classification of the elevated temperature range in the MEA

Table 1 Major transport and electrochemical processes in MEAs

Component	Role	Transport and electrochemical processes
CL	Assist HOR and ORR	Proton conduction in the ionomer network
		Electron conduction in the carbon network
		Water transport in the pore and ionomer networks
		Hydrogen/oxygen diffusion in the pore network and across the ionomer thin film
		$HOR: H_2 \rightarrow 2H^+ + 2e^-$
		ORR: $1/2O_2 + 2e^- + 2H^+ \rightarrow H_2O$
PEM	Permit flow of protons from the anode	Water transport (electro-osmotic drag, back diffusion, and hydraulic permeation)
	to the cathode; stop electrons from traveling	Proton conduction through the PEM
GDL	Ascertain gas diffusion from flow fields to active locations	Hydrogen and water vapor transport (usually diffusion dominates) in the anode
	Drain the water	Oxygen and water vapor transport (usually diffusion dominates) in the cathode Liquid water transport
GDL	active locations	Oxygen and water vapor transport (usually diffusion domin

involved: (i) mass transport processes, including the transport of reactant gases (H2 and O2) from the BP towards the CL, and water (gas or liquid) transport from the BP to CL, from the CL to

BP, and within the PEM and (ii) charged species conduction, including proton conduction from the ACL to CCL through the PEM layer and electron conduction from the ACL to CCL through the conductive network. Key changes in these transport processes and the load response characteristics at elevated temperatures are summarised in the following sections.

3.1. Gas and water mass transport

As illustrated in Fig. 4(a) and (b), compared with low-temperature operation, elevated temperature environments significantly alter the MEA transport of reactant gases and water. Tables 2 and 3 summarise the gas and water transport characteristics at different temperatures, respectively. At elevated temperatures, intrinsic gas diffusivities increase, with the oxygen diffusion coefficient rising from 3.896 \times 10⁻⁶ m² s⁻¹ at 80 °C to 5.347 \times 10⁻⁶ m² s⁻¹ at 90 °C.²⁰ The hydrogen diffusivity in the ionomer phase increased from 2.59 cm 2 s $^{-1}$ at 80 $^{\circ}$ C to 3.48 cm 2 s $^{-1}$ at 95 $^{\circ}$ C. 95 Song et al. 99 indicated that the apparent diffusion-limited current density for the cathode side increased from 1.32 to 2.76 A cm⁻², and the apparent diffusion-limited current density for the anode side increased from 3.01 to 9.63 A cm⁻² as the cell temperature rose from 23 °C to 120 °C. This implies enhanced reactant gas transport

at elevated temperatures, and this effect is more evident for hydrogen transport. Modelling calculations and experimental impendence analysis have also shown that gas transport is enhanced at elevated temperatures, 13,14,20,91,96,100 as shown in Fig. 5(b). Notably, elevated temperature also reduces the solubility of reactant gases in the ionomer, with oxygen solubility undergoing a 1.1-fold decrease from 9.34×10^{-6} mol cm⁻³ at 30 °C to $4.43 \times 10^{-6} \, \mathrm{mol \ cm^{-3}}$ at 80 °C. Zhang *et al.* ¹⁰¹ discovered that the gas transfer resistance increased when the cell temperature was raised from 160 °C to 200 °C, and they emphasised the importance of reactant gas solubility in the ionomer film in the CL. The permeability of reactant gases through the ionomer film is the product of the diffusion coefficient and solubility. Therefore, a higher gas diffusion resistance is observed when the effect of the reduced gas solubility is larger than the effect of the increased gas diffusivity. 101 However, this effect becomes significant only at cell operating temperatures above 160 °C.

The vapour saturation pressure rises exponentially with the temperature 102 because water vapourises at elevated temperatures,

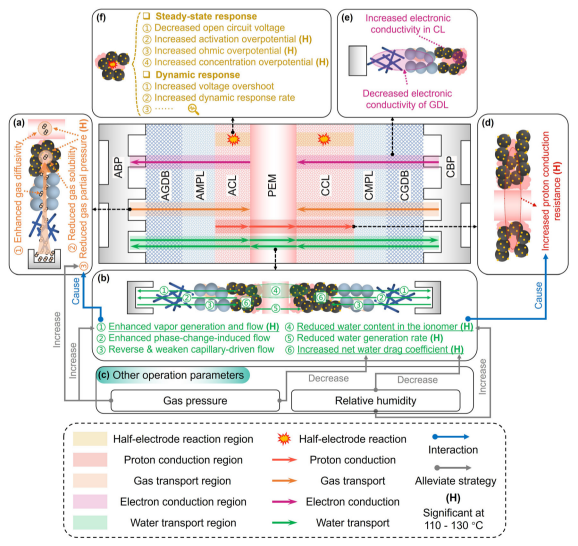


Fig. 4 Evolution patterns of the transport processes and load responses at elevated temperatures. (a) Gas transport. (b) Water transport. (c) Effects of other operation parameters on the gas and water transport. (d) Proton conduction. (e) Electron conduction. (f) Load responses.

Table 2 Gas transport characteristics at different temperatures

Serial number	Operation conditions (anode/cathode)	Items	Values	Ref.
1	_	Oxygen diffusion coefficient	$2.694 \times 10^{-6} \text{ m}^2 \text{ s}^{-1} \ \textcircled{@} \ 60 \ ^{\circ}\text{C}$ $3.896 \times 10^{-6} \ \text{m}^2 \ \text{s}^{-1} \ \textcircled{@} \ 80 \ ^{\circ}\text{C}$ $5.347 \times 10^{-6} \ \text{m}^2 \ \text{s}^{-1} \ \textcircled{@} \ 90 \ ^{\circ}\text{C}$	20
2	_	Oxygen diffusion coefficient in hydrated Nafion membranes	$9.95 \times 10^{-7} \text{ cm}^2 \text{ s}^{-1}$ (a) 30 °C $8.70 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ (a) 80 °C	94
3	_	Oxygen solubility in hydrated Nafion membranes Hydrogen diffusivity in the Nafion	$9.34 \times 10^{-6} \text{ mol cm}^{-3}$ ⓐ 30 °C $4.43 \times 10^{-6} \text{ mol cm}^{-3}$ ⓐ 80 °C $2.59 \text{ cm}^2 \text{ s}^{-1}$ ⓐ 80 °C	
4	Total pressure: 200 kPa, RH: 0.7;	117 membrane Gas partial pressure	3.48 cm ² s ⁻¹ @ 95 °C H ₂ O: 33 kPa @ 80 °C 71 kPa @ 100 °C 139 kPa @ 120 °C O ₂ : 35 kPa @ 80 °C 27 kPa @ 100 °C 13 kPa @ 120 °C N ₂ : 132 kPa @ 80 °C 102 kPa @ 100 °C 48 kPa @ 120 °C	95 16
5	Back pressure: 150 kPa	Gas partial pressure	48 KPa @ 120 °C H ₂ O: 51.87 kPa @ 80 °C 85.13 kPa @ 120 °C O ₂ : 14.81 kPa @ 80 °C 60.28 kPa @ 120 °C	91
6	RH: 0.4/0.4, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5;	Cell mass transfer resistance	At 1.8 A cm ⁻² $54.657 \text{ m}\Omega \text{ cm}^2 \text{ @ } 80 \text{ °C}$ $41.889 \text{ m}\Omega \text{ cm}^2 \text{ @ } 90 \text{ °C}$ $29.534 \text{ m}\Omega \text{ cm}^2 \text{ @ } 95 \text{ °C}$	96
7	RH: 1.0/1.0, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5;	Cell mass transfer resistance	At 1.8 A cm ⁻² : $68.945 \text{ m}\Omega \text{ cm}^2$ @ 80 °C $48.232 \text{ m}\Omega \text{ cm}^2$ @ 90 °C $34.122 \text{ m}\Omega \text{ cm}^2$ @ 95 °C	96
8	Back pressure: 100 kPa	Cell mass transfer resistance	137.3 s m ⁻¹ @ 80 °C 111.68 s m ⁻¹ @ 100 °C 85.75 s m ⁻¹ @ 120 °C	91
9	RH: 1.0/1.0, back pressure (psig): 30/30, flow rate (SLPM): 0.3/1.0;	Cell mass transfer resistance	At 0.91 A cm ⁻² : 0.077 Ω cm ² @ 80 °C 0.072 Ω cm ² @ 100 °C 0.072 Ω cm ² @ 120 °C	13

increasing its partial pressure in the MEA mass transfer structure. This causes a significant drop in the partial pressure of the reactant gases, particularly on the cathode side at high current densities (Fig. 5(a)). Tang et al. 13 reported that the mass transfer resistances were equal at 100 °C and 120 °C, implying a balance between the increased gas diffusivity in the diffusion medium and the reduced gas partial pressure and solubility in the ionomer phase for overall gas transport. Butori et al.16 found that the partial pressure of oxygen decreased significantly from 27 kPa to 13 kPa when the temperature rose from 100 °C to 120 °C, the total air pressure remained constant at 200 kPa, and the relative humidity (RH) remained constant at 70%. This is the main cause of cell voltage losses at a fixed current density. 103 Consequently, the significant role of the decreased gas partial pressure and solubility in the gas transport process at elevated temperature must be considered, especially in the (H) regime due to the complete conversion from liquid water to water vapour. Therefore, at elevated temperatures, the coupling between reactant gas and water transport becomes stronger, mainly due to the increased vapourization of liquid water.

Furthermore, this significant phase transition causes the equilibrium between the membranous water and the liquid water and water vapour in the pore structure to be disrupted at lower temperatures, especially in the (H) regime, where only the equilibrium between the membranous water and water vapour exists. This results in a lower ionomer water content based on Schroeder's paradox.⁸⁹ As depicted in Fig. 5(b), Wang et al.²⁰ suggested that the phase change-induced (PCI) flow increased significantly at high loads as the operating temperature increased to 90 °C due to an increase in the term $d(P_{sat}/T)/dT$ and the temperature gradient ∇T . They also found that the capillary-driven (CD) flow decreased significantly and even reversed due to the reduced content of liquid water within the CCL. The above results are supported by those of Xu et al. 104 and García-Salaberri et al. 10 In comparison, the enhanced PCI flow was stronger than the reverse CD flow, which promoted the expulsion of water in the CCL at elevated temperatures, 20,105 resulting in an insufficient environmental humidity faced by the ionomer phase. This further decreased the water content in the ionomer and made proton conduction difficult. Moreover,

 Table 3
 Water transport characteristics at different temperatures

Serial number	Items	Values	Ref.
1	Net water transport coefficient	At 1.1 A cm ⁻² :	93
	in the Nafion 212 R membrane	$0.016 \ @ \ 80 \ ^{\circ}\mathrm{C}$	
		0.103 @ 100 °C	
		0.394 @ 120 °C	
2	Net water transport coefficient	At 0.9 A cm ⁻² :	93
	in the Aquivion E87-05S membrane	-0.028 @ 80 °C	
		0.080 @ 100 °C	
	· · ·	0.398 @ 120 °C	
3	Water content	At 1.0 A cm ⁻² :	97
		Anode side (8.166 @ 70 °C, 8.537 @ 80 °C, 5.059 @ 90 °C)	
		Cathode side (15.435 @ 70 °C, 14.394 @ 80 °C, 7.836 @ 90 °C) Average (11.410 @ 70 °C, 11.182 @ 80 °C, 6.378 @ 90 °C)	
4	Water content in the membrane	At 0.75 V:	98
4	water content in the membrane	7.922 @ 70 °C	90
		7.862 @ 80 °C	
		7.803 @ 90 °C	
		At 0.65 V:	
		6.782	
		6.748 @ 80 °C	
		6.688 @ 90 °C	
		At 0.45 V:	
		5.407 @ 70 °C	
		5.252 a 80 °C	
		5.008 @ 90 °C	
		At 0.27 V:	
		4.784 @ 70 °C	
		4.530 @ 80 °C	
		4.155 @ 90 °C	
5	Water saturation in the cathode GDL	At 0.75 V:	98
		0.088 @ 70 °C	
		0.089 @ 80 °C	
		0.082 @ 90 °C	
		At 0.65 V:	
		0.133	
		0.128 @ 90 °C	
		At 0.45 V:	
		0.148 @ 70 °C	
		0.158 @ 80 °C	
		0.138 @ 90 °C	
		At 0.27 V:	
		0.151 @ 70 °C	
		0.156 @ 80 °C	
		0.133 @ 90 °C	
6	Water generation rate in the cathode side	At 60 °C:	88
	-	4.47 ml h^{-1} ⓐ a dew point temperature of 30 $^{\circ}$ C	
		6.71 ml h ⁻¹ ⓐ a dew point temperature of 60 °C 12.20 ml h ⁻¹ ⓐ a dew point temperature of 90 °C	
		At 90 °C:	
		1.39 ml h ⁻¹ @ dew point temperature of 30 °C	
		8.50 ml h ⁻¹ ⓐ a dew point temperature of 60 °C	
		11.97 ml h ⁻¹ ⓐ a dew point temperature of 90 °C	
7	Water content in the membrane	6.571 @ 80 °C	91
		6.511 @ 100 °C	
		6.029 @ 120 °C	

the increased net water drag coefficient at elevated temperatures suggests a more severe water shortage for the ionomer phase on the anode side (Fig. 5(f)), 93,97,106 especially in the (H) regime. When the output voltage dropped from 0.75 V to 0.27 V, the membrane dehydration effect caused by temperature increase was enhanced by 13%. Even if the water production increases under a high current density, this water production could not alleviate the dehydration of the ionomer phase at elevated temperatures. Therefore, relying on the back-diffusion of cathode-generated water to alleviate dehydration on the anode

side has a limited effectiveness. As shown in Fig. 5(e), Hou et al.88 found that the water generation rate decreased at elevated temperatures, as measured using a pH meter, especially in the (*H*) regime. Kellegoz *et al.*¹⁰⁷ also indicated that the active reaction area decreased when the cell temperature increased to 85 °C, which decreased the output performance. This was attributed to the presence of liquid water on the surface of the Pt catalyst at lower temperatures, which also provided active sites for the reaction. 108 Elevated temperature promotes the vapourization of liquid water, which reduces the

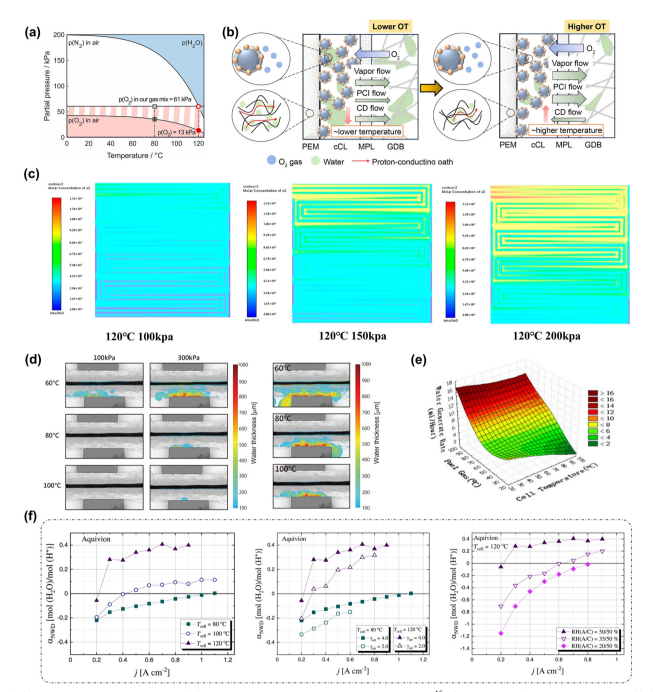


Fig. 5 (a) Partial pressure change patterns of O_2 and water vapor as the cell temperature increases. 16 (b) The impacts of operating temperature (OT) on the water transfer: lower OT and higher OT.²⁰ (c) Molar concentration distribution of oxygen at 120 °C and different pressures.⁹¹ (d) X-ray imaging of MEAs with 1.5 A cm⁻² at 100 and 300 kPa (abs.) and 60, 80, 100 °C and with 3.0 A cm⁻² at 300 kPa (abs.) and 60, 80, 100 °C. 92 (e) Water generation rate at different temperatures.⁸⁸ (f) Effects of operating parameters (temperature, stoichiometric ratio of air, and anode RH) on the net water transport coefficient (α_{NMD}) for cells with the Aguivion membrane.⁹³

active area and subsequently decreases the ORR rate and water production rate.

Other operating conditions, such as gas pressure, stoichiometric ratio, and RH, also influence the transport processes of reactant gas and water (Fig. 4(c)). As demonstrated in Fig. 5(c), an increase in the gas back pressure promotes the regulation of gas flow conditions within the cell, which slows the rate of decrease in the oxygen molar concentration and mitigates the

effect of concentration-differential polarization. 91 Akitomo et al. 92 used X-ray imaging to indicate that when the feed gas was pressurised, the accumulation of liquid water increased (Fig. 5(d)), leading to a decrease in the ohmic overpotential. Therefore, pressurizing reactant gases improves their supply by directly increasing their partial pressures and also suppresses the evaporation of liquid water, thereby alleviating the loss of water within the ionomer phase. However, higher gas pressures

are not always better, as they may pose challenges to the durability and dynamic response characteristics of the components. There must be a minimum gas pressure in the flow field, which depends on various factors, including current density, the structure of the MEA, and the design of the flow field. Increasing the gas RH alleviates the evaporation of liquid water within the MEA by externally supplying additional water vapour, but it also reduces the vapour concentration gradient between the MEA interior and the external environment, thereby mitigating vapour outflow. Altogether, these two aspects help relieve water loss from the ionomer phase and can restore the original electrochemically active surface area (Fig. 5(e)).88 As depicted in Fig. 5(f), at a constant flow rate of H_2 at the anode, α_{NMD} decreases upon decreasing the stoichiometric ratio of air (γ_{air}), but the effect of γ_{air} on α_{NMD} is relatively small. However, α_{NMD} can be significantly decreased by decreasing the RH of hydrogen supplied to the anode. Decreasing the anode RH can lower α_{NMD} at elevated temperatures. Notably, Lu et al. 21 indicated that increasing the inlet pressure also increased the α_{NMD} .

The reaction gas crossover also changes at elevated temperatures, as shown in Fig. 4(a). Butori *et al.* ¹⁰⁹ reported that the $\rm H_2$ permeability increased with temperature and RH, but lower $\rm H_2$ crossover was measured above 100 °C and under humid conditions due to low $\rm H_2$ partial pressure. Consequently, hydrogen crossover is more pronounced in the ($\it M$) regime and may lead to a reduction in the open-circuit voltage (OCV) of the fuel cell, which explains the observed decrease in OCV at elevated temperatures.

At elevated temperatures, the vapourization of liquid water is the fundamental driving force behind hindered reactant gas transport, reduced ionomer water content, decreased electrochemically active surface area, and increased $\rm H_2$ crossover. Therefore, elucidating the phase transition mechanism of water within the MEA and suppressing this process is necessary for achieving reliable water–gas management for operation at elevated temperatures. However, the phase transition of water within pores is also heavily influenced by capillary forces. Jatukaran *et al.* ¹¹⁰ observed that in pores smaller than 10 nm, the liquid evaporation pressure was much lower than that in larger pores, suggesting that the phase transition of water at the micro- and nanoscale may significantly differ from that at the macroscopic scale. It is imperative to thoroughly understand the phase transition between liquid water and water vapour in nanopores.

3.2. Charged species conduction

Proton conduction mainly occurs within the ionomer phase of the PEM and CL, which is closely associated with the internal water content, as illustrated in Fig. 6(a). ¹¹¹ Proton conduction is significantly influenced by water transport, as discussed above. Table 4 summarises the charged species (proton and electron) conduction properties at different temperatures. Yin *et al.* ⁹⁶ reported that as the cell operating temperature increased from 80 °C to 95 °C, the cell ohmic resistance, which is dominated by proton conduction, increased by 14.7% at 100% RH and by 25.1% at 40% RH. Yin *et al.* ¹¹¹ indicated a sharp drop (\sim 63.4% compared to 80 °C) in the proton conductivity when the

temperature increased to 130 °C (Fig. 6(b)). These results show that elevated temperatures can lead to water loss in the ionomer, causing it to dry out and significantly impair proton conductivity, especially for the (H) regime at a low RH. In contrast, Lufrano $et\ al.^{15}$ reported that the proton conductivity of a PEM increased with the temperature, even at 120 °C, as shown in Fig. 6(c). This increase was primarily attributed to the constant and sufficiently high RH environment faced by the PEM, and severe dehydration processes did not occur, which is supported by Butori $et\ al.^{109}$ and Eskandari $et\ al.^{113}$ The differences in proton conductivity between low temperatures and elevated temperatures can be attributed to the differences in thermally-activated properties such as proton mobility or diffusivity at a constant RH. 113

Therefore, maintaining a sufficiently high RH around the ionomer phase is essential. As explained above, increasing the gas pressure and inlet gas RH helps ensure a stable and sufficient humidity within both the CL and PEM to satisfy their constant hydration requirements. In addition, developing novel ionomer materials with enhanced water retention is a critical step in overcoming the limitations of elevated temperatures. Notably, the water absorption and desorption processes of nano-ionomer films within the CL remain poorly understood, with current models relying on empirical equations based on the absorption and desorption observed in the PEM. To alleviate ionomer dehydration in the CL at elevated temperatures, it is essential to first elucidate the phase transition kinetics of water in membranes within the CL and investigate the molecular structures responsible for absorption and desorption. The deformation of ionomer films induced by elevated temperatures also affects the continuity of the proton conduction paths within the CL, while the underlying mechanisms remain unclear.

Limited research has explored changes in electron conduction at elevated temperatures. As depicted in Fig. 4(e), electron conduction is affected by elevated temperature environments mainly within the GDL and CL due to the temperature dependence of electron movement and the altered electron conduction pathways. For the former, Fujita et al. 114 reported that the electronic resistance of carbon nanotubes generally decreased, suggesting a thermally activated process. It can be inferred that elevated temperatures activate and enhance electron transport within the MEA components. For the latter, Taş et al. 115 proposed that the increased electronic conductivity of GDB at higher temperatures was due to enhanced contact between neighbouring carbon fibres resulting from thermal expansion. Additionally, the electron conduction pathways in the CL were affected by the ionomer thermal expansion at elevated temperatures, leading to carbon particle reorientation or migration. This disrupted the conductive network and altered the electronic conductivity of the CL by inducing component deformation.¹⁷⁻¹⁹ Sun et al. 112 investigated variations in the electronic conductivity of GDB, MPL, and CL at different temperatures and found that the conductivity of the GDB and MPL decreased from 827.78 S m⁻¹ to 788.89 S m⁻¹ and 86.11 S m⁻¹ to 80.55 S m⁻¹, respectively, as the temperature increased from 70 °C to 90 °C, while the conductivity of CL increased from 16.67 S m⁻¹ to 22.22 S m⁻¹. Consequently, the

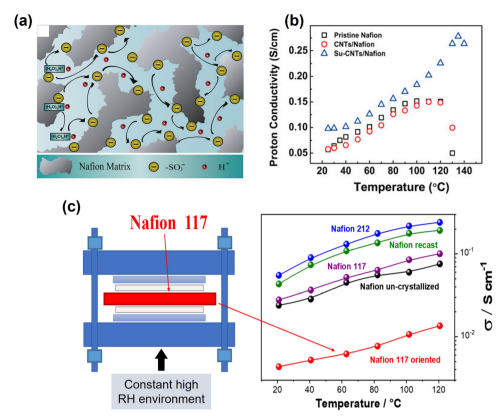


Fig. 6 (a) Schematic illustration of proton transportation in water-hydrated pristine Nafion membranes.¹¹¹ (b) Effect of operating temperatures on the proton conductivity.¹¹¹ (c) Proton conductivities of Nafion recast, Nafion 212, Nafion un-crystallized, and Nafion 117 as received and Nafion 117 oriented membranes as a function of temperature (20-120 °C) at 90% RH.¹⁵

impact of elevated temperatures on the electronic conductivity of MEA components remains controversial, while this effect is negligible compared with the restricted proton conductivity at elevated temperatures.

3.3. Load responses

3.3.1. Steady-state response. The steady-state performance of PEMFCs is typically characterised by the output voltage at a given current density. Shao et al.11 reported severe voltage collapse at elevated temperatures and low RH when the cell temperature was only slightly increased from 92 °C to 93.6 °C, as shown in Fig. 7(a), showing undesirable steady-state output of PEMFCs at elevated temperatures. Table 5 summarises the steady-state output of the cell at different temperatures. Although a few studies have indicated that increasing the cell temperature to 95 °C may slightly improve the output performance, as exemplified by case 14 in Table 5, it is significantly weakened in most cases at elevated temperatures (Fig. 4(f)). This is mainly attributed to the following four aspects: (1) reduction in OCV: the reversible equilibrium voltage tends to decrease at elevated temperatures, as shown in Fig. 7(b), but this decrease is not pronounced. Moreover, as previously mentioned, the substantial increase in hydrogen crossover in the (M) regime further contributes to a reduction in the OCV (Fig. 7(c)). As presented in Table 5, the ratio of OCV at elevated temperatures to that at low temperatures remains approximately 0.94-0.95.

- (2) Increased activation overpotential: elevated temperatures can activate the catalyst, thereby improving electrode reaction kinetics and enhancing the electrochemical reaction rate. 16,96,119 Conversely, elevated temperatures can lead to the evaporation of the water film covering the Pt catalyst, thereby reducing the ECSA, 103 especially in the (H) regime. As presented in Table 5, the ratio of cell voltage at a low current density at elevated temperatures to that at low temperatures is approximately 0.95-0.97.
- (3) Increased ohmic overpotential: elevated temperatures induce ionomer dehydration, which increases the proton conduction resistance. $^{20,96,119-121}$ especially in the (H) regime, but proton conduction is not impeded at a sufficient humidity. As presented in Table 5, the ratio of cell voltage at a moderate current density and elevated temperatures to that at low temperatures is approximately 0.78-0.96.
- (4) Increased concentration overpotential: elevated temperatures may accelerate the gas transport while decreasing the gas partial pressure and solubility in the transport medium, especially in the (H) regime. As presented in Table 5, the ratio of cell voltage at a high current density at elevated temperatures to that at low temperatures is approximately 0.6-0.87.

Increasing the gas pressure and gas RH may help mitigate the performance decline at elevated temperatures. Applying a gas backpressure at elevated temperatures is beneficial in reducing $R_{\rm mt}$, as presented in Fig. 7(d). Notably, the selfhumidification in the thinner membrane is stronger. As shown

Table 4 Charged species conduction properties at different temperatures

Serial number	Operation conditions (anode/cathode)	Items	Values	Ref.
1	_	Cell high-frequency resistance (HFR)	13.382 Ω @ 80 °C 15.441 Ω @ 100 °C 18.382 Ω @ 120 °C	91
2	_	Cell ionic resistance	At 0.8 A cm $^{-2}$: 77.98 mΩ cm 2 @ 70 °C	10
3	RH: 0.4/0.4, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5	Cell ohmic resistance	135.78 mΩ cm ² @ 90 °C At 1.8 A cm ⁻² 82.770 mΩ cm ² @ 80 °C 87.345 mΩ cm ² @ 85 °C 94.238 mΩ cm ² @ 90 °C	96
4	_	Cell ionic resistance	$103.470 \text{ m}\Omega \text{ cm}^2$ ⓐ 95 °C At 0.8 A cm ⁻² $32.83 \text{ m}\Omega \text{ cm}^2$ ⓐ 80 °C $70.21 \text{ m}\Omega \text{ cm}^2$ ⓐ 90 °C	97
5	RH: 1.0/1.0, back pressure (psig): 30/30, flow rate (SLPM): 0.3/1.0;	Cell ohmic transfer resistance	At 0.91 A cm ⁻² : 0.102 Ω cm ² @ 80 °C 0.095 Ω cm ² @ 100 °C 0.115 Ω cm ² @ 120 °C	13
6	RH: 0.4/0.4, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5;	Cell ohmic resistance	At 1.8 A cm ⁻² : 83.24 mΩ cm ² @ 80 °C 87.55 mΩ cm ² @ 85 °C 94.52 mΩ cm ² @ 90 °C 104.13 mΩ cm ² @ 95 °C	96
7	RH: 1.0/1.0, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5;	Cell ohmic resistance	At 1.8 A cm ⁻² : 64.15 mΩ cm ² @ 80 °C 67.30 mΩ cm ² @ 85 °C 70.20 mΩ cm ² @ 90 °C 73.59 mΩ cm ² @ 95 °C	96
8	Fully hydrated before measurement and without external humidification during measurement	Membrane proton conductivity	0.081 S m ⁻¹ @ 40 °C 0.134 S m ⁻¹ @ 80 °C 0.151 S m ⁻¹ @ 120 °C 0.049 S m ⁻¹ @ 130 °C	111
9	RH: constant at 0.9	Membrane proton conductivity	Nafion 117: In-plane $40.0 \pm 0.8 \text{ mS cm}^{-1}$ @ $40 ^{\circ}\text{C}$ $76.5 \pm 1.7 \text{ mS cm}^{-1}$ @ $80 ^{\circ}\text{C}$ $118.4 \pm 1.7 \text{ mS cm}^{-1}$ @ $80 ^{\circ}\text{C}$ $118.4 \pm 1.7 \text{ mS cm}^{-1}$ @ $120 ^{\circ}\text{C}$ Through-plane: $36.7 \pm 1.2 \text{ mS cm}^{-1}$ @ $40 ^{\circ}\text{C}$ $63.6 \pm 1.3 \text{ mS cm}^{-1}$ @ $80 ^{\circ}\text{C}$ $100.6 \pm 1.6 \text{ mS cm}^{-1}$ @ $120 ^{\circ}\text{C}$ Nafion 212: In-plane: $74.1 \pm 1.5 \text{ mS cm}^{-1}$ @ $40 ^{\circ}\text{C}$ $157.0 \pm 2.1 \text{ mS cm}^{-1}$ @ $40 ^{\circ}\text{C}$ $157.0 \pm 2.1 \text{ mS cm}^{-1}$ @ $120 ^{\circ}\text{C}$ Through-plane: $76.1 \pm 1.6 \text{ mS cm}^{-1}$ @ $40 ^{\circ}\text{C}$ $153.1 \pm 2.1 \text{ mS cm}^{-1}$ @ $80 ^{\circ}\text{C}$ $213.1 \pm 2.2 \text{ mS cm}^{-1}$ @ $80 ^{\circ}\text{C}$	15
10	Flow rate (ml min ⁻¹): 200/200, gas type: humidified nitrogen/humidified hydrogen	Electron conductivity	Carbon paper: 827.78 S m ⁻¹ @ 70 °C 780.56 S m ⁻¹ @ 80 °C 788.89 S m ⁻¹ @ 90 °C MPL: 86.11 S m ⁻¹ @ 70 °C 80.56 S m ⁻¹ @ 80 °C 80.55 S m ⁻¹ @ 90 °C CL: 16.67 S m ⁻¹ @ 70 °C 19.44 S m ⁻¹ @ 80 °C 22.22 S m ⁻¹ @ 90 °C	112

in Fig. 7(e), when the temperature is kept at 100 $^{\circ}\text{C}$ and the humidity is reduced from 43% to 30%, the HFR of the Nafion212based cell at a high current density increases by about 70% while that of the Gore12-based cell only increases by about 30%. Therefore, the stable operation of PEMFCs at elevated temperatures should also take membrane thickness into account.

Overall, water phase transitions within the MEA significantly impact the steady-state performance at elevated temperatures.

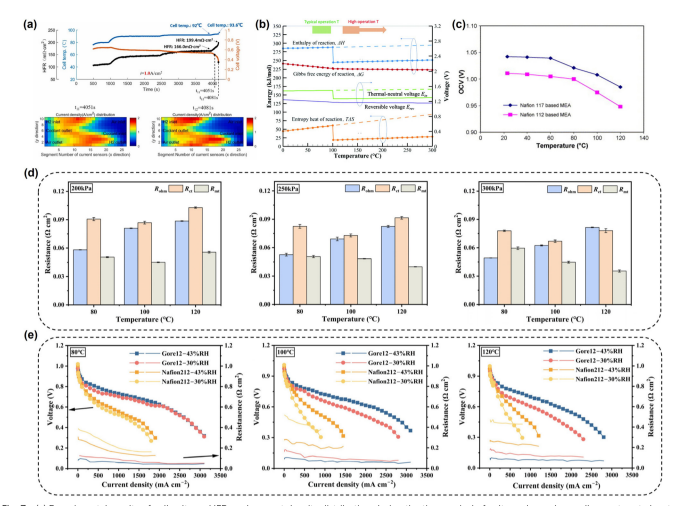


Fig. 7 (a) Experimental results of cell voltage, HFR, and current density distribution during the time period of voltage drop when cell operates at elevated temperatures.¹¹ (b) Correlations between different forms of voltage or energy and temperature based on data. The dashed lines represent the cases of water in the liquid state. 116 (c) Measured OCVs as a function of temperature. 117 (d) Under the conditions of 43% RH, 80/100/120 °C, and 200/250/300 kPa back pressures, respectively, the area resistance values at 0.65 V. 118 (e) Polarization and HFR curves of cells composed of Gore M788.12 and Nafion212 at 30%/43% RH and temperatures of 80 °C, 100 °C, and 120 °C. 118

While PEMFC water management has become more straightforward, the stronger escape tendency of water vapour and stronger coupled transport interactions between water vapour and reaction gases compared with liquid water make it difficult to ensure stable operation of PEMFCs at elevated temperatures.

3.3.2. Dynamic response. As illustrated in Fig. 4(f), the dynamic response characteristics of the PEMFC are also significantly influenced by elevated temperatures, but they have only been investigated within the (M) regime. Table 6 summarises the dynamic output performance of the cell at different temperatures. As illustrated in Fig. 8(a), Wang et al.²⁰ investigated the dynamic response of a PEMFC's output voltage at operating temperatures of 60 °C, 80 °C, and 90 °C and found that at 90 °C, the voltage exhibited the largest overshoot when the current density underwent a stepwise-increase. For instance, when the current density was increased stepwise from 10 mA cm⁻² to 1000 mA cm⁻², the voltage overshoot at 90 °C was 9.52 times greater than that at 60 °C, as shown in Table 6. This was explained by the significant increase in the ohmic resistance of the ACL. Lu et al. 21 also supported this conclusion (Fig. 8(b)), observing that at elevated temperatures, the voltage overshoot increased, and the dynamic response rate also accelerated. Lu et al.21 reported that when the current density was stepwise increased from 0 A cm⁻² to 0.2 A cm⁻², the dynamic response rate at 95 °C was 2.26 times greater than that at 75 °C, as shown in Table 6. The calculation formula for the dynamic response rate is illustrated in eqn (1):

Dynamic response rate =
$$\Delta V \times (t_1 + t_2)$$
 (1)

where ΔV represents the voltage difference between the minimum voltage during the loading process and the steady voltage after loading and t denotes the time for the cell to undergo the dynamic process, including two stages: from loading to voltage undershoot reaching the minimum point, t_1 , and from the minimum point to achieving a steady state again, t_2 . In other words, although the instantaneous voltage overshoot was larger at elevated temperatures, it recovered to stability more quickly.

Increasing the inlet gas pressure reduces the voltage overshoot and dynamic response rate due to an increased partial

Table 5 Steady-state output performance at different temperatures

Serial number	Operation conditions (anode/cathode)	Items	Ratio of elevated temperatures to low temperatures	Relevant processes	Ref.
1	Back pressure (abs, kPa): 200/200, RH: 0.7/0.7	OCV	120 °C to 80 °C: 0.94	Increased hydrogen crossover through PEM	16
2	Back pressure (abs, kPa): 100/100, stoichiometric ratio: very high, RH: 0.8/0.8	OCV	100 °C to 60 °C: 0.94		92
3	Back pressure (abs, kPa): 200/200; stoichiometric ratio: very high, RH: 0.8/0.8	OCV	100 °C to 60 °C: 0.95		92
4	Back pressure (abs, kPa): 300/300; stoichiometric ratio: very high, RH: 0.8/0.8	OCV	100 °C to 60 °C: 0.94		92
5	Back pressure (atm): 3.0/3.0, RH: 1.0/1.0	OCV	120 °C to 60 °C: 0.95		117
6	Back pressure (atm): 3.0/3.0, RH: 1.0/1.0	OCV	120 °C to 60 °C: 0.94		117
7	Back pressure (kPa): 130/120, stoichiometric ratio: 1.7/3, RH: 0.5/0.5	Cell voltage	90 °C to 60 °C:	1. Enhanced the phase transition from liquid water to water vapor in the MEA;	20
			0.95 @ 0.31 A cm ⁻²	2. Reduced water content of the ionomer in the PEM and CL	
			0.93 @ 0.95 A cm ⁻²		
			0.82 (a) 2 A cm ⁻²		
8	Back pressure (abs, bar): 1.0/1.0, RH: 0.5/0.5;	Cell voltage	90 °C to 70 °C:		103
		υ	0.97 (a) 0.2 A cm ⁻²		
			0.92 a 0.8 A cm ⁻²		
			0.60 (a) 1.4 A cm ⁻²		
9	Back pressure (kPa): 100/80, stoichiometric	Cell voltage	95 °C to 75 °C:		21
	ratio: 1.5/2.5, RH: 0.5/0.5;	_	0.97 @ 0.3 A cm ⁻²		
			0.96 @ 1 A cm ⁻²		
			0.87 @ 2 A cm ⁻²		
10	Back pressure (abs, kPa): 200/200,	Cell voltage	120 °C to 80 °C:		118
	RH: 0.43/0.43;		0.95 @ 0.5 A cm ⁻²		
			0.90 @ 1.3 A cm ⁻²		
			0.69 @ 2 A cm ⁻²		
11	Back pressure (abs, kPa): 250/250,	Cell voltage	120 °C to 80 °C:		118
	RH: 0.43/0.43;		0.97 @ 0.5 A cm ⁻²		
			0.93 @ 1.3 A cm ⁻²		
10	Parls (alan I-Pa) 200/200	Call scales as	0.83 @ 2 A cm ⁻² 120 °C to 80 °C:		110
12	Back pressure (abs, kPa): 300/300,	Cell voltage	0.97 (a) 0.5 A cm ⁻²		118
	RH: 0.43/0.43;		0.92 @ 1.3 A cm ⁻²		
			0.92 @ 1.3 A cm ⁻²		
13	Back pressure (abs, kPa): 300/300,	Cell voltage	0.87 (a) 2 A cm 120 °C to 80 °C:		118
10	RH: 0.3/0.3;	Cen voitage	0.99 @ 0.2 A cm ⁻²		110
	M1. 0.5/0.5,		0.78 (a) 1 A cm ⁻²		
14	RH: 0.4/0.4, back pressure (bar): 1.5/1.5, flow rate (SLPM): 0.7/2.5;	Cell voltage	95 °C to 80 °C:	1. Enhanced electrode reaction kinetics in the CL	96
	110W 14th (DDI 141). 0.7/2.3,		1.02 @ 0.3 A cm ⁻² 1.09 @ 2.1 A cm ⁻²	2. Enhanced gas diffusivity in the MEA	

pressure of reactants, which creates a more abundant supply of reactants and helps slow voltage undershoots. 21,122 Increasing the operating pressure induces the condensation of water vapour, alleviating the ionomer dehydration and the reduction in output performance.20 Overall, similar to the steady-state response characteristics, the phase transitions of water at elevated temperatures also need to be taken seriously during dynamic load. Notably, increasing the reactant gas pressure can optimise the dynamic response characteristics of PEMFCs. However, the interaction between elevated temperatures and variable local factors (e.g., temperature, water content, reactant concentration, and charge) during vehicle operation can considerably impair fuel cell components, particularly the PEM. A wellrounded strategy that optimally balances performance and durability is still lacking. It is crucial to comprehensively understand the sensitivity of the dynamic response characteristics of PEMFCs at elevated temperatures to various operational parameters, but such studies are limited. The main goal of such research is to

ensure optimal integration and compatibility of multiple components within a PEMFC, identify refined operational strategies and material designs, and ultimately achieve a stable internal state with minimal fluctuations.

4. Challenges in material and structure durability and their impacts on performance loss

To better understand how elevated temperatures affect MEA durability, MEA failure modes at elevated temperatures are discussed in this section in terms of individual components.

4.1. Catalyst layer durability

The CL of PEMFCs is a multi-component/multifunctional medium that consists of the tailored assembly of electrochemical catalyst and ionomer phase. Elevated temperatures can directly

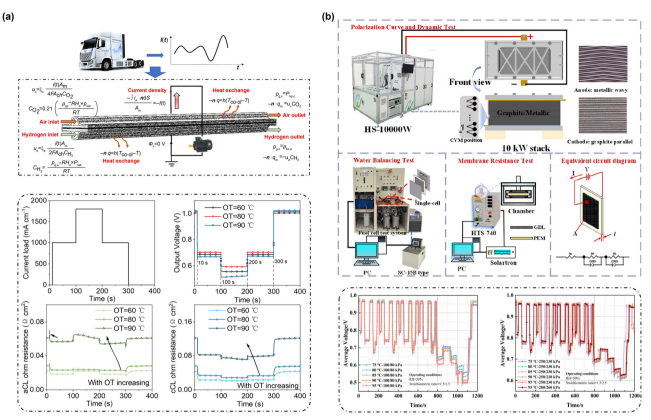
Table 6 Dynamic output performance at different temperatures

Serial number	Operation conditions (anode/cathode)	Items	Ratio of elevated temperatures to low temperatures	Relevant processes	Ref
1	Back pressure (kPa): 130/120, stoichiometric ratio: 1.7/3, RH: 0.5/0.5;	Voltage overshoot	90 °C to 60 °C:	Ionomer dehydration	20
	,		9.52 ⓐ 10 to 1000 mA cm ⁻² 6.80 ⓐ 1800 to 1000 mA cm ⁻²		
			$0.77 \ \text{@} \ 1000 \ \text{to} \ 10 \ \text{mA cm}^{-2};$		
2	Back pressure (kPa): 100/80, stoichiometric ratio: 1.5/2.5, RH: 0.5/0.5	Voltage overshoot	95 °C to 75 °C:	1. Ionomer dehydration	21
	111. 0.5/0.5		0.90, 1.10, 1.04, 1.22 @ 0 to 0.2 A cm ⁻²	2. Inadequate oxygen supply	
			1.06, 1.09, 1.18, 1.22 @ 0 to 0.4 A cm ⁻²	78 - 77	
			1.11, 0.76, 1.24, 1.29 @ 0 to 0.7 A cm ⁻²		
			1.43 @ 0 to 1.0 A cm ⁻²		
			1.31 @ 0.7 to 1.0 A cm ⁻²		
			1.44 @ 1.0 to 1.4 A cm ⁻²		
ā	D. J (I.D.) 250/220	77.10 · · · ·	1.27 @ 1.4 to 1.6 A cm ⁻² ;	4 I	24
3	Back pressure (kPa): 250/230, stoichiometric ratio: 1.5/2.5, RH: 0.5/0.5	Voltage overshoot	95 °C to 75 °C	1. Improved membrane hydration	21
	12.11 010,010		0.82, 1.00, 0.95, 1.00 @ 0 to 0.2 A cm ⁻²	2. Increased partial pressure of reactants	
			0.92, 1.00, 1.08, 1.00 @ 0 to 0.4 A cm ⁻²		
			0.91, 0.91, 0.91, 0.88 a 0 to 0.7 A cm ⁻²		
			0.83@ 0 to 1.0 A cm ⁻²		
			0.87		
			0.80 (a) 1.0 to 1.4 A cm 0.80 (a) 1.4 to 1.6 A cm ⁻² ;		
4	Back pressure (kPa): 100/80,	Dynamic	95 °C to 75 °C:	1. Ionomer dehydration	21
	stoichiometric ratio: 1.5/2.5, RH: 0.5/0.5;	response rate			
			2.26, 1.00, 0.73, 0.86 @ 0 to 0.2 A cm ⁻²	2. Inadequate oxygen supply	
			1.35, 1.00, 1.43, 0.88 @ 0 to 0.4 A cm ⁻²		
			$0.57, 0.35, 0.91, 2.06 \stackrel{\textcircled{a}}{=} 0 \text{ to } 0.7 \text{ A cm}^{-2}$		
			1.67 @ 0 to 1.0 A cm ⁻²		
			2.14 ⓐ 0.7 to 1.0 A cm ⁻² 1.78 ⓐ 1.0 to 1.4 A cm ⁻²		
			1.78 (a) 1.0 to 1.4 A cm 1.10 (a) 1.4 to 1.6 A cm ⁻²		
5	Back pressure (kPa): 250/230,	Dynamic	95 °C to 75 °C:		21
3	stoichiometric ratio: 1.5/2.5, RH: 0.5/0.5;	response rate	33 0 10 73 0.		21
	. ,		1.60, 0.84, 0.60, 0.72@ 0 to 0.2 A cm ⁻²	 Improved membrane hydration 	
			0.57, 0.45, 0.78, 0.67 @ 0 to 0.4 A cm ⁻²	2. Increased partial pressure of reactants	
			0.38, 0.45, 0.41, 0.67 (a) 0 to 0.7 A cm ⁻²		
			0.46 @ 0 to 1.0 A cm ⁻²		
			0.93 ⓐ 0.7 to 1.0 A cm ⁻² 0.79 ⓐ 1.0 to 1.4 A cm ⁻²		
			0.79 (a) 1.0 to 1.4 A cm 0.73 (a) 1.4 to 1.6 A cm ⁻²		
			0.73 @ 1.4 to 1.0 A CIII		

degrade the catalyst layer or synergistically exacerbate other degradation factors, leading to a reduction in the performance of its components through complex mechanisms. Table 7 reviews the stability of the catalyst layer at different temperatures. Consequently, it is imperative to examine the impact of elevated temperatures on the degradation of the Pt catalyst, carbon support, and ionomer phase.

4.1.1. Pt catalyst degradation. The following mechanisms have been identified as the most prevalent during the deterioration of the Pt catalyst: dissolution, agglomeration, detachment, and Ostwald ripening of Pt nanoparticles. 139,140 Existing research has mainly investigated the effect of elevated temperatures on Pt catalyst degradation in terms of Pt catalyst growth (Fig. 9(a)) and Pt dissolution (Fig. 9(b)).

As illustrated in Fig. 9(a), elevated temperatures can directly induce Pt catalyst growth, but this direct induction mechanism is significantly modulated by specific elevated temperature levels and the surrounding gas atmosphere. Sellin et al. 124 reported that increasing temperatures directly induced an increase in the Pt catalyst growth in the entire elevated temperatures range and a H2-containing atmosphere. The reducing effect of H2 gas removed surface oxides on Pt catalysts, which disrupted the protective oxide layer and supported the subsequent crystallite reconstruction and grain boundary diffusion. Sellin et al. 124 reported an approximately 4.9% loss of the Pt active surface area and a 60.0% increase in the Pt apparent mean size after thermal sintering at 100 °C. Hasché et al. 128 demonstrated significant Pt catalyst growth (~78.57%) under a



(a) Structure schematics of model and boundary condition settings and current load dynamic profile and dynamic responses of output voltage, ACL average proton ohmic resistance, and CCL average proton ohmic resistance at different OTs.²⁰ (b) Experimental system schematic diagram including the polarization curve test, water balancing test, membrane electric resistance test, and equivalent circuit diagram and dynamic response results at different operating temperatures and inlet pressures during the loading process.²¹

N₂ atmosphere at 160 °C for 155 h because the Pt catalyst crystallite growth was controlled by the interparticle distance, as indicated in Fig. 9(c). A decrease in the interparticle distance resulted in a shorter time for the Pt catalyst particle diffusion/ migration and coalescence. Therefore, the direct elevated temperature induced growth of Pt under a N2 atmosphere was only observed in the (E) regime. However, this direct induction was more challenging under an air atmosphere due to the protective effect of the Pt oxide layer. Shao et al. 141 reported no noticeable platinum particle sintering during long-term (3000 h) exposure to dry air at 125–195 $^{\circ}$ C. Overall, the direct elevated temperature induced growth of the Pt catalyst on the anode side in the PEMFC-matched elevated temperature range is acceptable, while it can be considered negligible on the cathode side.

Catalyst durability studies have also indicated that elevated temperatures can accelerate the Pt catalyst growth during potential cycling (Fig. 9(a)). 4,126,142 The influence of elevated temperatures on the accelerated growth of Pt catalysts under voltage cycling appears to increase linearly (Fig. 9(d)), suggesting that Pt catalyst growth is accelerated within the entire elevated temperature range. Elevated temperatures do not trigger explosive Pt growth, even though the underlying acceleration mechanism remains poorly understood. Bi et al. 143 reported that a low RH stimulates Pt growth at 60 °C. To our knowledge, the coupled effect of elevated temperatures and RH on Pt catalyst growth during operation has not been studied.

In terms of Pt dissolution at elevated temperatures, Fig. 9(b) shows that elevated temperature has an accelerating effect, exacerbating Pt catalyst dissolution under electrochemical potential operations within the entire elevated temperature range. 125,144-146 Yang et al. 125 observed a denser Pt band in the membrane after potential cycling at 90 °C (Fig. 9(e)). Borup et al. 138 indicated that the Pt dissolution reaction was endothermic, causing the Pt solubility to increase at elevated temperatures, following the Arrhenius relationship (Fig. 9(f)). Huang et al. 147 suggested that this was attributed to the impact of the operational temperature on the collective surface tension of Pt, which increased both the net rate of Pt decomposition and the net rate of particulate dissolution. Yue et al. 132 reported approximately 12.3% Pt mass loss after 1000 h on-road heavy-duty transportation with a coolant temperature of 92 °C. At low temperatures, a high RH promotes Pt catalyst dissolution due to accelerated Pt ion transport in the ionomer with larger and more abundant water channel networks. 143 Similar to Pt growth, the coupling effect of elevated temperature and RH levels on Pt dissolution during potential cycling has not yet been studied. Notably, Pt redeposition equally affects the available amount of Pt catalyst. For instance, at low temperatures, Đukić et al.148 reported that the rates of Pt catalyst dissolution and redeposition both increased with the temperature at a constant potential window, while the overall extent of Pt loss remained minimal. Nevertheless, it is

Review

 Table 7
 Stability of the CL at different temperatures

Serial number	Durability cases and corresponding conditions (anode/cathode)	Items	Values	Re
1	Durability test under a constant current density of 0.4 A cm ⁻² : operating temperature: 100 °C, back pressure (kPa) 150, gas type: H ₂ /air, RH: 0.25,	ECSA loss	15.6%	123
2	duration: 480 h; Thermal sintering test: temperature: 100 °C, gaseous atmosphere: 3% H ₂ /He, sample: Pt/Vulcan catalyst (40 wt% Pt);	Pt apparent mean size increase	60.0%	124
	,	Pt surface area loss	4.9%	
	Voltage cycling: square wave potential cycle: 10 s at 0.4 V, 10 s at 0.95 V (20 s per cycle), RH: 1.0, gas: 0.5 SLPM 4% H ₂ /0.5 SLPM N ₂ , pressure: atmospheric pressure, cell temperature: 80 °C and 90 °C;	ECSA loss	80 °C:	125
			42.2% @ 10 000 cycles	
			90 °C: 50.0% @ 10000 cycles;	
		Mass activity loss	80 °C:	
			33.9% @ 10 000 cycles 90 °C:	
			64.1% @ 10000 cycles;	
ļ	Voltage cycling: voltage range: 0.1–0.96 V, scan rate 10 mV s ⁻¹ , duration: 300 cycles, temperature: 60 °C,	Pt particle growth	60 °C:	126
	80 °C, 100 °C, 120 °C;		22.2% @ 300 cycles	
			80 °C:	
			49.4% @ 300 cycles 100 °C:	
			85.2% @ 300 cycles	
			120 °C:	
5	Voltage cycling: voltage range: 0.7–0.9 V, time of one cycle: 60 s, duration: 5000 cycles, RH: 0.25, temperature: 80 $^{\circ}$ C and 120 $^{\circ}$ C	Pt surface area loss	117.3% @ 300 cycles 8.7% @ 80 °C	127
	temperature: 80°C and 120°C		44.4% @ $120\ ^{\circ}\mathrm{C}$	
5	Thermally induced growth test: atmosphere: N_2 , duration: 155 h, temperature: 80 $^{\circ}$ C and 160 $^{\circ}$ C;	Pt particle growth	7.14% a 80 °C	128
7	Thermal corrosion test: atmosphere: air, duration: 3000 h, sample: Pt/BP2000 catalyst with different Pt loadings (40 wt%, 60 wt%, and 80 wt%), temperature: 125 °C;	Maximum fraction of carbon consumed	78.57% @ 160 °C; 17.78% @ 40 wt%	129
	,		33.43% @ 60 wt%	
8	Thermal treatment: atmosphere: air, sample:	Carbon weight loss	40.36% @ 80 wt%; 2.6%	130
)	Pt (40 wt%)/XC 72 catalyst temperature: 150 °C Durability test under a constant current density of	Carbon weight loss ECSA loss	9.59% @ 65 °C	131
	1.6 A cm $^{-2}$: operating temperature: 65 °C and 90 °C, gas type: H_2/O_2 , dew point temperature: 65 °C, stoichiometries: 1.5/2.5, duration: 100 h			
10	Durability test under a constant potential of 0.9 V:	Carbon mass loss rate	13.93% @ 90 °C 5.71 $\mu g \ h^{-1}$ @ 75 °C	4
	operating temperature: 75 $^{\circ}$ C, 80 $^{\circ}$ C and 90 $^{\circ}$ C		7.51 μg h ⁻¹ @ 80 °C	
			12.41 μg h ⁻¹ @ 90 °C	
11	On-road heavy-duty transportation: operating	Pt mass loss	12.3%	132
	temperature of 92 °C, duration: 1000 h;	Pt particle growth Ionomer degradation	3.92% Main-chain: 6.5%	
		ECSA loss	Side-chain: 14.2%; 44.75%	
12	NEDC near-water boiling temperature: operating temperature: 95 °C, inlet pressure (kPa): 280/260, duration: 100 h;	Pt particle growth	Anode:	133
	duration. 100 ii,		76.8–93.9% Cathode:	
		Catalant laws (1.1.1 a.s.	83.7-117.8%	
		Catalyst layer thickness	Anode: 3.87–37.8%	
			Cathode:	
		ECCA la	6.78-26.8%	
		ECSA loss	11.6-82.4%	

Table 7 (continued)

Serial number	Durability cases and corresponding conditions (anode/cathode)	Items	Values	Ref
13	A combined OCV hold and RH cycling testing: gas: H_2/air (1000/1000 sccm), pressure: ambient, duration: \sim 200 h, temperature: 95 °C;	Pt particle growth	Anode:	134
	, 1		47.6–85.7% Cathode: 195.2–180.9%	
14	Thermal annealing: sample: Nafion film, gas atmosphere: dry N_2 , duration: 1 h, temperature: 120 °C;	Ionomer proton conductivity loss	195.2–180.9% 20.91–27.6%	135
15	Thermal annealing: sample: Nafion film, gas atmosphere: vacuum, duration: 1 h, temperature: 110 °C and 146 °C;	Ionomer proton conductivity loss	81% @ 110 °C	136
			94% \textcircled{a} 146 $^{\circ}\text{C}$	
16	Thermal treatment: sample: MEA, gas atmosphere: vacuum, temperature and duration: 90 °C for 10 min, 140 °C for 10 min, 140 °C for 30 min	Decrease of oxygen transport resistance in the CCL	1.6% @ 90 °C for 10 min	137
	,		7.4% @ 140 °C for 10 min 16.0% @ 140 °C for 30 min;	
17	Thermal annealing: sample: Nafion film, gas atmosphere: vacuum, duration: 24 h, temperature: $60\ ^{\circ}\text{C}$ and $140\ ^{\circ}\text{C}$	Ionomer Young's mod- ulus increment	120.4% @ 60 °C	24
	00 0 and 140 0		214.1% (a) 140 $^{\circ}\mathrm{C}$	

essential to investigate the effect of elevated temperatures on Pt catalyst redeposition (e.g., degree and location) to explicitly understand the overall Pt dissolution.

4.1.2. Carbon support corrosion. Carbon blacks like Vulcan XC72, acetylene black, and Ketjen Black are widely used carbon supports for PEMFCs, but they often undergo carbon corrosion. The characteristics of carbon corrosion at elevated temperatures, as investigated in existing studies, 4,94,129-131,149-152 can be summarised as follows. Firstly, an environment containing oxygen or hydrogen is a prerequisite, such as air or hydrogen-containing composite gases. Secondly, elevated temperatures (both without and with a high electrode potential) significantly enhance the reaction rate between the carbon matrix and oxygen or hydrogen species to generate gaseous products that exacerbate the extent of carbon corrosion. Thirdly, carbon corrosion is significantly pronounced only during Pt catalysis. Overall, the effect of elevated temperatures on carbon corrosion can be categorised as either directly inducing chemical corrosion (Fig. 10(a)) or accelerating electrochemical corrosion (Fig. 10(b)).

As depicted in Fig. 10(a), under a H₂/O₂ atmosphere, elevated temperature promotes carbon corrosion, leading to the corrosion of the part of the carbon support in contact with Pt catalysts, which occurs within the (H) and (E) regimes but not at lower temperatures. Andersen et al. 149 found that the degradation of the support without a Pt catalyst was minimal (only $\sim 1\%$ weight loss) even at temperatures of up to 200 °C, while degradation of the carbon support with Pt was significantly higher, indicating that carbon corrosion was catalysed by Pt. Stevens et al. 129 reported that 40.36% of carbon mass was consumed after 3000 h of thermal corrosion at 125 °C when the Pt catalyst loading was 80 wt%. Sellin et al. 130 indicated that only 2.6% of the carbon support (Vulcan XC-72) was corroded after thermal

treatment at 150 °C, implying that the degree of thermal corrosion resistance of the carbon support itself is equally critical and that the Vulcan XC-72 support has good thermal corrosion resistance. Guterman et al. 151 suggested that narrowing the size dispersion of Pt and of support particles and improving the uniformity of Pt distribution increased the exposure of platinum to carbon and resulted in more uniform carbon oxidation. Under an O₂ atmosphere, the product of carbon corrosion is either CO or CO₂. 94,130,152 Sellin et al. 130 demonstrated that the carbon support could also be directly corroded at 423 K under a 3% H₂/ Ar flow, which generated methane or other hydrocarbons. Notably, the rate of change in the thermal environment faced by the carbon support also affects the degree of carbon corrosion. As depicted in Fig. 10(c), Baturina et al. 153 conducted thermogravimetric analysis (TGA) at heating rates from 0.1 to 10 °C min⁻¹ and showed that as the heating rate decreased, the percent mass loss increased. This indicated that Pt catalysed the oxidation of carbon during isothermal aging. Therefore, it can be concluded that the contribution of instantaneous thermal overshoot induced within the CL during dynamic processes is not a significant contributor to the thermal corrosion of carbon supports.

In terms of how elevated temperatures accelerate electrochemical corrosion, Fig. 10(b) shows that elevated temperatures further intensify corrosion at pre-existing electrochemical corrosion sites in carbon, thereby exacerbating the extent of electrochemical corrosion over the entire elevated temperature range (Fig. 10(d)). Lochner et al.4 reported that the carbon mass loss rate reached 5.71 $\mu g h^{-1}$ at 75 °C and 12.41 $\mu g h^{-1}$ at 90 °C during durability tests under a constant potential of 0.9 V. Zhang et al., 94 Roen et al., 154 and Lim et al. 152 supported the above results, and Stevens et al. 155 suggested that H₂O played a significant role in carbon corrosion. Therefore, high RH levels

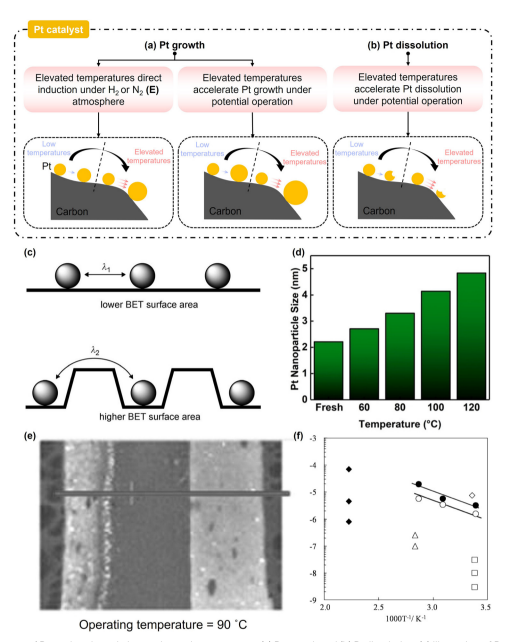


Fig. 9 Schemic diagram of Pt catalyst degradation at elevated temperatures: (a) Pt growth and (b) Pt dissolution. (c) Illustration of Pt particle growth as a function of the distance between the nanoparticles. 128 (d) Alterations in the Pt catalyst size as the cell temperature increases during voltage cycling (0.1-0.96 V).4 (e) Denser Pt microscopy images at the MEA cross-section. 125 (f) Logarithm of Pt solubility as a function of inverse temperature. 138

and enhanced water production at a high current density are unsuitable for ensuring that the carbon support remains stable at elevated temperatures, as shown in Fig. 10(a) and (b). Moreover, there has not been a comparative analysis at elevated temperatures between chemical corrosion and electrochemicalinduced corrosion of carbon supports.

4.1.3. Ionomer degradation. The ionomer has a phaseseparated morphology, formed by the hydrophobic nature of the carbon-fluorine backbone and the hydrophilic sulfonic acidfunctionalised side chains, as illustrated in Fig. 11(a). Within certain hydrophobic domains, enhanced backbone ordering facilitates the formation of crystalline phases, while the interconnected hydrophilic domains create continuous proton

conduction pathways. As depicted in Fig. 11(a), at low temperatures, the nanoscale structure of the ionomer does not undergo significant changes upon treatment, but recent studies have explored the effects of elevated temperatures on the structural and morphological properties of ionomer films. The results have revealed significant changes in their hydration, crystallinity, and hydrophilic-hydrophobic phase separation, 136,137,156-158 especially in the (H) and (E) regimes approaching or exceeding the glass transition temperature. Kusoglu et al. 157 indicated that annealing at 146 °C for 60 min induced the crystalline order corresponding to the CF2 chains of polytetrafluoroethylene (PTFE) crystallites, which formed crystallites and reduced the water uptake due to an increase in the backbone physical

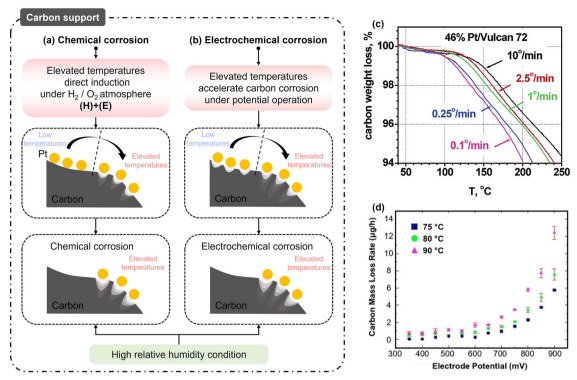


Fig. 10 Schemic diagram of carbon support corrosion at elevated temperatures: (a) chemical corrosion and (b) electrochemical corrosion. (c) TGA data for 46% Pt/C measured at heating rates of 0.1, 0.25, 1, 2.5, and 10 °C min^{-1.153} (d) Decline in carbon content against the cell potential at varying temperatures.

crosslinking. Modestino et al. 156 supported the above findings. Paul et al. 136 reported that the ionomer proton conductivity loss reached 81% and 94% at annealing temperatures of 110 °C and 146 °C, respectively. Consequently, elevated temperatures can increase the crystallinity of the hydrophobic region and shrink the hydrophilic region (Fig. 11(a)). Alentiev et al. 159 reported the presence of the ether (-S(O)2-O-) groups after annealing Nafion films at 95 °C, which further weakened the water retention capacity of the ionomer film. However, Wang et al.23 reported that the Nafion molecule was only dehydrated at a higher temperature of ~ 100 °C and the terminal SO_3^- group was protonated to SO_3H instead of forming an ether. Chu et al. 160 reported no significant differences in the infrared reflectance absorption spectra of the Dow 560 PFSA ionomer when heated below 300 $^{\circ}\text{C}.$ Therefore, the ionomer evolution within the (M) regime needs to be further investigated.

Nevertheless, elevated temperatures alter the properties of ionomers, including decreasing the thermal expansion coefficient (Fig. 12(a)),19 increasing Young's modulus (Fig. 12(b)),24 decreasing the proton conductivity (Fig. 12(c)), 136,158 increasing the surface hydrophobicity (Fig. 12(d)), 136,158,161 and improving the oxygen transmission capacity (Fig. 12(e)). 137 However, the underlying mechanisms between the evolution of phases, clusters, and molecular structures within ionomers and elevated temperatures are not fully understood, making it difficult to understand changes in ionomer properties. Most existing studies on directly-induced ionomer degradation at elevated temperatures have been conducted under vacuum or dry inert gases, but since the actual ionomer phase within the MEA typically exists in a humidified environment, the role of RH in this degradation process needs to be better understood.

Mechanical fatigue and chemical degradation processes are affected over the entire elevated temperature range. The mechanical degradation of ionomers is primarily caused by their frequent swelling and shrinkage during cyclic hydrothermal stress. Chang et al. 162 suggested that plastic strain within an ionomer could accumulate during cyclic changes in RH. Rong et al. 163 reported that the deformation of the Nafion ionomer at elevated temperatures during RH cycling was larger than that at lower temperatures, which was corroborated by Tang et al. 164 Therefore, a greater change in plastic strain occurs and propagates within the ionomer phase at elevated temperatures due to larger deformation, which accelerates mechanical degradation of the ionomer, as illustrated in Fig. 11(b). The accumulation of plastic strain may result in the internal breakdown of the ionomer when reaching the breakdown strain, which may significantly impair the structural stability of the CL. The mechanisms of ionomer chemical degradation in PEMFCs involve the formation of hydroxyl and hydroperoxyl radicals (OH• and OOH•) caused by hydrogen peroxide (H₂O₂) formation on the catalyst surface via the 2e ORR. Sethuraman et al. 165 reported the rates of H₂O₂ formation in the anode side of a PEMFC at different temperatures and showed that increasing the cell temperature from 25 °C to 95 °C significantly enhanced the rates of H₂O₂ formation, and the rate of H₂O₂ formation on the cathode side was the same. 165 The anode and cathode $\mathrm{H}_2\mathrm{O}_2$

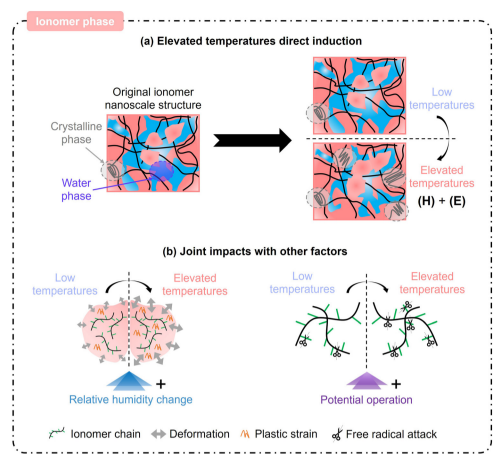


Fig. 11 Schemic diagram of ionomer degradation at elevated temperatures: (a) elevated temperature direct induction and (b) joint impacts with other factors

formation rates cannot be directly correlated with the ionomer degradation rates because there are several intermediate reactions between H2O2 formation and actual ionomer chemical degradation. However, it can be inferred that more significant ionomer chemical degradation may occur due to more severe free radical attack at higher concentrations of H2O2 generated at elevated temperatures, 166 as depicted in Fig. 11(b). The temperaturedependence of free-radical attack on ionomers remains unclear.

However, direct experimental observations or modelling calculations on the ionomer mechanical fatigue and ionomer chemical degradation at elevated temperatures have not been conducted. Additionally, thin, photo-sensitive, and low-density organic ionomer films are challenging to characterise and image, and ionomers in real CLs may exhibit non-uniform distribution, discontinuities, and variable thicknesses. The molecular chain configurations within an ionomer, spanning both in-plane and through-plane orientations, are strongly dependent on its thickness. 90,136 Domain changes in the ionomer phase, humidity fluctuations, and free radical attacks on ionomer molecular chains may occur simultaneously during PEMFC operation. The synergistic mechanisms underlying these processes and their effects on PEMFC performance are not yet fully characterised, and these uncertainties hinder a comprehensive understanding of ionomer degradation within the real CL at elevated temperatures.

4.1.4. Structural collapse. The CL functions as a critical component that facilitates electrochemical reactions and relies on the support from the PEM or GDL due to its intrinsically low structural strength. 170 Structural degradation of the CL can profoundly affect the electrochemical reactions and mass transport processes occurring within it. The CL exhibits distinct structural characteristics spanning from the microscale to the nanoscale, as shown in Fig. 13(a). Elevated temperatures complicate the structural evolution of the CL by introducing challenges to its stability and performance.

Existing studies have provided some insight into the structural evolution of the CL at the microscale, but have mainly focused on experimental phenomena. At the microscale, macroscopic features such as surface cracking and thickness decreases are observed. Han et al. 167 conducted two in situ fatigue experiments on a catalyst-coated membrane (CCM) at different operating temperatures and revealed that the CL surface exposed to 95 °C exhibited wider and deeper cracks than those formed at 85 °C, as shown in Fig. 13(b). They emphasised the significant contribution of the glass transition temperature of CCM (\sim 90 °C) on the CL surface cracking. An increase in the deformation of the PEM during the hydrothermal cycling, especially when the temperature exceeded the glass transition temperature, led to more severe cracking within

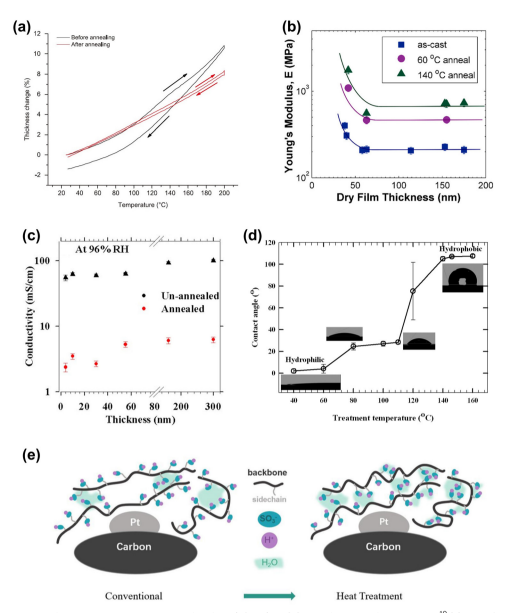


Fig. 12 (a) Thickness change of unannealed and annealed Nafion films (50 nm) on SiO₂ as a function of temperature.¹⁹ (b) Young's modulus at 30% RH: as cast, 60 °C annealing, and 140 °C annealing, 24 (c) Proton conductivity of unannealed and annealed Nafion thin films at 96% RH and 60 °C. 158 (d) Surface wettability of annealed ultrathin (10 nm) Nafion film.¹³⁶ (e) Mechanism of enhanced local transport after heat treatment.¹³⁷

the CL. 164,171 However, the structural stability of the CL was also significantly affected by elevated temperatures, particularly for ionomers. As the binder in the CL, an ionomer helps preserve the CL's surface integrity. 172 Therefore, further investigations into the evolution of ionomer adhesion properties on Pt/C catalyst agglomerates and their structural strength at elevated temperatures are warranted. More severe cracking can result in a range of detrimental effects, including an increased risk of PEM perforation, 173 water flooding within the CL, 173 high free-radical concentrations, 173 enhanced brittleness of the CL structure, 172 and the loss of certain agglomerates. 174 Investigating the deteriorated cracking mechanisms within the CL at elevated temperatures can provide valuable insights for improving the performance and extending the service life of PEMFCs.

Studies have shown that the thickness reduction of the CCL increases at elevated temperatures, 142,168,175 as displayed in Fig. 13(c) and (d). Kneer et al. 168 and Trogisch et al. 142 suggested that the observed reduction in CL thickness was negligible and primarily attributed to the compression effect of the MEA, but elevated temperatures may still substantially accelerate carbon corrosion rates, as discussed above. As shown in Fig. 13(b), Zhao et al. 131 reported that compared to 60 °C, the thickness of the CCL decreased by 9.3% after durability tests at 90 °C. The RH level also significantly affects the reduction of CL thickness. In region 9, because there was no water flooding during operation at 90 °C, the thickness reduction of the CCL at 90 °C was less than operation at 60 °C.131 Therefore, as the structural backbone of the CL, progressive carbon corrosion exacerbates thickness

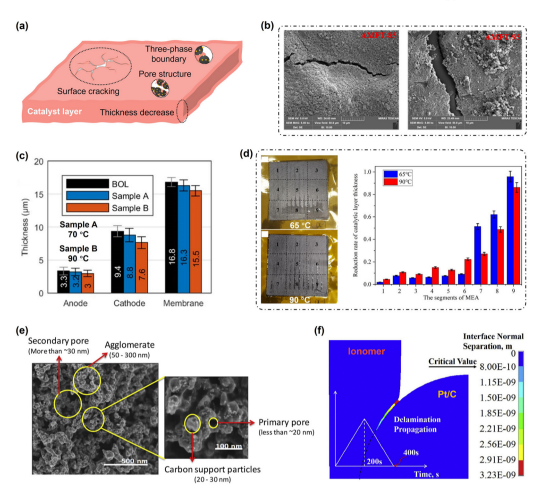


Fig. 13 Structure collapse features in the CL at elevated temperatures. (a) Diagram of collapse characteristics of CL structures at the micrometer and nanometer scales. (b) Cracking on the CL surface. 167 (c) Decrease in the thickness. 168 (d) Photographs of the MEA cathode and reduction rate of CL thickness in each area after 100 h operation at 1600 mA cm⁻² at 65 °C and 90 °C. ¹³¹ (e) Pore structure evolution. ¹⁶⁹ (f) TPB structure evolution. ¹⁶²

reduction, leading to increased catalyst loss and significant performance deterioration. Kneer et al. 168 also indicated the increased tortuosity of an aged MEA at elevated temperatures compared with lower temperatures. However, the effects and underlying mechanisms of elevated temperature-induced carbon corrosion on CL thickness and structure in the through-plane direction are unexplored. Additionally, the evolution of ionomer mechanical properties at elevated temperatures and their impact on CL thickness and structure in the through-plane direction warrant further investigation.

Nanoscale features include pore-based mass transport networks and three-phase boundary structures critical for electrochemical reactions. Among the microstructural properties of the CL, the pore size distribution and porosity are particularly critical because they govern the diffusion of reactant gases to active reaction sites and the efficient removal of generated water from the CL. Typically, two types of pores exist within the CL: (i) primary pores, with sizes smaller than approximately 20 nm, and (ii) secondary pores, ranging from 30 nm to 150 nm. Primary pores are located between carbon particles within agglomerates, whereas secondary pores represent the void spaces between agglomerates and are generally larger than

primary pores, as depicted in Fig. 13(e). Lu et al. 133 tested the durability of a PEMFC stack near the boiling point of water for 100 h and observed an increase in the pore size of the MEA in the range of 10-100 nm. As posited by Kneer et al., 168 the aged samples exhibited a shift in their pore size distribution towards larger pores, while the overall porosity of the cathode remained unchanged. The observed shift was interpreted as an onset of carbon corrosion that resulted in the collapse of the smallest pores and an increase in the number of larger pores. Yue et al. 132 demonstrated that the primary pores shrank while the secondary pores expanded after 1000 h of on-road heavyduty transportation with a coolant temperature of 92 °C, which was attributed to ionomer migration. Elevated temperatures accelerated carbon corrosion and also impacted the properties of ionomers, altering the pore structure within the CL.

Although Kneer et al. 168 suggested no obvious differences in the CL pore structure in the cells cycled at 90 °C versus those cycled at 70 °C, their conclusions were based on 3D reconstructions of the CL obtained using focused ion beam scanning electron microscopy, where the identified pores were generally considered secondary pores. 132 The evolution of primary pores, however, remains poorly understood, and additional research is needed to clarify the

mechanistic relationships between elevated temperatures, compositional changes, and the pore structure evolution.

The TPB structure consists of a Pt catalyst, ionomer/water films, and pore spaces and serves as the critical site for electrochemical reactions, as depicted in Fig. 13(f). Optimizing the TPB structure significantly enhances the power density and catalyst utilization, 176 and distortions in the TPB structure induced during operation can severely compromise the cell performance and durability. Current research on the evolution of the TPB structure primarily emphasises interfacial delamination between the ionomer and Pt/C agglomerates. 162,163,177-179 Cohesive zone models have demonstrated that the cyclic expansion and contraction of ionomers during hygrothermal cycling cause the partial detachment of ionomers from Pt/C agglomerates, leading to cracks in the CL and a reduction in the ECSA. 162,163,177-180 Changes in the water content of ionomers at lower temperatures are considered the primary driver of interfacial delamination. 162 Notably, Chang et al. 162 reported that lowtemperature cycling alone does not significantly induce interfacial delamination. In contrast, the extent of interfacial delamination is governed by the interplay between an ionomer's expansion-contraction and its adhesive properties, both of which are impacted by elevated temperatures. Moreover, Yue et al. 132 showed that the contact area between Pt catalysts and ionomers decreased by 37.3% due to ionomer migration at elevated temperatures, indicating significant changes in the TPB structure. However, the detailed distortion mechanisms of the TPB structure at elevated temperatures are unexplored. Notably, the role of humidity in the evolution of pore structure and the TPB structure at elevated temperatures significantly affects the degree of ionomer deformation.

4.2. Proton exchange membrane durability

The PEM plays a crucial role in transporting protons from the anode to the cathode, insulating negative electrons, and separating the reactants, hydrogen, and oxygen. Therefore, the membrane must be an efficient conductor of protons and an effective insulator of electrons and must exhibit both mechanical and chemical robustness. The most commonly employed membranes in PEM fuel cells are PFSA membranes with perfluoroether long side chains with sulfonic acid end groups. ¹³⁸ Recently, alternative PFSA membranes with short side chains have garnered significant attention for investigation. ¹⁸¹ Among these, the Nafion membrane is the most widely used due to its long-term stability in both oxidative and reductive environments, relatively low fabrication cost, and ease of processing. Nevertheless, the failure modes of PEMs, including perforation, irreversible reduction in proton conductivity, cracking, *etc.*, at elevated temperatures, emphasize the critical need to investigate and enhance their stability in such environments. Table 8 summarizes the stability of PEMs at different temperatures.

4.2.1. Elevated temperature direct induction. PEMs can undergo various changes in environments with elevated temperatures, and the extent of these changes strongly depends on both the temperature and duration of exposure, as shown in Fig. 14. This is considered to be a direct consequence of elevated temperature induction on PEMs.

In the absence of exposure duration to elevated temperatures, as illustrated in Fig. 14(a), TGA-MS can be used to determine the thermal decomposition process of PFSA ionomers at different temperatures. Typically, this occurs in three stages: $^{4,127,188,194-196}$ (i) loss of membranous water (100–200 \pm 20 °C), which culminates in the rupture of $\rm H_3O^+$ and the protonation of $\rm SO_3^-$ to $\rm SO_3H$, as well as the partial decomposition of $\rm SO_3H \rightarrow \rm SO_2 + OH^\bullet$ near 200 °C; (ii) cleavage of the C–S bond (280 \pm 30 °C to 400 \pm 20 °C) leading to sulfonate-group degradation; and (iii) the final decomposition of the perfluorinated matrix (400 \pm 20 °C to 600 \pm 40 °C). This demonstrates that in the elevated temperature range required for MEAs, only the loss of water occurs within the PEM, inducing phase transitions in clusters/ion channels without damaging its molecular structure. Rodgers $et~al.^{197}$ confirmed

Table 8 Stability of PEMs at different temperatures

Serial number	Durability cases and corresponding conditions (anode/cathode)	Items	Values	Ref.
1	NEDC near-water boiling temperature: operating temperature: 95 °C, inlet pressure (kPa): 280/260, duration: 100 h;	PEM thickness decrease	1.9-8.4%	133
2	A combined mechanical/chemical accelerated stress test: operating temperature: 90 °C, gas type: H ₂ /air, back pressure: no duration: M – 8 for 1512 h, M – 12 for 1488 h	Hydrogen crossover increase	M - 8: 348.4% @ 0 A cm ⁻²	182
	·		M - 12: 398.7% @ 0 A cm ⁻²	
3	Modelling of PEM chemical degradation	PEM mass loss rate	Ratio of 95 $^{\circ}$ C to 80 $^{\circ}$ C: 4.83	183
4	Modelling of PEM chemical degradation	Hydrogen crossover flux	Ratio of 90 $^{\circ}$ C to 70 $^{\circ}$ C: 1.53	184
5	Thermal annealing: sample: short side chain (SSC) membrane, gas atmosphere: vacuum, duration: 1 h, temperature: 140 °C;	PEM water uptake decrease	8.1–11.1%	185
	, ₁	PEM conductivity decrease	3.9-4.7%	
		Glass transition temperature	Increase from 134 °C to 143 °C (a) 3 M membrane	
			increase from 131 °C to 139 °C @ aquivion membrane	
6	Thermal annealing: sample: aquivion 870 membrane, duration: 72 h, temperature: 140 °C	Glass transition temperature	Increase from 124 °C to 131 °C @ aquivion 870 membrane	186
7	Modelling of PEM chemical degradation	Fluoride emission rate	Ratio of 90 °C to 80 °C: 4.9	187

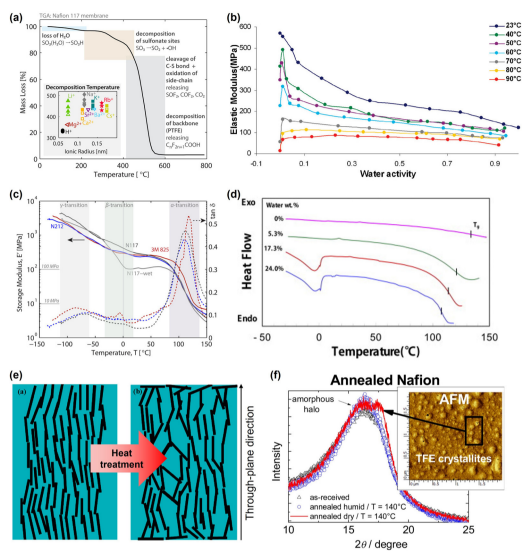


Fig. 14 (a) TGA curve of the Nafion 117 membrane. 188 (b) Instantaneous elastic response of Nafion N1110 as a function of water activity from 23 to 90 °C. 189 (c) Typical dynamic mechanical analysis (DMA) profiles of Nafion and 3 M PFSA membranes. 190 (d) Differential scanning calorimetry (DSC) data of the Nafion 117 membrane as a function of water content. 191 (e) Evolution of proton channels in the direction perpendicular to the membrane surface during the heat treatment process. 192 (f) Characterization results for N105 samples annealed at 140 °C, dry conditions and 140 °C, $RH = 100\%.^{193}$

this conclusion and suggested that the Nafion ionomer was thermally stable up to ~ 150 °C due to the presence of strong C-F bonds and a shielding effect of the electronegative fluorine atoms on the PTFE-like backbone.

Nevertheless, Majsztrik et al. 189 investigated the elastic modulus of Nafion N1110 as a function of water activity from 23–90 $^{\circ}\mathrm{C}$ and suggested that for any given level of hydration (water activity), the elastic modulus always decreased with temperature, as indicated in Fig. 14(b). This was because the internal structure of the PEM evolved, resulting in a decrease in its strength, which became apparent in the (M) regime. In practice, the internal molecular chain structure and domain structure of a PEM typically undergo a thermal transition process as the temperature increases, 198 which is captured by dynamic mechanical analysis (DMA).90 A typical DMA curve of the PEM

is demonstrated in Fig. 14(c), in which the characteristic thermal transition temperatures are noted.90 The PFSA membrane exhibits three thermal transitions corresponding to different relaxation behaviours: (i) an α relaxation at approximately T_{α} = 90–120 °C for acid-forming membranes; (ii) a broader β relaxation between $T_{\beta} = -40$ °C and 20 °C; and (iii) a γ relaxation approximately $T_{\gamma} = -120$ °C to -90 °C. Regarding the three relaxation mechanisms, Kusoglu et al.⁹⁰ reported that (i) T_{α} was a dynamic ionic-cluster transition associated with the onset of long-range mobility of the chains due to destabilization of the electrostatic network; (ii) T_{β} was the glass-transition temperature of the PFSA matrix attributed to the onset of segmental main-chain motion, facilitated by sidechains; (iii) the low-temperature transition T_{γ} was related to the local movement of the PTFE backbone. 90 When the

temperature exceeded T_{α} , the weakening of the electrostatic interactions within the ionic aggregates triggered the onset of long-range mobility of both the main and side chains, resulting in a significant increase in the chain mobility.¹⁹⁹

Due to the glass transition of the membrane, its mechanical properties were greatly altered. The mechanical behavior of a material, i.e., its ability to withstand deformation without rupturing, can be evaluated using its stress-strain curves. Numerous studies have demonstrated that the stress-strain curves of a membrane undergo a significant decline at elevated temperatures due to a reduction in both the elastic modulus and the yield stress, as well as a reduction in ultimate tensile strength and an increase in the strain at break. Marrony et al. 200 indicated that the ultimate tensile strength decreased by 54.1% when the temperature increased from 80 $^{\circ}\text{C}$ to 120 $^{\circ}\text{C}$ due to a decrease in intermolecular forces and the presence of amorphous domains. 90,164,201 Therefore, PEMs tend to soften in elevated temperature environments, especially close to or exceeding their glass transition temperatures. Notably, the membrane water content significantly influences the magnitude of the glass transition temperature. As depicted in Fig. 14(d), the glass transition temperature decreased from 132 $^{\circ}\text{C}$ to 108 $^{\circ}\text{C}$ when the membrane water content increased from 0% to 24.0%. 191 Therefore, the fragility of PEMs containing water needs to be given special consideration.

Considering the effect of the exposure duration on elevated temperature environments, the influence of elevated temperature environments on the PEM is typically referred to as thermal treatment or thermal annealing. It primarily focuses on its influence on the internal domain structure and the evolution of its operational characteristics (proton conductivity, mechanical properties, *etc.*), as illustrated in Fig. 14(e) and (f).

Studies have shown that thermal treatment of PEMs often decreases the proton conductivity and enhances the mechanical strength, particularly when the thermal treatment temperature falls within the (H) and (E) regimes. $^{185,186,191-193,202-205}$ This is attributed to increased crystallinity in the hydrophobic domains and changes in the ionic domain structure. An increase in crystallinity enhances the glass transition temperature, tensile strength, etc. Shin et al. 185 reported an increase in the glass transition temperature from 134 $^{\circ}$ C to 143 $^{\circ}$ C for a 3 M membrane after vacuum thermal treatment for 1 h at 140 °C. Factors leading to an increase in crystallinity are multifaceted, including increased chain mobility, increased long-range order, and the melting and re-crystallization of imperfect PTFE-like crystallites at elevated temperatures. 186,191,202 However, Jeon et al.206 suggested that the crystallization of Nafion 212 and Aquivion E87-05S membranes decreased after thermal treatment at 90-140 $^{\circ}$ C. As demonstrated in the seminal work conducted by Hensley et al., 203 annealing had a marked effect on the crystallinity of films, with a decline observed in the crystallinity of thick films concomitant with an increase in the crystallinity of thin films, as determined by wide-angle X-ray measurements. The heat treatment duration in Jeon's study was much longer than that in the other four studies, which may have re-damaged the already-formed crystalline regions.

Regardless, the mechanical properties of the membrane exhibited fluctuations as the exposure time increased at elevated temperatures, which pose a significant challenge to the reliable sealing and stable operation of PEMFCs.

As illustrated in Fig. 14(e), the through-plane conductivity of a PEM significantly decreased after treatment at 120 °C for 30 h. This reduction was caused by the reorientation of proton transport pathways near the membrane surface from perpendicular to parallel to the surface as the annealing temperature approached the glass transition temperature of the membranes, thereby impeding proton conduction. 192 Shin et al. 185 also suggested a proton conductivity loss in the range of 3.9-4.7% after vacuum annealing for 1 h at 140 °C. Notably, Matos et al. 193 reported that the properties of a PEM were significantly less affected when annealing was carried out at high RH, as illustrated in Fig. 14(f). Despite that, the internal evolution of PEMs at elevated temperatures remains unclear, and low-RH conditions pose a significant challenge to the stable operation of PEMs at elevated temperatures. Furthermore, the extent of phase separation in the hydrophobic matrix and the hydrophilic region increases due to the thermal treatment based on the small-angle X-ray scattering measurements. 185 This can increase the risk and degree of hydrogen or oxygen crossover through the PEM, thereby reducing the OCV of the PEMFC and inducing chemical degradation.

4.2.2. Acceleration assistance for other degradation processes. Even though PEMs are highly stable, they also suffer from serious degradation at low temperatures such as mechanical failure and chemical degradation, which become worse at elevated temperatures. PEMs are readily available and easy to characterise, allowing accelerated degradation experimental phenomena to be observed at elevated temperatures. In contrast, research on the degradation effects in ionomer films is limited.

Elevated temperatures can accelerate the creep deformation of PEMs due to increased molecular mobility, especially when the temperature is close to or exceeds the glass transition temperature, 138,208 that is in the (H) and (E) regimes. Excessive creep deformation forms wrinkles and induces membrane thinning. Lu et al. 133 observed a 1.9-8.4% PEM thickness decrease after a 95 °C durability test. In addition to creep behaviour, elevated temperature can accelerate membrane fatigue, including but not limited to pinholes and cracks. Alavijeh et al. 171 conducted two accelerated mechanical stress tests on the CCMs at cell temperatures of 80 $^{\circ}\text{C}$ and 95 $^{\circ}\text{C}$ and reported that larger and more frequent cracks were observed within the sample at 95 °C. This was attributed to the increased fatigue amplitude and reduced mechanical strength of the membrane at elevated temperatures, suggesting that the MEAs at 95 °C underwent more severe mechanical failure, as shown in Fig. 15(a). Han et al. 167 conducted in situ hygrothermal cycling durability tests on a PEM with a reinforced layer of e-PTFE above its glass transition temperature and suggested that the crosslinking degree between molecular chains inside the membrane was weakened after exceeding the glass transition temperature. This resulted in a significant decrease in the mechanical

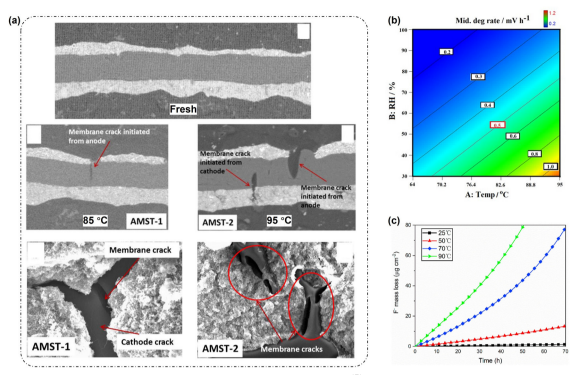


Fig. 15 (a) SEM images of fresh and degraded CCMs from the AMST-1 and AMST-2 tests.¹⁷¹ (b) Contour plot of effects of temperature and humidity on the Mid. membrane degradation rate.²⁰⁷ (c) Evolution of cumulative fluoride loss at different operating temperatures.¹⁸⁴

properties of the PEM after durability tests. More severe silver lines and earlier fractures appeared during tensile tests.

Some studies have conducted modelling of ex situ and in situ fatigue behaviour. 209,210 The application of constitutive models and failure criteria to describe materials remains a fundamental challenge in understanding fatigue mechanisms and accurately predicting material lifetimes at elevated temperatures. To achieve high-fidelity computational simulations of the fatigue process, it is essential to incorporate material models that capture the nonlinear viscoelastic and/or viscoplastic effects, particularly those that consider the influence of elevated temperatures. Particular attention should be given to the (H) and (E)regimes, as membrane failure during operation will be significantly accelerated under such conditions. At low temperatures, RH fluctuations and pressure differences between the anode and cathode sides are the main driving forces for membrane fatigue failure. At elevated temperatures, the contribution of RH and pressure difference variation characteristics to membrane mechanical failure requires further investigation to understand membrane degradation during dynamic operation of PEMFCs at elevated temperatures. Regardless, more severe membrane failures significantly increase the crossover leak rate, risking the stable operation of PEMFCs. 208

As illustrated in Fig. 15(b), Zhao et al.207 investigated the impact of operating temperature on the durability of membranes using the OCV accelerated stress tests (AST). Their findings revealed a direct proportionality between the membrane degradation rate and the operating temperature. The study identified an optimal operational region, which was determined without changing the membrane materials, suggesting that operation at >90 °C was not feasible. A decrease in RH levels further accelerated the chemical degradation of the PEM. Failure of the Nafion XL membrane was observed only after ~ 250 h of the steady-state OCV AST at 30% RH and 90 °C. 211 Therefore, elevated temperature dramatically affected the membrane chemical degradation, especially in a low-RH environment. Modelling calculations also confirmed that elevated temperatures accelerate the chemical degradation of the membrane, as evidenced by a significant increase in PEM mass loss and the fluoride emission rate. Chen et al. 183 reported that increasing the operating temperature from 80 °C to 90 °C resulted in an approximately 4.83fold increase in PEM mass loss. Frensch et al. observed a 4.9-fold increase in the fluoride emission rate upon raising the operating temperature from 80 °C to 90 °C. Similarly, as shown in Fig. 15(c), Zheng et al. 184 found a dramatic increase in the fluoride ion release rate and hydrogen crossover flux (1.53-fold) at 90 °C compared with 25 °C, indicating the severe chemical degradation degree at elevated temperatures. The PEM chemical degradation at low temperatures is induced by free-radical attack. As discussed in Section 4.1.3, the rates of H₂O₂ formation significantly increase at elevated temperatures, offering a significant number of sources for free radical generation. 165 Chen et al. 183 also supported the above conclusion and indicated that the chemical degradation of the PEM was accelerated due to an accelerated chemical reaction in the cell at elevated temperatures, exhibiting an exponential relationship with temperature (60 $^{\circ}$ C $\leq T \leq$ 95 °C). Furthermore, trace metal ions resulting from the corrosion of bipolar or end plates, as well as the leaching of less-noble

metals from an alloy catalyst, may also accumulate within the membrane. These ions can catalyse radical attacks previously mentioned and accelerate chemical degradation of the membrane, but there are several intermediate reactions between H2O2 formation and actual ionomer chemical degradation. The effect of elevated temperatures on these intermediate reactions, including free radical formation, subsequent attack of free radicals on membrane polymer chains, and leaching of metal ions, is unclear. Notably, the mechanism between mechanical failure and chemical degradation of membranes has been investigated, 212-214 but the role of elevated temperatures is poorly understood. The interplay between elevated temperatures and mechanical and chemical degradation needs further investigation to provide a theoretical basis for extending the lifetime of PEMFCs.

4.3. Gas diffusion layer durability

The primary function of the GDL within a PEMFC is twofold. First, it must facilitate the transport of reactant gases from the flow channels to the CL and also facilitate the drainage of water produced at the CL.217,218 The GDL must also serve as an efficient electron conductor to collect the current. Various types of GDLs are under development, but they are typically comprised of porous carbon fibres and carbon particles, 127 and a typical GDL consists of a GDB and a MPL. The primary function of the GDB is the distribution of reactant gases, in conjunction with the current collection. The MPL, which is composed of carbon powder and hydrophobic materials such as PTFE, is responsible for managing water flow. Consequently, it can be concluded that mechanical strength, conductivity, and hydrophobicity are fundamental properties of the GDL.

Several mechanisms can cause degradation of the GDL, including chemical degradation, erosion resulting from gas flow, dissolution by water, freezing/thawing effects, and mechanical compression. There is limited research into the impact of elevated temperature on GDL degradation. 127,133,215,216,219 Table 9 shows the stability of the GDL at elevated temperatures. TGA has shown that the decomposition or volatilization of the GDL occurs only when the temperature reaches 500-650 °C due to thermal degradation of fluorinated polymers. This temperature is much higher than the elevated temperature range required for PEMFCs, indicating that the GDL materials exhibit strong resistance to thermal decomposition. Nevertheless, elevated temperature environments still have a considerable detrimental effect on the operational performance of the GDL, primarily due to thermal softening^{133,215} and thermal oxidation.^{127,133,216} The main degradation characteristics include carbon fibre fracture, 133,215

reduction in the porosity, 212 filler loss, 133 thickness reduction, 133 PTFE loss, 133,216 and more cracking in the MPL. 133

Thermal softening describes the phenomenon in which there is a decrease in the storage elastic modulus of binder resins (phenolic resin, epoxy resin, cellulose acetate, etc.) at elevated temperatures, 220-224 leading to a decrease in the compressive stiffness of the GDB layer. This also demonstrates the marked temperature dependence on the mechanical stiffness of PTFE at low temperatures. Indeed, the mechanical stiffness of pure PTFE sharply decreases as the temperature increases from −150 °C to 150 °C.²²⁵ This temperature-dependent behaviour of these adhesive resins and PTFE is the main cause of the reduction in compressive stiffness of the GDL at elevated temperatures. This softening effect results in more extensive and severe carbon fiber breakage within the GDB, 215 as depicted in Fig. 16(a), accompanied by a reduction in the porosity. ²¹⁵ The localised fracture of carbon fibres may lead to the erosion of fillers within the GDB due to water and gas flushing, ¹³³ leading to the localised appearance of large pores, as illustrated in Fig. 16(b). The reduction in the GDL thickness (anode side: 1.5-10.7%, and cathode side: 7.7-23.2%) after a durability test at 95 °C may have been induced by the thermal softening, ¹³³ as shown in Fig. 16(c). However, the mechanism by which thermal softening affects filler loss and thickness reduction remains unclear. Regardless, the compressive resistance and mass transfer capability of the GDL are compromised at elevated temperatures, amplifying the differences in electrochemical reactions beneath the flow channels and ribs and exacerbating the inplane non-uniformity of the reaction.

Thermal oxidation refers to the degradation of GDL materials resulting from their reaction with oxygen at elevated temperatures. Qu et al. 216 reported that under an air atmosphere at 120 °C, PTFE undergoes thermal oxidation, resulting in a looser surface structure. This loosening was attributed to the degradation of molecular chains during thermal-oxidative aging, with the escape of small molecules, which weakened interactions between chains. The thermal oxidative aging mechanism is shown in Fig. 16(e). As thermal oxidation progresses, PTFE is gradually oxidised from its end groups, which releases volatile small molecules such as CF₂O and decreases the PTFE content. PTFE loss was also observed after a 100 h, 95 °C New European Driving Cycle (NEDC) durability test, after which the loss of PTFE in the cathode was more pronounced than in the anode, 133 as shown in Fig. 16(d). However, the contribution of thermal oxidation to PTFE loss at elevated temperatures is unclear, as PTFE may also detach due to erosion caused by the

Table 9 Stability of the gas diffusion layer at elevated temperatures

Durability cases and the corresponding conditions	Items	Values	Ref.
NEDC near-water boiling temperature: operating temperature: 95 °C, inlet pressure (kPa)	GDL thickness decreases	Anode: 1.5–10.7% Cathode: 7.7–23.2%;	133
(anode/cathode = 280/260), duration: 100 h;	GDB graphitization degree decreases	Anode: 286.3% Cathode: 493.2%;	
	MPL graphitization degree decreases	Anode: 2.0% Cathode: 9.4%;	

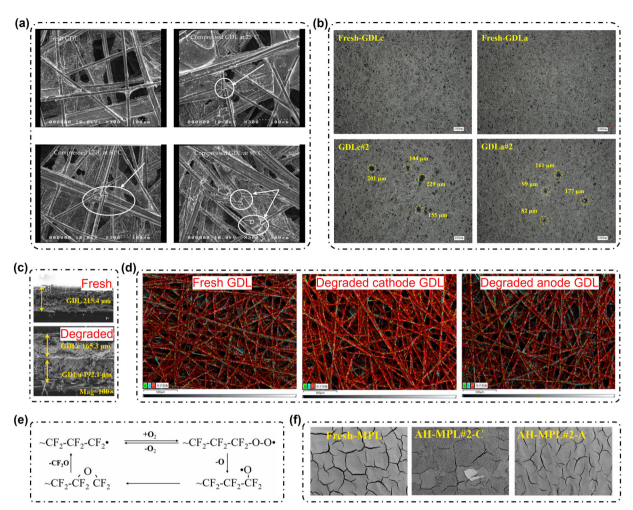


Fig. 16 Durability issues for the GDL at elevated temperatures. (a) SEM images of the uncompressed GDL and compressed GDL at 25 °C, 60 °C, and 90 °C. 215 (b) Changes in pinhole distribution on the GDL surface, 133 (c) thickness features of the GDL 133 and (d) changes in elemental content of GDL surfaces before and after elevated temperature testing. 133 (e) Molecular structure change during the thermal oxidation process of PTFE. 216 (f) Surface morphologies of MPL before and after elevated temperature testing. 133

reactive flow. 226 Additionally, carbon particles and the surfaces of carbon fibres can also undergo oxidation. 127,133 Lu et al. 133 reported significant GDB graphitization degree decreases of 286.3% at the anode side and 493.2% at the cathode side after a durability test at 95 °C. Similarly, the MPL graphitization degree also decreased, but not significantly, as presented in Table 9. Nevertheless, there are a few studies on the oxidation of these materials at elevated temperatures.

Overall, the reduction in the PTFE content or the oxidation of the GDL diminishes the hydrophobicity of the GDL, 133 obstructing the liquid water drainage and increasing the risk of flooding. Lu et al. 133 reported increased cracking in the MPL surface after a 100-hour, 95 °C NEDC durability test, as shown in Fig. 16(f). This was attributed to the water and gas transport, but the underlying mechanism was unclear. The more pronounced cracking, coupled with the hydrophilization of the surface, exacerbates the risk of water flooding. The degradation mechanisms of the mass transport properties of GDL at elevated temperatures warrant further investigation. Given the distinct compositions

of the GDB and MPL, the degradation of the two components at elevated temperatures should not be generalised.

4.4. Impacts on performance loss

The degradation of MEA components induced by elevated temperatures results in substantial PEMFC performance loss. Fig. 17 presents the performance degradation characteristics of PEMFCs after various durability tests at elevated temperatures, including on-road heavy-duty driving conditions (Fig. 17(a)), NEDC cycling (Fig. 17(b)), in situ hygrothermal fatigue testing (Fig. 17(c)), constant operation at a high current density (Fig. 17(d)), voltage cycling (Fig. 17(e)), and temperature cycling (Fig. 17(f)). PEMFCs experience more severe performance degradation at elevated temperatures due to the harsher deterioration of the MEA. For instance, Ruiu et al. 22 demonstrated that short-term temperature cycling between 90 $^{\circ}\mathrm{C}$ and 120 $^{\circ}\mathrm{C}$ caused more pronounced degradation in MEA components compared with long-term stable operation under automotiverelevant conditions at 80 °C. This resulted in a fivefold increase

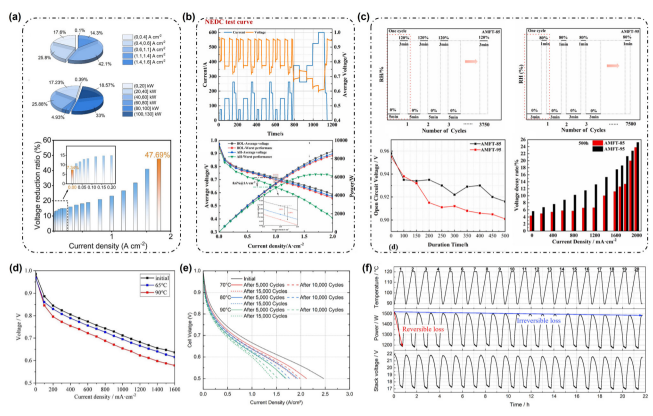


Fig. 17 (a) Time percentages of operating current and power during the 1000 h on-road heavy-duty transportation with a coolant temperature of 92 °C and voltage reduction ratio at each current density. 132 (b) NEDC durability protocol and polarization curves of the stack before and after the durability testing with a coolant temperature of 95 °C for 100 h.133 (c) In situ accelerated fatigue conditions of RH cycling: accelerated mechanical fatigue tests at the operating temperature of 85 °C and 95 °C, the OCV trends, and voltage decay rate after 500 h of RH cycling. 167 (d) Polarization curves before and after 100 h of continuous operation at different temperatures. ¹³¹ (e) Polarization curves after voltage cycling at different operating temperatures. ¹⁴⁷ (f) Stack temperature, power and voltage observed during 20 thermal cycles at 70 A.²²

in the rate of irreversible performance loss of the PEMFC. Tables 10 and 11 summarise the performance degradation rates of PEMFCs at different temperatures and the dominant degradation mechanisms of the major MEA components at elevated temperatures, respectively. Other operating conditions also have varying impacts on the service life of the MEA at elevated temperatures. For instance, as shown in Case 10 of Table 10, when the operating temperature is set to 95 °C, the lifetime under 30% RH is only 82 h, which is significantly shorter than 168 h observed at 60% RH. 207 The degradation of different MEA components has a profound impact on the performance loss. It is well-established that the output performance loss of PEMFCs is governed by different overpotential increments, including the leakage loss overpotential η_{loss} , the charge-transfer overpotential (η_{act}), the ohmic overpotential $(\eta_{\rm ohm})$, and the concentration overpotential $(\eta_{\rm con})^{230}$ Of these, $\eta_{\rm loss}$ is mainly affected by gas permeation and electronic shortcircuiting. As illustrated in Fig. 18(a), elevated temperature can increase $\eta_{\rm loss}$ due to the PEM degradation due to increased gas crossover flow caused by enhanced phase separation, more severe cracking, pinholes, etc. Furthermore, increased Pt growth and dissolution, more severe carbon support chemical and electrochemical corrosion, and more severe distortion of

the ionomer and the TPB structure at elevated temperatures are the main causes that further decrease the ECSA, leading to a higher η_{act} , as shown in Fig. 18(b). Additionally, the increased crystallinity of the hydrophobic region and decrease in the hydrophilic region in the ionomer phase within the CL and PEM at elevated temperatures further inhibit proton transport, stimulating an increased $\eta_{\rm ohm}$, as shown in Fig. 18(c). Moreover, the increased gas transport pathways induced by the increased Pt catalyst dissolution, more severe carbon support corrosion, the distortion of ionomer and pore structure in the CL, and reduced porosity of the GDL lead to an increased η_{con} , as shown in Fig. 18(d).

Given the significant impact of elevated temperatures on the degradation of MEA components, it is crucial to recognise and address the associated performance losses in PEMFCs. Establishing a quantifiable relationship between component degradation and performance decline is necessary to optimise the PEMFC durability at elevated temperatures. Because these degradation processes are exacerbated at elevated temperatures, maintaining optimal operating conditions and developing materials with enhanced thermal stability are essential for improving the long-term performance and reliability of a PEMFC.

Review

 Table 10
 Performance loss of PEMFCs at different temperatures

Serial number	Durability cases and corresponding conditions (anode/cathode)	Items	Values	Degradation causes	Re
l	On-road heavy-duty transportation: operating temperature of 92 °C, dura- tion: 1000 h;	Cell voltage loss	7.34% @ 0 A cm ⁻²	1. Pt catalyst growth	13
	don 1000 n,		47.69% @ 1.8 A cm ⁻²	Pt catalyst loss Carbon support corrosion Ionomer migration and chemical degradation	
	<i>In situ</i> hydrothermal cycling: operating temperature: 85 °C and 95 °C, inlet pressure (kPa): 80/60, gas type: H ₂ /N ₂ , dew point temperature: 90 °C, stoichiometric ratio: 1.5/2.5, duration: 500 h	Cell voltage loss	85 °C	1. Cracks within the CCM	16
			4.38% @ 0 A cm ⁻²	2. Delamination between the CL and PEM	
			23.74% (a) 2 A cm ⁻² 95 °C: 5.65% (a) 0 A cm ⁻²		
•	NEDC near-water boiling temperature: operating temperature: 95 °C, inlet pressure (kPa): 280/260, duration: 100 h	Cell voltage loss	25.17% ⓐ 2 A cm ⁻² MEA#1:	1. Polymer melting	13
	pressure (ki a). 200/200, duration. 100 ii		3.34% @ 0 A cm ⁻² 1.07% @ 1.0 A cm ⁻²	 Carbon support corrosion Aggregation and growth of Pt nanoparticles 	
			10.85% @ 2.0 A cm ⁻² MEA#2: 13.89% @ 0 A cm ⁻² 56.51% @ 0.4 A cm ⁻²	4. MEA structural breakdown	
			MEA#3: 11.26% @ 0 A cm ⁻² 49.13% @ 0.4 A cm ⁻² MEA#4:		
			0.27% @ 0 A cm ⁻² 0.19% @ 1.0 A cm ⁻² 19.36% @ 2.0 A cm ⁻² MEA#5: 3.12% @ 0 A cm ⁻² 8.61% @ 1.0 A cm ⁻²		
	A combined mechanical/chemical accelerated stress test: operating temperature: 90 °C, gas type: H ₂ /air, back pressure: no duration: M - 8 for 1512 h, M - 12 for 1488 h;	Cell voltage loss	58.33% @ 2.0 A cm ⁻² M - 8:	Cracks or pinholes within the PEM	1
			12.48% (a) 0 A cm ⁻² M - 12: 8.39% (a) 0 A cm ⁻²		
	Load-cycle: operating temperature: 120 °C, gas type: H_2/O_2 , stoichiometric ratio: 1/1.5, RH: 40%, range of potential: 0.6–1 V, PEM type: Nafion [®] , duration: 7 h;	Cell voltage loss	8.39% @ 0 A cm 16.57 μV h ⁻¹ @ 0.5 A cm ⁻²	1. Hydrogen crossover increase through the PEM	22
	Load-cycle: operating temperature: $120 ^{\circ}\text{C}$, gas type: H_2/O_2 , stoichiometric ratio: $1/1.5$, RH: 40% , range of potential: $0.6{\text{-}}1 ^{\text{TM}}$, PEM type: Aquivion M, duration: $50 ^{\text{h}}$	Cell voltage loss	$2.22~\mu V~h^{-1}~ @~0.5~A~cm^{-2}$	2. Pt catalyst growth	
	Voltage cycle: sample A: square wave (2 s) , 0.4 – 0.95 V , 100% RH, $70 ^{\circ}\text{C}$, 460 h , sample B: square wave (2 s) , 0.6 – 0.95 V ,	Cell voltage loss	Sample A:	1. Pt catalyst growth	1
	100% RH, 90 °C, 400 h;		0.10% @ $0~{\rm A~cm^{-2}}$	2. Redistribution of Pt in the MEA	
			1.84% @ 1.5 A cm ⁻² 6.11% @ 2.5 A cm ⁻² Sample B: 2.79% @ 0 A cm ⁻² 13.00% @ 1.5 A cm ⁻² 30.00% @ 2.5 A cm ⁻²	3. Carbon support corrosion	

Table 10 (continued)

Serial number	Durability cases and corresponding conditions (anode/cathode)	Items	Values	Degradation causes	Ref
8	Hydrothermal cycle under OCV: RH: 100%, operating temperature: 50 °C, 70 °C, and 90 °C duration: 100, 200, 500, and 1000 h	Cell voltage loss	100 h, 0.6 A cm ⁻²	1. Destroyed Nafion structure	228
			8.6% @ 50 °C 4.7% @ 70 °C 23.2% @ 90 °C; 200 h, 0.6 A cm ⁻² : 14.5% @ 50 °C 7.7% @ 70 °C, 29.9% @ 90 °C; 500 h, 0.6 A cm ⁻² : 23.3% @ 50 °C 14.7% @ 70 °C 38.8% @ 90 °C; 1000 h, 0.6 A cm ⁻² : 32.2% @ 50 °C 21.9% @ 70 °C	Carbon support Catalyst aggregation	
9	Aging testing: operating temperature: 70 °C, 80 °C, 90 °C, 100 °C, and 120 °C; PEM types: sulfonated polyether therketone (SPEEK), Nafion [®] 212, Aquivion™ E790-03S, Aquivion™ E790-03S ⁺ (Gen 2), Aquivion™ E790-03S TT	Lifetime	48.1% @ 90 °C; SPEEK	1. PEM thinning	200
			$>\!2655$ h @ 70 $^{\circ}\mathrm{C}$	2. Delamination between the	
			<1558 h @ 80 °C Nafion® 212: > 2505 h @ 80 °C 685 h @ 90 °C; Aquivion™ E790-03S: > 269 h @ 90 °C < 107 h @ 100 °C Aquivion™ E790-03S⁺ (Gen 2): < 200 h @ 90 °C < 270 h @ 120 °C; Aquivion™ E790-03S TT: > 269 h @ 100 °C	CL and PEM 3. Pt catalyst dissolution 4. CL cracking	
10	OCV hold accelerated stress test: back pressure (bar): 3/2.9, RH: 30% and 60%, operating temperature: 64 $^{\circ}$ C, 80 $^{\circ}$ C, and 95 $^{\circ}$ C	Lifetime	> 240 h @ 120 °C 1100 h @ 64 °C and 60% RH	PEM chemical degradation	207
			339 h @ 80 °C and 60% RH 168 h @ 95 °C and 60% RH 215 h @ 80 °C and 30% RH 82 h @ 95 °C and 30% RH		
11	<i>In situ</i> accelerated aging procedure: operating temperature: 64 °C, 80 °C, and 95 °C; PEM types: Nafion [®] 212, Nafion [®] 115	Lifetime	Nafion [®] 212:	PEM degradation	229
			> 2820 h @ ≤80 °C <769 h @ 90 °C; Nafion [®] 115 > 2820 h @ ≤90 °C		
12	Thermal cycling test: dynamic temperature cycles between 90 and 120 °C, cell numbers: 30, active cell area: 142 cm², gas type: H ₂ /air, stoichiometric ratio: 1.5/2.5	Cell voltage loss	714 μV h ⁻¹ cell ⁻¹ (a) 70 A	1. PEM mechanical degradation	22
				 PEM chemical degradation CL structural damage CL ionomer degradation Delamination between the CL and PEM 	

Table 11 Major degradation mechanisms of MEA components at elevated temperatures

Degradation types	Degradation features		Degradation causes
Pt catalyst degradation	Growth	Direct induction	(i) Entire elevated temperature range, reduced H ₂ atmosphere, the removal of surface oxides on Pt catalysts, and subsequent crystallite reconstruction and grain boundary diffusion (ii) N ₂ atmosphere, 160 °C, Pt oxide reduction by the support
		Acceleration assistance	Indeterminate
	Dissolution	Acceleration assistance	 (i) Endothermic Pt dissolution reaction (ii) Impact on the Pt catalyst collective surface tension (i) Environment containing oxygen or hydrogen (H₂/O₂)
Carbon support corrosion	Chemical corrosion	Direct induction	(ii) Enhanced the reaction rate between the carbon matrix and oxygen or hydrogen species (iii) Catalysis of the Pt catalyst (iv) High RH accelerates
Ionomer degradation	Electrochemical corrosion	Acceleration assistance	(i) Intensified the carbon electrochemical corrosion sites (ii) High RH accelerates
	Ionomer nanoscale structure evolution	Direct induction	(i) Increased crystallinity of the hydrophobic region(ii) Shrank the hydrophilic region(iii) Increased extent of phase separation
	Mechanical fatigue Chemical degradation	Acceleration assistance Acceleration assistance	Increased plastic strain induced by increased ionomer deformation change Enhanced hydrogen peroxide formation rate
CL structural collapse PEM degradation	Surface cracking	Acceleration assistance	(i) Increased in PEM deformation (ii) Altered CL structural stability
	Thickness decrease Pore structure collapse TPB distortion Phase transitions in ion channels	Indeterminate Indeterminate Indeterminate Direct induction	Compression effect, carbon corrosion, or other factors Ionomer migration or other factors Ionomer migration or other factors Loss of water
	Glass transition	Direct induction	Destabilization of the electrostatic network
	Internal domain structure evolution	Direct induction	(i) Altered crystallinity of the hydrophobic region(ii) Alerted proton conduction pathway(ii) Increased extent of phase separation
	Chemical degradation Creep deformation	Acceleration assistance Acceleration assistance	Enhanced hydrogen peroxide formation rate Increased molecular mobility especially exceeding the glass transition
	Mechanical fatigue	Acceleration assistance	temperature (i) Increased fatigue amplitude (ii) Reduced mechanical strength of the membrane
CDI	Thermal softening	Direct induction	(i) Reduced storage elastic modulus of binder resins (ii) Decreased mechanical stiffness of PTFE
GDL degradation	Thermal oxidation MPL cracking	Direct induction Indeterminate	Oxidation of PTFE, carbon particles, and carbon fibre Water and gas transport, or other factors

5. Progress in overcoming the elevated temperature limitations

5.1. Development of PEMs with high proton conductivity at elevated temperatures

As previously discussed, when a MEA operates at elevated temperatures, particularly in the superheated (H) regime, the ionomer phase undergoes significant dehydration, thereby severely hindering proton transport. Concurrently, the substantial expulsion of water vapour markedly reduces the internal RH level within the MEA, resulting in a pronounced decline in the overall performance of the PEMFC. To overcome these performance losses associated with elevated temperature operation, considerable efforts have been devoted to developing new proton conductors in low RH and elevated temperature environments. To date, research progress in this field can be broadly classified into three approaches: (1) composite PFSA membranes with water-retentive fillers; (2) composite PFSA membranes doped with fillers with a high sulfonate concentration; and (3) modified PFSA membranes with additional proton conduction pathways.

For route (1), researchers have introduced inorganic fillers such as SiO2, TiO2, and CeO2 into electrolyte membranes using conventional casting and solvent evapouration methods. By enhancing the water retention of the membrane at elevated temperatures, these fillers effectively improve the proton conductivity of the PEM in elevated temperature environments. 235-238 However, the overall enhancement in proton conductivity is limited due to the disruption of continuous proton transport pathways and the agglomeration of inorganic fillers within the polymer matrix. To address the above issues, Nam et al.231 uniformly patterned mesoporous TiO₂ microporous plates (PTMPs) and embedded them onto the anode-facing surface of the Nafion membrane by employing microporous templates and an ionomer spraying technique, as shown in Fig. 19(a). This innovative approach protected the proton conduction pathways by precisely positioning TiO₂ microporous plates on the membrane surface while eliminating agglomeration typically associated with the introduction of hygroscopic inorganic fillers. As a result, the power density was more than 35.2% higher than that of the commercial Nafion 211 membrane at 120 °C and 35% RH, with a 24.5% reduction in the ohmic resistance. The hygroscopic mesoporous

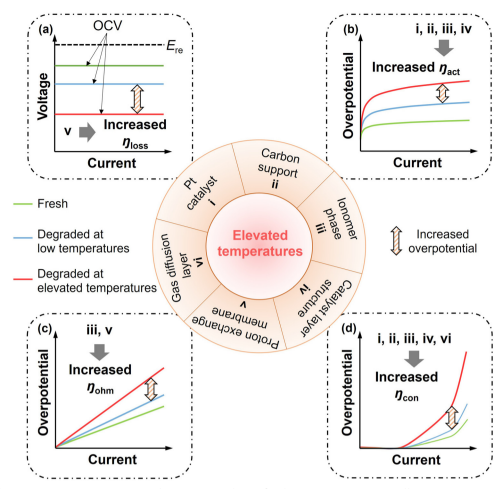
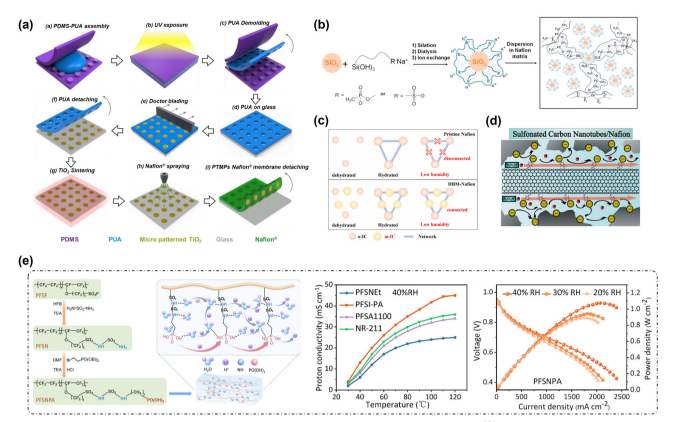


Fig. 18 Impacts of MEA component durability on performance loss of PEMFCs after the durability test at different temperatures

PTMPs embedded within the membrane attracted water molecules at elevated temperatures (>100 °C), helping mitigate the reduction in proton conductivity caused by membrane dehydration. Highly proton-conductive nanocomposite membranes were synthesised by incorporating surface-modified silica nanoparticles bearing different acidic functional groups (phosphonate or sulfonate ester groups) in the Nafion matrix, 232 as shown in Fig. 19(b). The short oligomeric corona containing phosphonate or sulfonate functional groups facilitated the formation of non-aggregated, discrete nanoparticles that synergistically interacted with the polymer matrix to form a membrane with significantly enhanced proton conductivity. The sulfonate-based nanocomposite exhibited a higher proton conductivity of 0.045 S cm⁻¹ at 130 °C and 30% RH. Nicotera et al.²³⁹ also integrated a Nafion matrix with TiO₂ nanoparticles directly grown on the surface of multi-walled carbon nanotubes (MWCNTs) to fabricate a Nafion/MWCNTs-TiO2 composite membrane. This synthesis approach enabled the nanoscale dispersion of TiO2 within the host matrix, thereby contributing to superior water-retention capability. The elongated carbon nanotubes provided additional pathways for proton conduction, and the assembled H₂/air single cell (with an active electrode area of 5 cm²) achieved a power output of 307.7 mW cm⁻² when operated at 120 °C and 30% RH.

For route (2), doping polymers with high sulfonic acid group density or functionalised carbon nanomaterials can be used to form a dense network of proton transport sites and bridging pathways within a membrane. For instance, Xu et al.233 synthesised a hyperbranched polyamide (HBM) featuring a high concentration of sulfonic acid groups and a nanoparticle morphology and incorporated it into a Nafion matrix without structural damage, as depicted in Fig. 19(c). The HBM was designed to imitate ion clusters and accelerate proton transport at elevated temperatures. The carboxyl groups of the HBM formed strong hydrogen bonds with the sulfonate groups of the Nafion matrix, thereby enhancing its structural compatibility and proton transport. As a result, the composite membrane exhibited a proton conductivity of 0.047 S cm⁻¹ at 110 °C and 60% RH, which was 1.9 times higher than that of pristine Nafion. Furthermore, the corresponding power density increased by 30%, demonstrating the feasibility of this approach for extending the operating temperature range of PFSA-based fuel cells. Yin et al. 12 prepared a composite membrane by incorporating sulfonated carbon nanotubes (Su-CNTs) into a Nafion matrix, as illustrated in Fig. 19(d). This significantly changed the membrane's microstructure, leading to the formation of numerous proton transport pathways, particularly under elevated temperature and low



(a) Schematic illustrations of the fabrication process of PTMPs embedded in the Nafion[®] membrane.²³¹ (b) Schematic illustration of the synthesis of functionalized nanoparticles and the final nanocomposite membranes.²³² (c) Proton conduction mechanism of pristine Nafion and HBM Nafion at different humidities.²³³ (d) Schematic illustration of enhanced proton transportation in Su-CNTs/Nafion membranes with proton highways along Su-CNTs. 12 (e) Schematic diagram of the synthesis process and proton conduction for the PFSNPA membrane. 234

RH conditions. The carbon nanotubes bridged isolated hydrated clusters, thereby greatly enhancing proton transport efficiency, giving the Su-CNTs/Nafion composite membrane a proton conductivity of $\sim 0.03 \text{ S cm}^{-1}$ at 130 °C and 50% RH, whereas that of the pristine Nafion remained below 0.01 S cm⁻¹ under the same conditions.

For route (3), Liao et al. 234 designed a novel PEM by precisely engineering the side-chain chemical structure of perfluorosulfonyl fluoride by introducing sulfonyl imide and phosphonic acid functional groups, as illustrated in Fig. 19(e). The sulfonyl imide groups exhibited super-acidic properties, while the phosphonic acid groups served as both proton donors and acceptors, and these two functional groups synergistically established unique and additional proton transport channels. Strong interactions between the acidic groups and water molecules enhanced water retention and molecular diffusion, thereby significantly improving the proton conductivity. The resulting perfluorosulfonimide-phosphonic acid (PFSNPA) membrane demonstrated a conductivity of 263 mS cm⁻¹ at 120 °C, and in H₂/air fuel cell tests, it delivered a power density of 1076.1 mW cm⁻² at 120 °C and 40% RH.

5.2. Optimization of operational strategies

To address the significant reduction in reactant partial pressures at elevated temperatures, along with the pronounced

dehydration of the ionomer phase within the MEA, particularly in low-RH environments, researchers have enhanced the performance of PEMFCs at elevated temperatures by solely optimizing operating strategies, without needing to develop new materials. As shown in Fig. 20(a), Wang et al.20 reported that increasing the operating pressure enhances the output voltage of PEMFCs by 50-150 mV, and this improvement was attributed to the higher oxygen concentration and improved membrane hydration. They found that a higher pressure reduced the heat generation, resulting in a 0.6-2 °C decrease in the temperature of the CCL, which also facilitated the accumulation and transport of water vapour and liquid water via CD flow. A higher operating pressure also helped minimise the mismatch in response rates between the ORR and the transport of oxygen and protons. Especially under higher loads, the Damköhler numbers of both oxygen and hydrogen decrease by up to 40% and 54%, respectively. Yang et al.8 also indicated that increases in the operating temperature must be accompanied by an increase in the operating pressure. As illustrated in Fig. 20(b), Lu et al. 21 demonstrated that to mitigate the impact of elevated temperatures on the saturated vapour pressure, increasing the inlet pressure of the fuel cell stack enhances both the steady-state and dynamic performance. Nevertheless, increasing the pressure is not a perfect solution, with Lu et al.²¹ demonstrating that α_{NMD} also increased with both temperature and pressure, making it

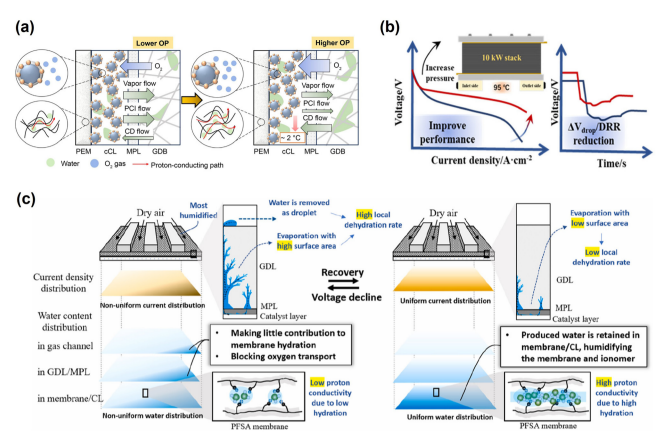


Fig. 20 (a) Schematic of operating pressure (OP) impacts on local mass transfer at higher loads: lower OP and higher OP.²⁰ (b) Schematic of the PEMFC performance improvement at elevated temperatures by increasing the inlet pressure.²¹ (c) Schematic of the performance loss and recovery mechanism, where the 'undesirable' and 'desirable' water distributions are shown on the left and right respectively. 105

challenging to mitigate ionomer dehydration on the anode side. Although increasing the cathode-side humidity can enhance back-diffusion and help alleviate excessive membrane drying at the anode, it simultaneously reduces the oxygen partial pressure. Furthermore, dynamic durability often deteriorates under a high pressure, resulting in limited performance improvements, and the associated trade-offs, particularly the potential reduction in fuel cell lifespan, have not been adequately addressed. Shao et al. 105 developed an optimised sequence for current loading and temperature elevation during PEMFC operation at elevated temperatures, as illustrated in Fig. 20(c). By first increasing the current before raising the temperature, they utilised the selfhumidification effect on the cathode side, which enabled a more uniform in-plane distribution of both current density and membrane hydration, even in dry environments.

5.3. Design of materials with enhanced durability at elevated temperatures

At elevated temperatures, catalyst growth and dissolution, carbon support corrosion, and the pronounced degradation of ionomers near or above their glass transition temperature collectively underscore the need to develop MEA components with enhanced thermal stability. As shown in Fig. 21(a), the development of stable core-shell spatially confined structures, the enhancement of interactions between Pt nanoparticles and

carbon supports, and the use of thermally corrosion-resistant graphitised carbon supports can improve the thermal stability of Pt/C catalysts. 9,244 Ji et al. 240 leveraged the corrosion resistance of TiO₂ and the strong metal-support interactions between Pt nanoparticles and TiO2 nanofibers, in combination with the high electrochemical conductivity of carbon nanofibers (CNFs), to synthesise a composite electrode consisting of CNT-wrapped Pt/TiO2 nanofibers. This design mitigated Pt dissolution and exhibited remarkable durability even at 120 °C and 40% RH. Zhang et al. 245 also exploited the thermal stability of TiO2 and introduced a nanolayer composed of TiO2 nanoparticles between the Pt catalyst and the carbon support, which protected the carbon from corrosion and established strong metal-support interactions between Pt and TiO2. After an AST at 150 °C, the maximum power density of the Pt/C@ TiO2-MEA decreased by only 8.6%, which was significantly lower than the 23.9% reduction observed for the conventional Pt/C-MEA. Therefore, TiO₂ may serve as an alternative support material at elevated temperatures due to its stability, but its inherently low electrical conductivity remains a critical limitation. One potential strategy to overcome this issue is to form composites with highly conductive materials.

Research on optimizing the thermal stability of ionomer materials in the CL is limited, but Yang et al.241 developed a novel α-aminoketone-linked covalent-organic framework (COF)

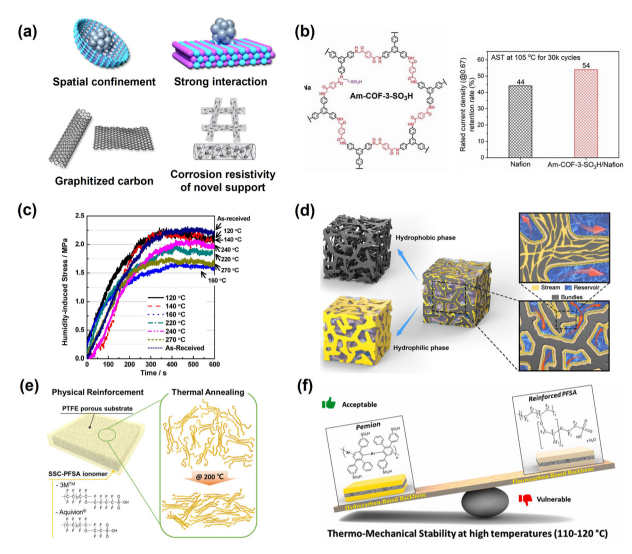


Fig. 21 (a) Mitigation strategies for catalyst and support degradation at elevated temperatures. 9,240 (b) Open framework ionomer with enhanced durability at elevated temperatures.²⁴¹ (c) Humidity-induced stress of the as-received Nafion 211 membrane and annealed Nafion 211 membranes.²⁰² (d) Three-dimensional structure scheme of the SSC-PFSA membrane with high durability at elevated temperatures. ²⁴² (e) Improved durability of short-sidechain perfluorinated polymer electrolyte membranes by the annealing process. 185 (f) High-temperature stability of hydrocarbon-based Pemion® proton exchange membranes.²⁴³

ionomer and then interweaved it with Nafion to function as "breathable" proton conductors. As depicted in Fig. 21(b), this innovative proton conductor retained 54% of its rated current density after 30k AST cycles at 105 °C, outperforming the 44% retention rate of Nafion ionomers. Further research is needed to enhance the thermal stability of ionomers within the CL because they play a pivotal role in facilitating electrochemical reactions and governing mass and proton transport throughout the CL.

To enhance the thermal stability of PEMs, existing research has primarily focused on three optimization strategies: (1) increasing the PEM crystallinity through thermal treatment; (2) employing short side-chain PEMs; and (3) incorporating thermally stable aromatic rings into the polymer backbone. While these approaches differ, they all help increase the glasstransition temperature of the membrane. Li et al. 202 reported that annealing Nafion membranes in the range of 120-160 °C

reduced the stress induced by humidity variations, as shown in Fig. 21(c). In addition, short side chain (SSC) PFSA membranes exhibited superior thermal stability compared with Nafion membranes due to their higher glass-transition temperatures.²⁰⁶ As depicted in Fig. 21(d), Guan et al.242 optimised the structure of SSC-PFSA membranes to form an internal "stream-reservoir" morphology, in which the hydrophobic domains served as physical crosslinking centers to maintain a high mechanical strength. As a result, the SSC-PFSA membranes are considered ideal candidates for high-power-density and heavy-duty fuel cells that typically operate in elevated temperature and lowhumidity environments. Moreover, Shin et al. 185 fabricated physically-reinforced SSC-PFSA membranes via annealing and incorporating porous substrates, which demonstrated significantly improved durability, as shown in Fig. 21(e). Specifically, a 3M 729 reinforced composite membrane annealed at 200 $^{\circ}\mathrm{C}$

withstood 16 600 cycles, markedly outperforming its non-annealed counterpart. Furthermore, Mirfarsi et al.243 indicated that the Pemion membrane, a commercially available mechanically reinforced sulfo-phenylated polyphenylene-based PEM, exhibited superior thermo-mechanical stability compared with a reinforced PFSA membrane in a temperature range of 30-120 °C and RH range of 10-90% RH, as illustrated in Fig. 21(f). The Young's modulus and strain hardening of the Pemion membrane remained largely insensitive to temperature, whereas the reinforced PFSA membrane exhibited substantial degradation in its mechanical properties above 90 °C. The Pemion membrane maintained an acceptable yield resistance across the entire test range, while the reinforced PFSA underwent spontaneous yielding above 110 °C under relatively small mechanical stress (e.g., 1 MPa). The pronounced stability of the Pemion membrane was primarily attributed to its rigid-rod polyphenylene backbone, which reduced its segmental mobility, resulting in a high glass transition temperature of 161 °C.

5.4. Diagnosis technique advancements

To gain deeper insights into the structural evolution of MEA during degradation, considerable efforts have recently been devoted to developing advanced characterization techniques. Key areas of progress include non-destructive spectroscopic analysis methods, novel electron microscopy, electrochemical analysis techniques based on impedance–relaxation time correlations, and non-destructive magnetic alignment-assisted imaging approaches.

In terms of non-destructive spectroscopic analysis methods, Janssen et al. 246 combined in situ small-angle and wide-angle Xray scattering (SAXS and WAXS) to establish a correlation between the structural parameters of the high-activity skeletal PtCo catalyst and its degradation during start-up/shutdown (SU/SD) cycling, as shown in Fig. 22(a). This integrated approach enabled the detailed in situ monitoring of structural changes within each crystalline phase and revealed two primary degradation mechanisms of the catalyst under SU/SD conditions: Co dissolution and crystallite growth. Bogar et al. 252 developed a SAXS-based analytical method coupled with electrochemical characterization to resolve the microstructural features of all functional layers within a MEA, including the ionomer, catalyst support, Pt nanoparticles, and the GDL. The Voigt peak was used to represent the ionomer peak and the matrix inflection point associated with the PEM. The Vulcan carbon support was modeled using the DAB model, while the CL was characterised by a population of Schulz-distributed spheres that aggregated into a mass fractal network that corresponded to Pt nanoparticle clusters. The GDL was represented using a power-law function to account for its larger structural features. Chen $\it et~al.^{253}$ employed synchrotron-based X-ray fluorescence (XRF) and X-ray absorption spectroscopy (XAS) to investigate the catalyst migration mechanism. A carefully designed MEA cross-sectional exposure method was developed by optimizing a PTFE gasket and minimizing interference through an integrated X-ray window. This configuration enabled the direct and in-depth observation of the internal of a fuel cell, facilitating in situ characterization. Interestingly, Li et al.254 designed a fuel cell compatible with operando XAS to investigate the structural evolution of manganese spinel oxide electrocatalysts in anion-exchange membrane fuel cells (AEMFCs). Notably, ionomer membranes may undergo beaminduced damage caused by X-rays, neutrons, or electron beams, thereby leading to morphological or chemical changes. Yu et al. 247 demonstrated a novel approach for distinguishing ionomer phases from carbon supports in a PEMFC by using carbon signals from scanning transmission electron microscopy coupled with electron energy loss spectroscopy (STEM-EELS), as presented in Fig. 22(b). This method enabled the high-spatialresolution mapping of both ionomer distributions and carbon supports. Although electron beam irradiation may damage samples, the study minimised such damage by carefully controlling the electron dose, even though it was not entirely eliminated. As shown in Fig. 22(c), Chabot et al. 248 employed small-angle neutron scattering (SANS) to investigate the nanoscale structural features of fuel cell electrodes, including the size distribution of Pt nanoparticles, the morphology of 2–3 nm thick ionomer films coating the catalyst surface, and the specific locations of water uptake within an electrode. This work highlighted the unique capability of SANS for resolving multiphase interfacial structures and water distribution in complex electrode architectures.

In terms of novel electron microscopy techniques, identicallocation transmission electron microscopy (IL-TEM) is a powerful technique for tracking structural evolution. Shokhen et al. 249 employed IL-TEM imaging based on ionic liquids to monitor the atomic-scale degradation of the top surface of a CL in an actual PEMFC. As shown in Fig. 22(d), this method was used to precisely track the same location over multiple imaging sessions, providing direct insight into nanoscale structural changes during degradation. Strandberg et al. combined identical-location scanning electron microscopy (IL-SEM) and IL-TEM to investigate the degradation of the cathode Pt/C catalyst layer in an actual operating PEMFC. Under SU/SD conditions, IL-SEM was used to observe the initiation and propagation of cracks within the electrode layer. Nanoscale IL-TEM imaging also showed an ~20% reduction in the average diameter of the primary carbon support particles, accompanied by an \sim 63% increase in platinum nanoparticle size due to their agglomeration and growth. These morphological changes corresponded to a 65% loss in the ECSA. The study identified the corrosion of structurally vulnerable regions of the carbon support as the key factor leading to catalyst layer collapse. Soleymani et al.255 developed an epoxy-free ultramicrotomy technique that, when combined with TEM, enabled the first direct visualization of an ionomer network, carbon particle morphology, and Pt nanoparticle distribution. This method also provided a detailed observation of structural changes during degradation.

The distribution of relaxation time (DRT) analysis based on electrochemical impedance spectroscopy allows for the *in situ* and non-destructive tracking of the evolution of characteristic peaks associated with different reactions and transport processes within a MEA. This provides insights into the degradation mechanisms of the internal components, ^{250,256} as shown in Fig. 22(e). Interestingly, Sun *et al.* ²⁵¹ developed a magnetic array imaging system

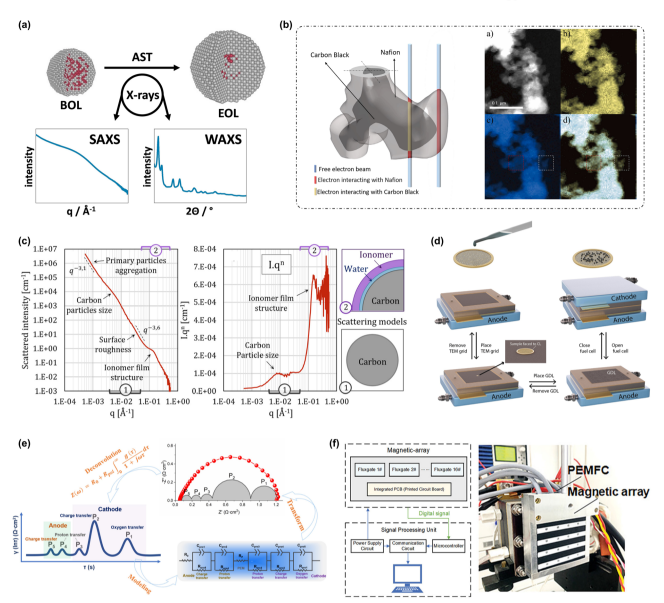


Fig. 22 (a) In situ SAXS & WAXS techniques monitoring the morphological changes of catalysts. ²⁴⁶ (b) Identification of ionomers in PEMFCs by EELS. ²⁴⁷ (c) Structural information details of the three-phase boundary accessible from the SANS profile of MEA.²⁴⁸ (d) Schematic illustration of identical location transmission electron microscopy.²⁴⁹ (e) DRT technology decoupling of polarization process of PEMFCs.²⁵⁰ (f) Schematics of a designed magnetic array system and a working diagram of a magnetic array. 251

based on magnetic sensing principles to detect variations in external magnetic fields induced by internal current changes to visualise the heterogeneity of degradation in PEMFCs, as presented in Fig. 22(f). This technique was non-destructive, easy to implement, and did not require any modification of the tested structures or materials. It has particular advantages for characterizing spatial heterogeneities in structural degradation within fuel cell stacks, but its accuracy depends on eliminating interference from magnetic contaminants adhered to the PEMFC.

5.5. Exploration of temperature effects on the MEA of anion exchange membrane water electrolyzers

Anion-exchange membrane water electrolyzers (AEMWEs) use the anion-exchange membrane (AEM) to conduct OH ions and separate hydrogen and oxygen, as illustrated in Fig. 23(a). Their structural configuration is similar to that of PEMFCs, with both systems based on MEAs, but the materials differ. Given the significant influence of elevated temperatures on a PEMFC's performance, this section briefly explores how temperature affects the efficiency and stability of MEA operation in an AEMWE. Table 12 summarises the performance and durability data of AEMWEs at different temperatures.

In AEMWE systems, the operating temperature is typically limited to below 100 °C and in the range of 50-90 °C. Temperature is a critical operational parameter, where a higher temperature leads to a lower thermodynamic equilibrium potential, enhanced electrochemical reaction kinetics, and improved ionic conductivity, all of which further improve the

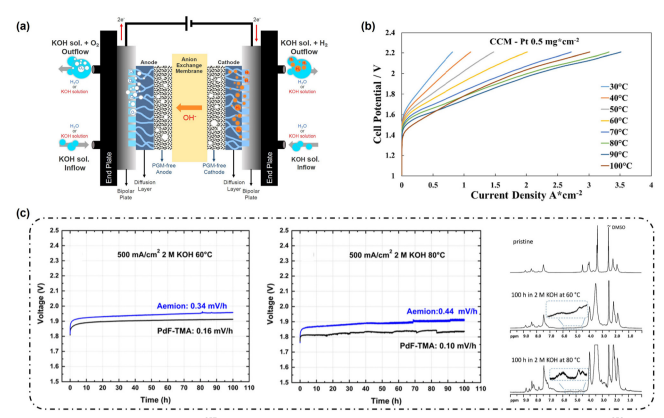


Fig. 23 (a) Schematic diagram of the AEMWE.²⁵⁷ (b) Polarization curves for AEMWE on the CCM configuration at different temperatures.²⁵⁸ (c) Stability evaluations at 500 mA cm⁻² in 2 M KOH at 60 °C and 80 °C and ¹H NMR spectra of the Aemion membrane before and after long-term stability evaluation in the electrolysis cell.²⁵⁹

output performance, as shown in Table 12. Lim et al. 261 systematically investigated the influence of temperature within the 50-90 °C range and reported that under a constant overpotential of 591 mV, the current density increased by approximately 50 mA cm⁻² for every 10 °C increase. This enhancement was attributed to improved ionic conductivity in the membrane, ionomer, and electrolyte, as well as accelerated electrochemical kinetics. As shown in Fig. 22(b), raising the cell temperature to 90 °C consistently resulted in higher current densities due to improved oxygen and hydrogen evolution reaction kinetics and elevated ionic conductivity. 258 However, further increasing the temperature to 100 $^{\circ}\mathrm{C}$ caused the performance to decline due to an increase in series resistance (R_S) from 0.09 Ω cm² at 90 °C to $0.10 \Omega \text{ cm}^2$ at 100 °C and a rise in charge-transfer resistance (R_{ct}) from 0.095 Ω cm² to 0.105 Ω cm². These increases indicate a deterioration of the interface between the PEM and the CL due to dehydration at 100 °C. Therefore, there is an optimal operating temperature range for AEMWEs, and avoiding operation near the evapouration temperature of water is crucial to maintaining their performance and interfacial stability.

Temperature also significantly impacts the stability of MEAs, especially AEMs, as presented in Table 12. Lin et al. 266 evaluated the stability of AEMs by immersing membrane samples in aqueous KOH solutions at 60 °C and 80 °C and monitoring the hydroxide ion conductivity over time. After 480 h, the hydroxide conductivity decreased by 6.7% at 60 $^{\circ}$ C and by 11.1% at 80 $^{\circ}$ C, indicating that higher temperatures accelerated membrane degradation. Furthermore, as shown in Fig. 22(c), the performance of Aemion cells significantly declined over time, and the degradation rate at 80 °C (0.44 mV h⁻¹) was higher than that at 60 °C (0.34 mV h⁻¹). ¹H NMR analysis of membranes before and after degradation revealed detectable structural changes, further confirming that higher temperatures accelerated polymer degradation, leading to faster performance losses.

6. Conclusions and perspectives

Elevated temperatures within MEAs have become a significant focus of research to advance PEMFCs, but current methodologies face significant challenges in ensuring the stable, long-term operation of PEMFCs at elevated temperatures. These limitations include difficulties in mass transfer, charge conduction, electrochemical reactions, component degradation, and water and gas management within the MEA. This review provided a comprehensive overview of the challenges associated with ensuring stable MEAs at elevated temperatures and highlighted key gaps and potential future directions (Fig. 24).

(1) Study on the correlation between the microscale and macroscale thermal environments. At different elevated temperature levels, the transport and degradation of MEA components exhibit different characteristics. A fundamental prerequisite for

Table 12 Performance and durability data of AEMWE at different temperatures

Serial number	MEA configurations	Items	Values	Ref.
1	FAA3-50 membrane, IrO ₂ (anode catalyst), Pt/C (cathode catalyst), 1 M KOH (feed);	Performance	At 1.8 V	258
	(cathode catalyst), 1 M ROH (leed);		0.6 A cm ⁻² @ 60 °C	
			0.9 A cm ⁻² @ 70 °C	
			1.05 A cm ⁻² @ 80 °C	
			At 2 V:	
			1.2 A cm ⁻² @ 60 °C	
			1.7 A cm ⁻² @ 70 °C	
			2.1 A cm ⁻² @ 80 °C	
2	FAA3-50 membrane, IrO ₂ (anode catalyst), Pt/C	Performance	At 1.8 V	260
_	(cathode catalyst), 1 M KOH (feed);	Terrormance	At 1.0 V	200
	(cathode catalyst), I w Roll (leed),		0.73 A cm ⁻²	
			$0.73~\mathrm{A~cm}^{-2}$ @ $50~^{\circ}\mathrm{C}$ $1.15~\mathrm{A~cm}^{-2}$ @ $70~^{\circ}\mathrm{C}$	
			At 2 V	
			1.42 A cm ⁻² @ 50 °C,	
			1.83 A cm ⁻² @ 70 °C	
3	FAA-3-PK-75 membrane, IrO ₂ (anode catalyst), Pt/C	Performance	At 1.8 V	261
	(cathode catalyst), 0.5 M KOH (feed);			
	<i>,</i> ,,		$0.46~\mathrm{A~cm^{-2}}$ @ $50~^\circ\mathrm{C}$	
			0.607 A cm ⁻² (a) 60 °C	
			0.607 A cm^{-2} @ $60 ^{\circ}\text{C}$ 0.625 A cm^{-2} @ $70 ^{\circ}\text{C}$	
			0.87 A cm ⁻² @ 80 °C	
			0.87 A cm ⁻² @ 80 °C 0.983 A cm ⁻² @ 90 °C	
4	Fumatech FAA-3-50 membrane, IrO ₂ (anode	Performance	At 2 V	262
	catalyst), Pt/C (cathode catalyst), 1 M KOH (feed)			
			$3.5~\mathrm{A~cm^{-2}}$ @ $80~^\circ\mathrm{C}$	
		Durability	50 h	
5	Aemion membrane, NiFe ₂ O ₄ (anode catalyst),	Durability	$0.34~\mathrm{mV~h^{-1}}$ @ $60~^{\circ}\mathrm{C}$	259
	RANEY [®] nickel (cathode catalyst), 2 M KOH (feed)			
			$0.44~\mathrm{mV~h^{-1}}$ @ 80 $^{\circ}\mathrm{C}$	
6	A201 membrane, IrO ₂ (anode catalyst), Pt/C (cath-	Performance	At 1.8 V	263
	ode catalyst), 1 M KOH (feed)			
			$0.399~\mathrm{A~cm}^{-2}$ @ $50~^{\circ}\mathrm{C}$;	
		Durability	500 h	
7	AF1-HNN8-50 membrane, NiS ₂ /Ni ₃ S ₄ (anode	Performance	At 2.0 V	264
	catalyst), Pt/C (cathode catalyst), 1 M KOH (feed)			
			1.8 A @ 60 °C	
		Durability	120 mV kh $^{-1}$ @ 60 $^{\circ}$ C	
8	PBPA membrane, NiFe ₂ O ₄ -X (anode catalyst), Pt/C	Performance	At 1.77 V	265
	(cathode catalyst), 1 M KOH (feed)			
			2.0 A @ 80 °C	
		Durability	240 mV kh ⁻¹ @ 80 °C	

clarifying these mechanisms is to accurately characterise the thermal environment experienced by each functional layer and its internal nanoscale constituents. The CL, whose nanoscale structure is comprised of a Pt catalyst, carbon support, and ionomer, is highly sensitive to local thermal conditions that directly affect electrochemical reaction kinetics, mass transport pathways, and material durability. However, existing temperature sensing techniques, such as thin-film microsensors, 267 only provide localised average temperatures at the CL surface and cannot resolve heterogeneous thermal fields within the nanostructured interior. To address this, systematic investigations into the multi-scale heat transfer mechanisms within MEAs are needed. Specifically, an understanding of the thermal conduction across functional layers and at interfaces between nanoscale materials is essential. Establishing a quantitative correlation between microscopic thermal environments and macroscopic operating temperatures will enable the precise determination of the actual elevated temperatures experienced by MEA components in various operating scenarios.

(2) Development of ionomers with high proton conductivity and anti-dehydration optimization. At elevated temperatures,

particularly in the super-boiling regime (H), the ionomer undergoes substantial dehydration, which significantly hinders proton transport, especially on the anode side. To address this, it is essential to elucidate the equilibrium mechanisms governing the interconversion between liquid water, membranous water, and water vapour. Special attention should be given to identifying the key structural features that dictate dehydration behaviour in nanoscale ionomer films. It is also necessary to establish quantitative correlations with high-frequency resistance to enable the real-time assessment of an ionomer's hydration state. From a material innovation perspective, the design of microporous architectures that promote localised condensation, along with highly water-retentive and multi-channel fast proton-conducting ionomer phases, is a promising approach. In terms of optimizing operation, the utilization of cathode-generated water and enhancing the cathode-anode humidity gradient improve the backdiffusion of water, thereby helping replenish water at the anode. Optimizing the control between current density and temperature rise is critical for mitigating severe ionomer dehydration at elevated temperatures.

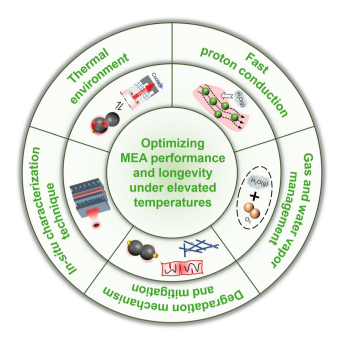


Fig. 24 Approaches to optimizing MEA performance and longevity at elevated temperatures

(3) Optimization of cathode gas and water vapour management: at elevated temperatures, excess water evapouration is the primary cause of reduced reactant gas partial pressures. On the cathode side, although humidification can alleviate ionomer dehydration, it also lowers the partial pressure of reactant gases. Therefore, it is crucial to identify the minimum flow channel pressure threshold required to ensure an adequate gas supply, but maintaining this threshold often necessitates a high operating pressure for PEMFCs at elevated temperatures, which may compromise durability under dynamic conditions. This highlights the need for advanced pressure control techniques to manage the anode-cathode pressure differences. The development of MEA porous networks with low mass transport resistance and the optimization of flow field designs are essential to ensure rapid and efficient delivery of reactant gases at elevated temperatures.

(4) Study on MEA coupled degradation mechanisms and mitigation strategies: the degradation of MEA components is markedly accelerated at elevated temperatures, which includes the growth and dissolution of Pt catalysts, the corrosion of carbon supports, and the deterioration of both the ionomer and GDL. Within the CL and PEM, the ionomer phase undergoes significant structural changes in its internal chain and domain structures, which cause increased mechanical fatigue and chemical decomposition. The GDL is similarly susceptible to thermal softening and oxidative degradation. Notably, the coupled degradation mechanisms of Pt catalysts at elevated temperatures and varying electrochemical potentials remain insufficiently understood. While elevated temperatures induced glass transitions have been confirmed in PEMs, the glass transition of ionomers confined within the CL remains poorly characterised due to their nanoscale thickness and

heterogeneous morphologies. The coupled chemo-mechanical degradation of ionomers near or above their glass transition temperatures also remains largely unexplored. Furthermore, the implications of increased molecular mobility on the deformation and adhesion properties of ionomers warrant in-depth investigations, as these factors are critical for understanding the structural collapse of the CL, where ionomers serve as functional binders. Humidity and pressure also play pivotal roles in the degradation of both CL and PEM materials and should be systematically examined. For the GDL, the long-term thermal stability of bonding resins, hydrophobic PTFE, and the MPL needs to be thoroughly investigated. Although the MPL shows more pronounced cracking at elevated temperatures, the increased water vapour fraction reduces its ability to uniformly expel liquid water. Removing the MPL from the GDL structure provides an option for the next-generation design of MEA operating at elevated temperatures to simplify the fabrication, reduce ohmic losses, and enhance the overall durability of MEA components. Importantly, the individual contributions of each MEA component to performance degradation at elevated temperatures are not yet clearly delineated, complicating efforts to identify the primary degradation-limiting factor. Establishing a quantitative relationship between component degradation and cell performance loss is essential for guiding the rational design of PEMFCs with enhanced durability during operation at elevated temperatures.

In terms of material innovations, enhancing the catalyst durability at elevated temperatures necessitates the development of thermally stable catalysts with spatial confinement, strong metal-support interactions, and corrosion-resistant supports (graphitised carbon, TiO₂, etc.). In parallel, the development of ionomer materials with low segmental mobility (i.e., high glass transition temperatures) is essential for maintaining mechanical integrity and proton conductivity at elevated temperatures. Strategies such as appropriate thermal treatment, short side-chain structures, or the incorporation of aromatic rings into the polymer backbone can enhance the structural stability of ionomers. However, attention must be paid to preserving the proton conduction channels. Regarding the GDL, future efforts should elucidate the underlying degradation mechanisms in elevated temperature environments to inform the development of targeted durability enhancement strategies.

(5) Development of in situ characterization techniques: highresolution characterization of MEAs, particularly within the CL, is critical to unravel the complex degradation mechanisms associated with the TPB and porous transport structures. However, conventional techniques face several intrinsic limitations. Most techniques are destructive, providing only ex situ postmortem analysis with limited relevance to real-time degradation dynamics. The nanoscale dimensions of CL components demand exceptionally high spatial resolution, and the carbonaceous nature of both supports and ionomers hampers clear distinction due to low material contrast. Ionomer structures are often vulnerable to damage or changes under high-energy particle beams, which compromises the reliability of observations. These challenges highlight the need to develop in situ

and non-destructive multi-dimensional characterization techniques. Promising directions include non-destructive in situ spectroscopic methods, integrating electron microscopy with optimised sample preparation protocols and visualization techniques based on electrical, magnetic, or acoustic principles. Such integrated approaches may provide deeper insights into the dynamic structural evolution and degradation pathways of MEA components under realistic operating conditions.

Author contributions

Caizheng Yue: writing - original draft, investigation, visualization, formal analysis; Weibo Zheng: writing - reviewing and editing, supervision, funding acquisition; Qiuya Wang: investigation, formal analysis; Zhendong Wang: investigation; Bing Li: supervision. Cunman Zhang: supervision; and Pingwen Ming: writing - reviewing and editing, investigation, supervision, funding acquisition.

Data availability

Data will be made available upon request.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

This work was supported by the National Key Research and Development Program of China (2023YFB2504201) and National Natural Science Foundation of China (no. 52276210 and 52106255).

References

- 1 X. Lü, Y. Qu, Y. Wang, C. Qin and G. Liu, Energy Convers. Manage., 2018, 171, 1273-1291.
- 2 N. L. Panwar, S. C. Kaushik and S. Kothari, Renewable Sustainable Energy Rev., 2011, 15, 1513-1524.
- 3 M. A. Sadeghi, Z. A. Khan, M. Agnaou, L. Hu, S. Litster, A. Kongkanand, E. Padgett, D. A. Muller, T. Friscic and J. Gostick, Appl. Energy, 2024, 353, 122004.
- 4 T. Lochner, R. M. Kluge, J. Fichtner, H. A. El-Sayed, B. Garlyyev and A. S. Bandarenka, ChemElectroChem, 2020, 7, 3545-3568.
- 5 J. Xu, C. Zhang, Z. Wan, X. Chen, S. H. Chan and Z. Tu, Renewable Sustainable Energy Rev., 2022, 155, 111908.
- 6 E. Jannelli, M. Minutillo and A. Perna, Appl. Energy, 2013, **108**, 82-91.
- 7 K. Jiao, J. Xuan, Q. Du, Z. Bao, B. Xie, B. Wang, Y. Zhao, L. Fan, H. Wang, Z. Hou, S. Huo, N. P. Brandon, Y. Yin and M. D. Guiver, Nature, 2021, 595, 361-369.

- 8 P. C. C. Yang, S. Srinivasan, J. Benziger and A. B. Bocarsly, J. Power Sources, 2001, 103, 1-9.
- 9 Q. Meyer, C. Yang, Y. Cheng and C. Zhao, Electrochem. Energy Rev., 2023, 6, 16.
- 10 P. A. García-Salaberri and A. Sánchez-Ramos, Appl. Energy, 2024, 367, 123332.
- 11 Y. Shao, L. Xu, Z. Hu, L. Xu, X. Zhang, G. Zhao, J. Li and M. Ouyang, J. Power Sources, 2023, 581, 233520.
- 12 G. Xu, X. Dong, B. Xue, J. Huang, J. Wu and W. Cai, Energies, 2023, 16, 1565.
- 13 J. Z. Yanghua Tang, C. Song, H. Liu, J. Zhang, H. Wang, S. Mackinnon, T. Peckham, J. Li, S. McDermid and P. Kozak, J. Electrochem. Soc., 2006, 153, A2036-A2043.
- 14 T. S. T.-C. Jao, S. Uemura, T. Yoshida and S. Hirai, ECS Trans., 2016, 75, 179-188.
- 15 E. Lufrano, C. Simari, M. L. Di Vona, I. Nicotera and R. Narducci, Polymers, 2021, 13, 359.
- 16 M. Butori, B. Eriksson, N. Nikolić, C. Lagergren, G. Lindbergh and R. W. Lindström, J. Power Sources, 2023, 563, 232803.
- 17 D. R. P. Morris, S. P. Liu, D. Villegas Gonzalez and J. T. Gostick, ACS Appl. Mater. Interfaces, 2014, 6, 18609-18618.
- 18 M. Tesfaye, D. I. Kushner, B. D. McCloskey, A. Z. Weber and A. Kusoglu, ACS Macro Lett., 2018, 7, 1237-1242.
- 19 C. Zhang, M. Davies and K. Karan, J. Polym. Sci., Part B: Polym. Phys., 2019, 57, 343-352.
- 20 Q. Wang, J. Lu, W. Zheng, B. Li, J. P. Zheng, G. Cui, L. Hao and P. Ming, Fuel, 2025, 381, 133623.
- 21 Y. Lu, H. Wu, D. Yang, W. Zhuge, P. Ming, Y. Zhang, J. Chen and X. Pan, Energy Fuels, 2024, 38, 7331-7343.
- 22 T. Ruiu, A. M. Dreizler, J. Mitzel and E. Gülzow, J. Power Sources, 2016, 303, 257-266.
- 23 Q. Wang, B. Li, D. Yang, H. Dai, J. P. Zheng, P. Ming and C. Zhang, J. Power Sources, 2021, 492, 229613.
- 24 B. R. Frieberg, K. A. Page, J. R. Graybill, M. L. Walker, C. M. Stafford, G. R. Stafford and C. L. Soles, ACS Appl. Mater. Interfaces, 2016, 8, 33240-33249.
- 25 J. Ramousse, S. Didierjean, O. Lottin and D. Maillet, Int. J. Therm. Sci., 2008, 47, 1-6.
- 26 K. Suárez-Alcántara, A. Rodríguez-Castellanos, R. Dante and O. Solorza-Feria, *J. Power Sources*, 2006, **157**, 114–120.
- 27 M. V. V. D. B. Sepa, J. Electrochem. Soc., 1987, 134, 845-848.
- 28 Y. S. Hui Xu, H. Russell Kunz and J. M. Fenton, J. Electrochem. Soc., 2005, 152, A1828-A1836.
- 29 H. Yano, J. Inukai, H. Uchida, M. Watanabe, P. K. Babu, T. Kobayashi, J. H. Chung, E. Oldfield and A. Wieckowski, Phys. Chem. Chem. Phys., 2006, 8, 4932.
- 30 V. Ramani, H. R. Kunz and J. M. Fenton, J. Membr. Sci., 2004, 232, 31-44.
- 31 S. Bose, T. Kuila, T. X. H. Nguyen, N. H. Kim, K.-T. Lau and J. H. Lee, Prog. Polym. Sci., 2011, 36, 813-843.
- 32 R. Haider, Y. Wen, Z.-F. Ma, D. P. Wilkinson, L. Zhang, X. Yuan, S. Song and J. Zhang, Chem. Soc. Rev., 2021, 50, 1138-1187.
- 33 J. N. Adam and Z. Weber, J. Electrochem. Soc., 2006, 153, A2205-A2214.
- 34 K. P. Peter Berg, J. St. Pierre, J. Stumper and B. Wetton, J. Electrochem. Soc., 2004, 151, A341-A335.

- 35 R. H. Qingfeng Li, J. Oluf Jensen and N. J. Bjerrum, *Chem. Mater.*, 2003, **15**, 4896–4915.
- 36 A. Liu, X. Liu, L. Liu, Y. Pu, K. Guo, W. Tan, S. Gao, Y. Luo, S. Yu, R. Si, B. Shan, F. Gao and L. Dong, ACS Catal., 2019, 9, 7759–7768.
- 37 R. H. Qingfeng Li, J.-A. Gao, J. Oluf Jensen and N. J. Bjerrum, *J. Electrochem. Soc.*, 2003, **150**, A1599–A1605.
- 38 S. G. Kandlikar and Z. Lu, *Appl. Therm. Eng.*, 2009, **29**, 1276–1280.
- 39 N. Alhazmi, M. S. Ismail, D. B. Ingham, K. J. Hughes, L. Ma and M. Pourkashanian, J. Power Sources, 2013, 241, 136–145.
- 40 O. S. Burheim, H. Su, H. H. Hauge, S. Pasupathi and B. G. Pollet, *Int. J. Hydrogen Energy*, 2014, 39, 9397–9408.
- 41 M. Ahadi, M. Tam, M. S. Saha, J. Stumper and M. Bahrami, *J. Power Sources*, 2017, 354, 207–214.
- 42 O. S. Burheim and J. G. Pharoah, *Curr. Opin. Electrochem.*, 2017, 5, 36–42.
- 43 B. G. P. Odne and S. Burheim, ECS Trans., 2018, 86, 97–109.
- 44 J. Zhou, S. Shukla, A. Putz and M. Secanell, *Electrochim. Acta*, 2018, **268**, 366–382.
- 45 A. Goshtasbi, P. García-Salaberri, J. Chen, K. Talukdar, D. G. Sanchez and T. Ersal, *J. Electrochem. Soc.*, 2019, 166, F3154–F3179.
- 46 R. Bock, H. Karoliussen, B. G. Pollet, M. Secanell, F. Seland, D. Stanier and O. S. Burheim, *Int. J. Hydrogen Energy*, 2020, 45, 1335–1342.
- 47 Q. Chen, G. Zhang, X. Zhang, C. Sun, K. Jiao and Y. Wang, *Appl. Energy*, 2021, **286**, 116496.
- 48 W. Wang, Z. Qu, X. Wang and J. Zhang, *Membranes*, 2021, 11, 148.
- 49 Y. Huang, X. Xiao, H. Kang, J. Lv, R. Zeng and J. Shen, *Energy Convers. Manage.*, 2022, **254**, 115221.
- 50 L. Yang, N.-N. Nik-Ghazali, M. A. H. Ali, W. T. Chong, Z. Yang and H. Liu, Renewable Sustainable Energy Rev., 2023, 187, 113737.
- 51 A. U. Yakubu, J. Zhao, Q. Jiang, X. Ye, J. Liu, Q. Yu and S. Xiong, *Heliyon*, 2024, **10**, e38556.
- 52 Z. Zhang, J. Mao, H. Wei, C. Cheng and Z. Liu, *Appl. Therm. Eng.*, 2024, **239**, 122083.
- 53 G. Unsworth, N. Zamel and X. Li, *Int. J. Hydrogen Energy*, 2012, 37, 5161–5169.
- 54 O. S. Burheim, H. Su, S. Pasupathi, J. G. Pharoah and B. G. Pollet, *Int. J. Hydrogen Energy*, 2013, **38**, 8437–8447.
- 55 J. G. P. Odne, S. Burheim, H. Lampert, P. J. S. Vie and S. Kjelstrup, *J. Fuel Cell Sci. Technol.*, 2011, **8**, 021013.
- 56 B. H. Robert Bock, M. Onsrud, H. Karoliussen, F. Seland and O. S. Burheim, *ECS Trans.*, 2019, **92**, 223–245.
- 57 Z. L. A. Radhakrishnan and S. G. Kandlikar, *ECS Trans.*, 2010, 33, 1163–1176.
- 58 G. Karimi, X. Li and P. Teertstra, *Electrochim. Acta*, 2010, 55, 1619–1625.
- 59 V. Leduc, G. Sdanghi, R. Bligny, J. Dillet, S. Didierjean, T. Schmitt, M. Hanauer, U. Sauter and G. Maranzana, *Electrochim. Acta*, 2024, 505, 145001.

- 60 F. Liu, Q. Wang, B. Li, C. Zhang and P. Ming, eTransportation, 2022, 12, 100162.
- 61 I. Nitta, O. Himanen and M. Mikkola, *Fuel Cells*, 2008, **8**, 111–119.
- 62 M. Ahadi, M. Andisheh-Tadbir, M. Tam and M. Bahrami, *Int. J. Heat Mass Transfer*, 2016, **96**, 371–380.
- 63 Y. Wang and M. Gundevia, Int. J. Heat Mass Transfer, 2013, 60, 134–142.
- 64 A. Pfrang, D. Veyret, F. Sieker and G. Tsotridis, *Int. J. Hydrogen Energy*, 2010, 35, 3751–3757.
- 65 O. Burheim, P. J. S. Vie, J. G. Pharoah and S. Kjelstrup, J. Power Sources, 2010, 195, 249–256.
- 66 O. S. Burheim, G. Ellila, J. D. Fairweather, A. Labouriau, S. Kjelstrup and J. G. Pharoah, *J. Power Sources*, 2013, 221, 356–365.
- 67 C. Csoklich, M. Sabharwal, T. J. Schmidt and F. N. Büchi, *J. Power Sources*, 2022, **540**, 231539.
- 68 E. Sadeghi, N. Djilali and M. Bahrami, *J. Power Sources*, 2011, **196**, 246–254.
- 69 S. Wang, J. Wang and Y. Zhu, *J. Power Sources*, 2023, 575, 233179.
- 70 G. Xu, J. M. LaManna, J. T. Clement and M. M. Mench, J. Power Sources, 2014, 256, 212-219.
- 71 N. Zamel and X. Li, Prog. Energy Combust. Sci., 2013, 39, 111–146.
- 72 L. Chen, Y.-F. Wang and W.-Q. Tao, *Therm. Sci. Eng. Prog.*, 2020, **19**, 100616.
- 73 H. Sadeghifar, N. Djilali and M. Bahrami, *J. Power Sources*, 2015, 273, 96–104.
- 74 T.-F. Cao, Y.-T. Mu, J. Ding, H. Lin, Y.-L. He and W.-Q. Tao, *Int. J. Heat Mass Transfer*, 2015, **87**, 544–556.
- 75 O. S. Burheim, ECS Trans., 2017, 80, 509-525.
- 76 Q. Wang, F. Tang, B. Li, H. Dai, J. P. Zheng, C. Zhang and P. Ming, J. Power Sources, 2021, 488, 229419.
- 77 Q. Wang, F. Tang, B. Li, H. Dai, J. P. Zheng, C. Zhang and P. Ming, *Appl. Energy*, 2022, 308, 118377.
- 78 Q. Wang, F. Tang, X. Li, J. P. Zheng, L. Hao, G. Cui and P. Ming, *Chem. Eng. J.*, 2023, 463, 142286.
- 79 H. Pourrahmani, A. Yavarinasab, M. Siavashi, M. Matian and J. Van herle, *Energy Rev.*, 2022, **1**, 100002.
- 80 F. Cai, S. Cai and Z. Tu, Energy Convers. Manage., 2024, 307, 118348.
- 81 Z. Z. Mufan Li, T. Cheng, A. Fortunelli, C.-Y. Chen, R. Yu, Q. Zhang, L. Gu, B. V. Merinov, Z. Lin, E. Zhu, T. Yu, Q. Jia, J. Guo, L. Zhang, W. A. Goddard, Y. Huang and X. Duan, *Science*, 2016, 354, 1414–1419.
- 82 T. J. Bvumbe, P. Bujlo, I. Tolj, K. Mouton, G. Swart, S. Pasupathi and B. G. Pollet, *Hydrogen Fuel Cells*, 2016, 1, 1–20.
- 83 J. N. Qiao, H. Guo, F. Ye and H. Chen, *Int. J. Heat Mass Transfer*, 2024, **226**, 125456.
- 84 W. Yoon and A. Z. Weber, *J. Electrochem. Soc.*, 2011, **158**, B1007.
- 85 D. Banham, J. Zou, S. Mukerjee, Z. Liu, D. Yang, Y. Zhang, Y. Peng and A. Dong, *J. Power Sources*, 2021, 490, 229515.
- 86 S. Liu, S. Yuan, Y. Liang, H. Li, Z. Xu, Q. Xu, J. Yin, S. Shen, X. Yan and J. Zhang, *Int. J. Hydrogen Energy*, 2023, 48, 4389–4417.

- 87 S. Kreitmeier, P. Lerch, A. Wokaun and F. N. Büchi, J. Electrochem. Soc., 2013, 160, F456-F463.
- 88 K.-H. Hou, C.-H. Lin, M.-D. Ger, S.-W. Shiah and H.-M. Chou, Int. J. Hydrogen Energy, 2012, 37, 3890-3896.
- 89 L. Chen, Y. Chen and W.-Q. Tao, Renewable Sustainable Energy Rev., 2023, 173, 113050.
- 90 A. Kusoglu and A. Z. Weber, Chem. Rev., 2017, 117, 987-1104.
- 91 J. Zhang, Y. Zhang, Z. Xiao, J. Tan, H. Zhang and J. Yu, Membranes, 2024, 14, 72.
- 92 F. Akitomo, T. Sasabe, T. Yoshida, H. Naito, K. Kawamura and S. Hirai, J. Power Sources, 2019, 431, 205-209.
- 93 H. Ito, T. Mimoto, S. Someya and T. Munakata, J. Electrochem. Soc., 2021, 168, 124505.
- 94 J. Zhang, Z. Xie, J. Zhang, Y. Tang, C. Song, T. Navessin, Z. Shi, D. Song, H. Wang, D. P. Wilkinson, Z.-S. Liu and S. Holdcroft, J. Power Sources, 2006, 160, 872-891.
- 95 V. Ionescu, Phys. Scr., 2020, 95, 034006.
- 96 T. Yin, D. Chen, T. Hu, S. Hu, R. Li, T. Wei, Y. Li, Y. Li, X. Xu and P. Pei, Energy Convers. Manage., 2024, 315, 118740.
- 97 J. Shen, C. Zhang, L. Li, S. Liu, H. Liu, B. Chen and C. Du, Energy, 2024, 308, 132851.
- 98 A. Martín-Alcántara, J. Pino and A. Iranzo, Int. J. Hydrogen Energy, 2023, 48, 13987-13999.
- 99 C. Song, Y. Tang, J. L. Zhang, J. Zhang, H. Wang, J. Shen, S. McDermid, J. Li and P. Kozak, Electrochim. Acta, 2007, 52, 2552-2561.
- 100 H. Sun, C. Xie, H. Chen and S. Almheiri, Appl. Energy, 2015, **160**, 937-944.
- 101 J. Zhang, Y. Tang, C. Song and J. Zhang, J. Power Sources, 2007, 172, 163-171.
- 102 J. Zhang, Y. Tang, C. Song, Z. Xia, H. Li, H. Wang and J. Zhang, Electrochim. Acta, 2008, 53, 5315-5321.
- 103 S. Wang, X. Hu, Z. Zhou, Q. Wang, K. Ye, S. Sui, M. Hu and F. Jiang, Sustainable Energy Technol. Assess., 2023, 60, 103529.
- 104 Y. Xu, R. Fan, G. Chang, S. Xu and T. Cai, Energy Convers. Manage., 2021, 248, 114791.
- 105 Y. Shao, L. Xu, L. Xu, X. Zhang, Z. Wang, G. Zhao, Z. Hu, J. Li and M. Ouyang, eTransportation, 2023, 18, 100285.
- 106 J. Zhang, Y. Tang, C. Song, X. Cheng, J. Zhang and H. Wang, Electrochim. Acta, 2007, 52, 5095-5101.
- 107 I. O. M. Kellegoz and M. S. Kilickaya, J. Optoelectron. Adv. Mater., 2008, 10, 369-372.
- 108 X. Li, F. Tang, Q. Wang, B. Li, H. Dai, G. Chang, C. Zhang, W. Zheng and P. Ming, Int. J. Hydrogen Energy, 2024, 50, 1228-1238.
- 109 M. Butori, B. Eriksson, N. Nikolić, C. Lagergren, G. Lindbergh and R. Wreland Lindström, Int. J. Hydrogen Energy, 2024, 95,
- 110 A. Jatukaran, J. Zhong, A. H. Persad, Y. Xu, F. Mostowfi and D. Sinton, ACS Appl. Nano Mater., 2018, 1, 1332-1338.
- 111 C. Yin, J. Li, Y. Zhou, H. Zhang, P. Fang and C. He, ACS Appl. Mater. Interfaces, 2018, 10, 14026-14035.
- 112 R. Sun, Z. Xia, C. Yang, F. Jing, S. Wang and G. Sun, Prog. Nat. Sci.: Mater. Int., 2020, 30, 912-917.
- 113 H. Eskandari, D. K. Paul, A. P. Young and K. Karan, ACS Appl. Mater. Interfaces, 2022, 14, 50762-50772.

- 114 S. Fujita and A. Suzuki, J. Appl. Phys., 2010, 107, 1.
- 115 M. Taş and G. Elden, Fuel Cells, 2020, 20, 531-539.
- 116 Y. Wang, Y. Pang, H. Xu, A. Martinez and K. S. Chen, Energy Environ. Sci., 2022, 15, 2288-2328.
- 117 J. Zhang, Y. Tang, C. Song, J. Zhang and H. Wang, J. Power Sources, 2006, 163, 532-537.
- 118 Y. Zhang, Z. Xiao, X. Zhao, J. Wang, Y. Wang and J. Yu, Membranes, 2025, 15, 72.
- 119 J.-C. Shyu, K.-L. Hsueh and F. Tsau, Energy Convers. Manage., 2011, 52, 3415-3424.
- 120 Q. Yan, H. Toghiani and H. Causey, J. Power Sources, 2006, 161, 492-502.
- 121 Q. Wang, F. Tang, B. Li, H. Dai, J. P. Zheng, C. Zhang and P. Ming, Chem. Eng. J., 2022, 433, 133667.
- 122 H. Chen, Y. Liu, C. Deng and J. Chen, Int. J. Hydrogen Energy, 2023, 48, 1075-1089.
- 123 M. W. H. Xu, Y. Liu, V. Mittal, R. Vieth, H. R. Kunz, L. J. Bonville and J. M. Fenton, ECS Trans., 2006, 3, 561–568.
- 124 C. D. G. Remy Sellin, S. Arrii-Clacens, S. Pronier, J.-M. Clacens, C. Coutanceau and J.-M. Leger, J. Phys. Chem. C, 2009, 113, 21735-21744.
- 125 Z. Yang, S. Ball, D. Condit and M. Gummalla, J. Electrochem. Soc., 2011, 158, B1439.
- 126 R. L. Borup, J. R. Davey, F. H. Garzon, D. L. Wood and M. A. Inbody, J. Power Sources, 2006, 163, 76-81.
- 127 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.
- 128 F. Hasché, M. Oezaslan and P. Strasser, ChemPhysChem, 2012, 13, 828-834.
- 129 D. A. Stevens and J. R. Dahn, Carbon, 2005, 43, 179-188.
- 130 R. Sellin, J.-M. Clacens and C. Coutanceau, Carbon, 2010, 48, 2244-2254.
- 131 J. Zhao, X. Huang, H. Chang, S. Hwa Chan and Z. Tu, Energy Convers. Manage.: X, 2021, 10, 100087.
- 132 C. Yue, W. Zheng, Y. Lian, J. Kang, S. Chen, X. Dong, B. Li, C. Zhang and P. Ming, J. Energy Storage, 2024, 104, 114426.
- 133 Y. Lu, D. Yang, H. Wu, L. Jia, J. Chen, P. Ming and X. Pan, Renewable Energy, 2024, 234, 121166.
- 134 S. Lee, J. Nam, J. Ahn, S. Yoon, S. C. Jeong, H. Ju and C. H. Lee, Int. J. Hydrogen Energy, 2024, 96, 333-342.
- 135 X. Gao, K. Yamamoto, T. Hirai, T. Uchiyama, N. Ohta, N. Takao, M. Matsumoto, H. Imai, S. Sugawara, K. Shinohara and Y. Uchimoto, Langmuir, 2020, 36, 3871-3878.
- 136 D. K. Paul and K. Karan, J. Phys. Chem. C, 2014, 118, 1828-1835.
- 137 Y. Liang, Y. Feng, H. Li, M. He, D. Yang, S. Shen, X. Yan, J. Zhang, X. Xu and R. Masrour, presented in part at the International Conference on Optoelectronic Information and Functional Materials (OIFM 2024), 2024.
- 138 R. Borup, J. Meyers, B. Pivovar, Y. S. Kim, R. Mukundan, N. Garland, D. Myers, M. Wilson, F. Garzon and D. Wood, Chem. Rev., 2007, 107, 3904-3951.
- 139 J. C. Meier, C. Galeano, I. Katsounaros, J. Witte, H. J. Bongard, A. A. Topalov, C. Baldizzone, S. Mezzavilla, F. Schüth and K. J. J. Mayrhofer, Beilstein J. Nanotechnol., 2014, 5, 44-67.

- 140 A. P. Soleymani, L. R. Parent and J. Jankovic, Adv. Funct. Mater., 2021, 32, 2105188.
- 141 Y. Shao, G. Yin and Y. Gao, *J. Power Sources*, 2007, **171**, 558–566.
- 142 N. Trogisch, D. Babik, A. Orfanidi, G. Jegert, A. Albert and R.-A. Eichel, *J. Electrochem. Soc.*, 2024, 171, 104506.
- 143 W. Bi, Q. Sun, Y. Deng and T. F. Fuller, *Electrochim. Acta*, 2009, **54**, 1826–1833.
- 144 G. Inzelt, B. Berkes and Á. Kriston, *Electrochim. Acta*, 2010, 55, 4742–4749.
- 145 Z. Zheng, F. Yang, C. Lin, F. Zhu, S. Shen, G. Wei and J. Zhang, ACS Appl. Mater. Interfaces, 2020, 12, 35088–35097.
- 146 T. F. F. Wu Bi, ECS Trans., 2007, 11, 1235.
- 147 Z. Huang, Z. Zhou, J. Zhao, W.-T. Wu, L. Wei, C. Hu, Y. Yang, Y. Li and Y. Song, *Int. J. Hydrogen Energy*, 2024, 53, 1107–1122.
- 148 T. Đukić, L. J. Moriau, L. Pavko, M. Kostelec, M. Prokop, F. Ruiz-Zepeda, M. Šala, G. Dražić, M. Gatalo and N. Hodnik, ACS Catal., 2021, 12, 101–115.
- 149 S. M. Andersen, M. Borghei, P. Lund, Y.-R. Elina, A. Pasanen, E. Kauppinen, V. Ruiz, P. Kauranen and E. M. Skou, *Solid State Ionics*, 2013, 231, 94–101.
- 150 S. Maass, F. Finsterwalder, G. Frank, R. Hartmann and C. Merten, *J. Power Sources*, 2008, **176**, 444–451.
- 151 V. E. Guterman, S. V. Belenov, V. V. Krikov, L. L. Vysochina, W. Yohannes, N. Y. Tabachkova and E. N. Balakshina, J. Phys. Chem. C, 2014, 118, 23835–23844.
- 152 K. H. Lim, H.-S. Oh, S.-E. Jang, Y.-J. Ko, H.-J. Kim and H. Kim, *J. Power Sources*, 2009, **193**, 575–579.
- 153 S. R. A. Olga, A. Baturina and K. J. Wynne, *Chem. Mater.*, 2006, **18**, 1498–1504.
- 154 C. H. P. L. M. Roen and T. D. Jarvi, *Electrochem. Solid State Lett.*, 2004, 7, A19–A22.
- 155 M. T. H. D. A. Stevens, G. M. Haugen and J. R. Dahn, J. Electrochem. Soc., 2005, 152, A2309–A2315.
- 156 M. A. Modestino, A. Kusoglu, A. Hexemer, A. Z. Weber and R. A. Segalman, *Macromolecules*, 2012, **45**, 4681–4688.
- 157 A. Kusoglu, D. Kushner, D. K. Paul, K. Karan, M. A. Hickner and A. Z. Weber, *Adv. Funct. Mater.*, 2014, 24, 4763–4774.
- 158 D. K. Paul, H. K. K. Shim, J. B. Giorgi and K. Karan, J. Polym. Sci., Part B: Polym. Phys., 2016, 54, 1267–1277.
- 159 A. Alentiev, J. Kostina and G. Bondarenko, *Desalination*, 2006, 200, 32–33.
- 160 D. Chu, D. Gervasio, M. Razaq and E. Yeager, J. Appl. Electrochem., 1990, 20, 157–162.
- 161 R. P. Dowd, Jr., Y. Li and T. Van Nguyen, *J. Appl. Electro-chem.*, 2020, **50**, 993–1006.
- 162 Y. Chang, J. Zhao, S. Shahgaldi, Y. Qin, Y. Yin and X. Li, *Int. J. Hydrogen Energy*, 2020, 45, 29904–29916.
- 163 F. Rong, C. Huang, Z.-S. Liu, D. Song and Q. Wang, *J. Power Sources*, 2008, **175**, 699–711.
- 164 Y. Tang, A. M. Karlsson, M. H. Santare, M. Gilbert, S. Cleghorn and W. B. Johnson, *Mater. Sci. Eng.*, A, 2006, 425, 297–304.
- 165 J. W. W. Vijay, A. Sethuraman, A. T. Haug, S. Motupally and L. V. Protsailo, *J. Electrochem. Soc.*, 2008, **155**, B50–B57.

- 166 C. Yue, W. Zheng, S. Chen, B. Li, C. Zhang, F. Tang and P. Ming, *Int. J. Hydrogen Energy*, 2024, **50**, 1515–1525.
- 167 S. Han, W. Zheng, Y. Lu, Y. Lian, B. Li, P. Ming and C. Zhang, *Int. J. Hydrogen Energy*, 2024, **58**, 279–288.
- 168 A. Kneer, J. Jankovic, D. Susac, A. Putz, N. Wagner, M. Sabharwal and M. Secanell, *J. Electrochem. Soc.*, 2018, **165**, F3241–F3250.
- 169 A. Malekian, S. Salari, J. Stumper, N. Djilali and M. Bahrami, *Int. J. Hydrogen Energy*, 2019, 44, 23396–23405.
- 170 S. Kim, J.-H. Kim, J.-G. Oh, K.-L. Jang, B.-H. Jeong,
 B. K. Hong and T.-S. Kim, ACS Appl. Mater. Interfaces,
 2016, 8, 15391–15398.
- 171 A. S. Alavijeh, R. M. H. Khorasany, Z. Nunn, A. Habisch, M. Lauritzen, E. Rogers, G. G. Wang and E. Kjeang, J. Electrochem. Soc., 2015, 162, F1461–F1469.
- 172 J.-B. Pyo, J. H. Kim and T.-S. Kim, *J. Mater. Chem. A*, 2020, **8**, 24763–24773.
- 173 S. Kundu, M. W. Fowler, L. C. Simon and S. Grot, *J. Power Sources*, 2006, **157**, 650–656.
- 174 J. Zhao, S. Shahgaldi, X. Li and Z. Liu, *J. Electrochem. Soc.*, 2018, **165**, F3337–F3345.
- 175 R. L. Borup, A. Kusoglu, K. C. Neyerlin, R. Mukundan, R. K. Ahluwalia, D. A. Cullen, K. L. More, A. Z. Weber and D. J. Myers, *Curr. Opin. Electrochem.*, 2020, 21, 192–200.
- 176 C. Li, K. Yu, A. Bird, F. Guo, J. Ilavsky, Y. Liu, D. A. Cullen, A. Kusoglu, A. Z. Weber, P. J. Ferreira and J. Xie, *Energy Environ. Sci.*, 2023, 16, 2977–2990.
- 177 Y. Zhang, X. Li and A. Klinkova, *Int. J. Hydrogen Energy*, 2021, **46**, 11071–11083.
- 178 F. Rong, C. Huang, Z.-S. Liu, D. Song and Q. Wang, *J. Power Sources*, 2008, **175**, 712–723.
- 179 J. Liu, Y. Yin, J. Zhang, T. Zhang, X. Zhang and H. Chen, *J. Power Sources*, 2021, 512, 230487.
- 180 Y. Chang, J. Liu, R. Li, J. Zhao, Y. Qin, J. Zhang, Y. Yin and X. Li, *Energy Convers. Manage.*, 2019, **189**, 24–32.
- 181 A. Stassi, I. Gatto, E. Passalacqua, V. Antonucci, A. S. Arico, L. Merlo, C. Oldani and E. Pagano, *J. Power Sources*, 2011, 196, 8925–8930.
- 182 Z. Yao, F. Zhou, C. Tu, J. Tan and M. Pan, *Int. J. Hydrogen Energy*, 2024, **50**, 200–208.
- 183 S. Chen, M. Hao, Y. Hu, K. Liu and Y. Li, *J. Power Sources*, 2024, **599**, 234238.
- 184 W. Zheng, L. Xu, Z. Hu, Y. Ding, J. Li and M. Ouyang, J. Power Sources, 2021, 487, 229367.
- 185 S.-H. Shin, P. J. Nur, A. Kodir, D.-H. Kwak, H. Lee, D. Shin and B. Bae, *ACS Omega*, 2019, 4, 19153–19163.
- 186 S. Giancola, R. A. B. Arciniegas, A. Fahs, J.-F. Chailan, M. L. Di Vona, P. Knauth and R. Narducci, *Membranes*, 2019, 9, 134.
- 187 S. H. Frensch, G. Serre, F. Fouda-Onana, H. C. Jensen, M. L. Christensen, S. S. Araya and S. K. Kær, *J. Power Sources*, 2019, 420, 54–62.
- 188 M. Feng, R. Qu, Z. Wei, L. Wang, P. Sun and Z. Wang, Sci. Rep., 2015, 5, 9859.
- 189 A. B. B. Paul, W. Majsztrik and J. B. Benziger, *Macromole-cules*, 2008, **41**, 9849–9862.

- 190 S. Shi, A. Z. Weber and A. Kusoglu, J. Membr. Sci., 2016, 516, 123-134.
- 191 H.-Y. Jung and J. W. Kim, Int. J. Hydrogen Energy, 2012, 37, 12580-12585.
- 192 S. W. Osung Kwon and D.-M. Zhu, J. Phys. Chem. B, 2010, **114**, 14989-14994.
- 193 B. R. Matos, M. A. Dresch, E. I. Santiago, L. P. R. Moraes, D. J. Carastan, J. Schoenmaker, I. A. Velasco-Davalos, A. Ruediger, A. C. Tavares and F. C. Fonseca, Electrochim. Acta, 2016, 196, 110-117.
- 194 C. A. Wilkie, J. R. Thomsen and M. L. Mittleman, J. Appl. Polym. Sci., 2003, 42, 901-909.
- 195 M. Pan, C. Pan, C. Li and J. Zhao, Renewable Sustainable Energy Rev., 2021, 141, 110771.
- 196 A. M. Dafalla, L. Wei, B. T. Habte, J. Guo and F. Jiang, Energies, 2022, 15, 9247.
- 197 M. P. Rodgers, L. J. Bonville, H. R. Kunz, D. K. Slattery and J. M. Fenton, Chem. Rev., 2012, 112, 6075-6103.
- 198 Y. K. Hamdy, F. M. Mohamed, C. S. Kuroda and A. Ohira, Macromol. Chem. Phys., 2011, 212, 708-714.
- 199 K. M. C. Kirt, A. Page and R. B. Moore, Macromolecules, 2005, 38, 6472-6484.
- 200 M. Marrony, D. Beretta, S. Ginocchio, Y. Nedellec, S. Subianto and D. J. Jones, Fuel Cells, 2013, 13, 1146-1154.
- 201 P. Choi, N. H. Jalani, T. M. Thampan and R. Datta, J. Polym. Sci., Part B: Polym. Phys., 2006, 44, 2183-2200.
- 202 J. Li, X. Yang, H. Tang and M. Pan, J. Membr. Sci., 2010, 361, 38-42.
- 203 J. E. Hensley, J. D. Way, S. F. Dec and K. D. Abney, J. Membr. Sci., 2007, 298, 190-201.
- 204 D. DeBonis, M. Mayer, A. Omosebi and R. S. Besser, Renewable Energy, 2016, 89, 200-206.
- 205 M. Casciola, G. Alberti, M. Sganappa and R. Narducci, J. Power Sources, 2006, 162, 141-145.
- 206 Y. Jeon, H.-k Hwang, J. Park, H. Hwang and Y.-G. Shul, Int. J. Hydrogen Energy, 2014, 39, 11690-11699.
- 207 N. Zhao, Y. Chu, Z. Xie, K. Eggen, F. Girard and Z. Shi, Fuel Cells, 2020, 20, 176-184.
- 208 D. Qiu, L. Peng, X. Lai, M. Ni and W. Lehnert, Renewable Sustainable Energy Rev., 2019, 113, 109289.
- 209 M. N. Silberstein and M. C. Boyce, J. Power Sources, 2011, **196**, 3452-3460.
- 210 A. Kusoglu, M. H. Santare and A. M. Karlsson, J. Polym. Sci., Part B: Polym. Phys., 2011, 49, 1506-1517.
- 211 R. Mukundan, A. M. Baker, A. Kusoglu, P. Beattie, S. Knights, A. Z. Weber and R. L. Borup, J. Electrochem. Soc., 2018, 165, F3085-F3093.
- 212 Y. Singh, F. P. Orfino, M. Dutta and E. Kjeang, J. Electrochem. Soc., 2017, 164, F1331-F1341.
- 213 S. Shi, X. Sun, Q. Lin, J. Chen, Y. Fu, X. Hong, C. Li, X. Guo, G. Chen and X. Chen, Int. J. Hydrogen Energy, 2020, 45, 27653-27664.
- 214 S. Bhattacharya, J. Leung, M. V. Lauritzen and E. Kjeang, Electrochim. Acta, 2020, 352, 136489.
- 215 Y. Chen, C. Jiang and C. Cho, Polymers, 2018, 10, 971.

- 216 F. Qu, L. Liu, G. Tao, W. Zhan, S. Zhan, Y. Li, C. Li, X. Lv, Z. Shi and H. Duan, Tribol. Int., 2023, 188, 108850.
- 217 Q. Meyer, Y. Zeng and C. Zhao, Adv. Mater., 2019, 31, 1901900.
- 218 P. Irmscher, D. Qui, H. Janßen, W. Lehnert and D. Stolten, Int. J. Hydrogen Energy, 2019, 44, 23406-23415.
- 219 F. Lapicque, M. Belhadj, C. Bonnet, J. Pauchet and Y. Thomas, J. Power Sources, 2016, 336, 40-53.
- 220 R. Taherian, M. Matboo Ghorbani and S. R. Kiahosseini, J. Electroanal. Chem., 2018, 815, 90-97.
- 221 A. Kausar, I. Rafique and B. Muhammad, Polym.-Plast. Technol. Eng., 2016, 55, 1167-1191.
- 222 M. C. Gutiérrez, M.-A. De Paoli and M. I. Felisberti, Ind. Crop. Prod., 2014, 52, 363-372.
- 223 A. P. G. Agarwal and R. K. Sharma, J. Eng. Sci. Technol., 2014, 9, 590-604.
- 224 J. Zhou, Z. Yao, Y. Chen, D. Wei, Y. Wu and T. Xu, Polym. Compos., 2013, 34, 1245-1249.
- 225 P. K. Z. Zhang and K. Friedrich, Compos. Sci. Technol., 2002, 62, 1001-1009.
- 226 Y. Lian, W. Zheng, C. Yue, S. Han and P. Ming, Int. J. Heat Mass Transfer, 2024, 223, 125239.
- 227 Y. Jeon, H. Na, H. Hwang, J. Park, H. Hwang and Y.-G. Shul, Int. J. Hydrogen Energy, 2015, 40, 3057-3067.
- 228 H. Lee, S. Jeong, J. Song, M. Kim, C. Chu, Y. Lee, D. Kim and M. Kim, J. Mech. Sci. Technol., 2023, 37, 2095-2108.
- 229 M. Marrony, R. Barrera, S. Quenet, S. Ginocchio, L. Montelatici and A. Aslanides, J. Power Sources, 2008, 182, 469-475.
- 230 P. Ren, P. Pei, Y. Li, Z. Wu, D. Chen and S. Huang, Prog. Energy Combust. Sci., 2020, 80, 100859.
- 231 L. V. Nam, E. Choi, S. Jang and S. M. Kim, Renewable Energy, 2021, 180, 203-212.
- 232 J.-S. Park, M.-S. Shin and C.-S. Kim, Curr. Opin. Electrochem., 2017, 5, 43-55.
- 233 G. Xu, A. Ke, G. Xu, Y. Liu, Y. Zuo, X. Yang, Y. Dong, J. Wang, J. Zheng, J. Li and W. Cai, Int. J. Hydrogen Energy, 2024, 56, 330-337.
- 234 Y. Liao, S. Zhao, G. Liu, H. Li, J. Shuai, L. Wang, B. Liu and H. Tang, Chem. Eng. J., 2024, 488, 150971.
- 235 G. Gnana Kumar, A. R. Kim, K. Suk Nahm and R. Elizabeth, Int. J. Hydrogen Energy, 2009, 34, 9788-9794.
- 236 J. Niu, S. Zhang, Y. Li, X. Li, J. Zhang, S. Lu and Q. He, Polymer, 2023, 273, 125869.
- 237 S. Vengatesan, H. Kim, S. Lee, E. Cho, H. Yongha, I. Oh, S. Hong and T. Lim, Int. J. Hydrogen Energy, 2008, 33, 171-178.
- 238 A. A. Wani, N. Shaari, S. K. Kamarudin, N. F. Raduwan, Y. N. Yusoff, A. M. Khan, S. Yousuf and A. Mohamed Nainar Mohamed, Energy Fuels, 2024, 38, 18169-18193.
- 239 I. Nicotera, L. Coppola and C. Simari, Mater. Renewable Sustainable Energy, 2024, 13, 307-318.
- 240 Y. Ji, Y. i Cho, Y. Jeon, C. Lee, D.-H. Park and Y.-G. Shul, Appl. Catal., B, 2017, 204, 421-429.
- 241 H. X. Jianwei Yang, J. Li, K. Gong, F. Yue, X. Han, K. Wu, P. Shao, Q. Fu, Y. Zhu, W. Xu, X. Huang, J. Xie, F. Wang,

- W. Yang, T. Zhang, Z. Xu, X. Feng and B. Wang, Science, 2024, 385, 1115-1120.
- 242 Y. Z. Panpan Guan, M. Zhang, W. Zhong, J. Xu, J. Lei, H. Ding, W. Feng, F. Liu and Y. Zhang, Sci. Adv., 2023, 9, eadh1386.
- 243 S. H. Mirfarsi, A. Kumar, J. Jeong, M. Adamski, S. McDermid, B. Britton and E. Kjeang, Int. J. Hydrogen Energy, 2024, 50, 1507-1522.
- 244 C. Li, W. Wang, S. Zhu, H. Pan, Q. Xu, P. Shi and Y. Min, Chem. Eng. J., 2023, 466, 143105.
- 245 W. Zhang, Y. Chen, Y. Jin, H. Liu, Q. Ma, Q. Xu and H. Su, Int. J. Hydrogen Energy, 2024, 65, 829-836.
- 246 M. Janssen, J. Drnec, I. Martens, J. Quinson, R. Pittkowski, D. Park, P. Weber, M. Arenz and M. Oezaslan, Chem-SusChem, 2024, 17, e202400303.
- 247 K. Yu, J. L. Hart, J. Xie, M. L. Taheri and P. Ferreira, Nano Energy, 2023, 111, 108393.
- 248 F. Chabot, P. Lionel, L. Guétaz, S. Rosini and A. Morin, J. Electrochem. Soc., 2024, 171, 124506.
- 249 V. Shokhen, L. Strandberg, M. Skoglundh and B. Wickman, J. Mater. Chem. A, 2023, 11, 21029-21035.
- 250 C. Wang, J. Li, S. Zhang, X. Li, X. Duan, Y. Wu, Q. Zhang, T. Yang and J. Liu, Appl. Energy, 2024, 367, 123377.
- 251 Y. Sun, L. Mao, Z. Hu, X. Zhang and R. Peng, Adv. Sci., 2024, **11**, 2403631.
- 252 M. Bogar, Y. Yakovlev, J. Nováková, A. M. Darabut, M. Kriechbaum, H. Amenitsch, R. Taccani and I. Matolínová, Int. J. Hydrogen Energy, 2024, 58, 1673-1681.
- 253 J. Chen, M. Perez-Page, C. M. A. Parlett, Z. Guo, X. Yang, Z. Zhou, H. Zhai, S. Bartlett, T. S. Miller and S. M. Holmes, Chem. Eng. J., 2024, 487, 150670.

- 254 Q. Li, C. J. Pollock, J. Soto, A. M. Villarino, Z. Shi, M. R. Krumov, D. A. Muller and H. D. Abruña, Nat. Commun., 2025, 16, 3008.
- 255 A. P. Soleymani, M. Reid and J. Jankovic, Adv. Funct. Mater., 2022, 33, 2209733.
- 256 Y. Ao, Z. Li, S. Laghrouche, D. Depernet, D. Candusso and K. Zhao, J. Power Sources, 2024, 603, 234420.
- 257 C. Liu, Z. Geng, X. Wang, W. Liu, Y. Wang, Q. Xia, W. Li, L. Jin and C. Zhang, J. Energy Chem., 2024, 90, 348-369.
- 258 I. Gatto, A. Caprì, C. Lo Vecchio, S. Zignani, A. Patti and V. Baglio, Int. J. Hydrogen Energy, 2023, 48, 11914-11921.
- 259 M. Rossini, D. Pan, B. Koyutürk, S. Chen, A. Khataee, G. Lindbergh, P. Jannasch and A. Cornell, J. Mater. Chem. A, 2024, **12**, 12826–12834.
- 260 J. E. Park, S. Y. Kang, S.-H. Oh, J. K. Kim, M. S. Lim, C.-Y. Ahn, Y.-H. Cho and Y.-E. Sung, Electrochim. Acta, 2019, 295, 99-106.
- 261 A. Lim, H.-J. Kim, D. Henkensmeier, S. Jong Yoo, J. Young Kim, S. Young Lee, Y.-E. Sung, J. H. Jang and H. S. Park, J. Ind. Eng. Chem., 2019, 76, 410-418.
- 262 S. Y. Kang, J. E. Park, G. Y. Jang, O.-H. Kim, O. J. Kwon, Y.-H. Cho and Y.-E. Sung, Int. J. Hydrogen Energy, 2022, 47, 9115-9126.
- 263 Y. Leng, G. Chen, A. J. Mendoza, T. B. Tighe, M. A. Hickner and C.-Y. Wang, J. Am. Chem. Soc., 2012, 134, 9054-9057.
- 264 L. Xia, W. Jiang, H. Hartmann, J. Mayer, W. Lehnert and M. Shviro, ACS Appl. Mater. Interfaces, 2022, 14, 19397-19408.
- 265 X. Wang, Z. Jiang, Y. Ma, X. Su, X. Zhao, A. Zhu and Q. Zhang, J. Power Sources, 2024, 591, 233819.
- 266 C. X. Lin, X. Q. Wang, L. Li, F. H. Liu, Q. G. Zhang, A. M. Zhu and Q. L. Liu, J. Power Sources, 2017, 365, 282-292.
- 267 L. Yuan, Q. Wang, F. Tang, B. Li, P. Ming and C. Zhang, Etransportation, 2022, 13, 100178.