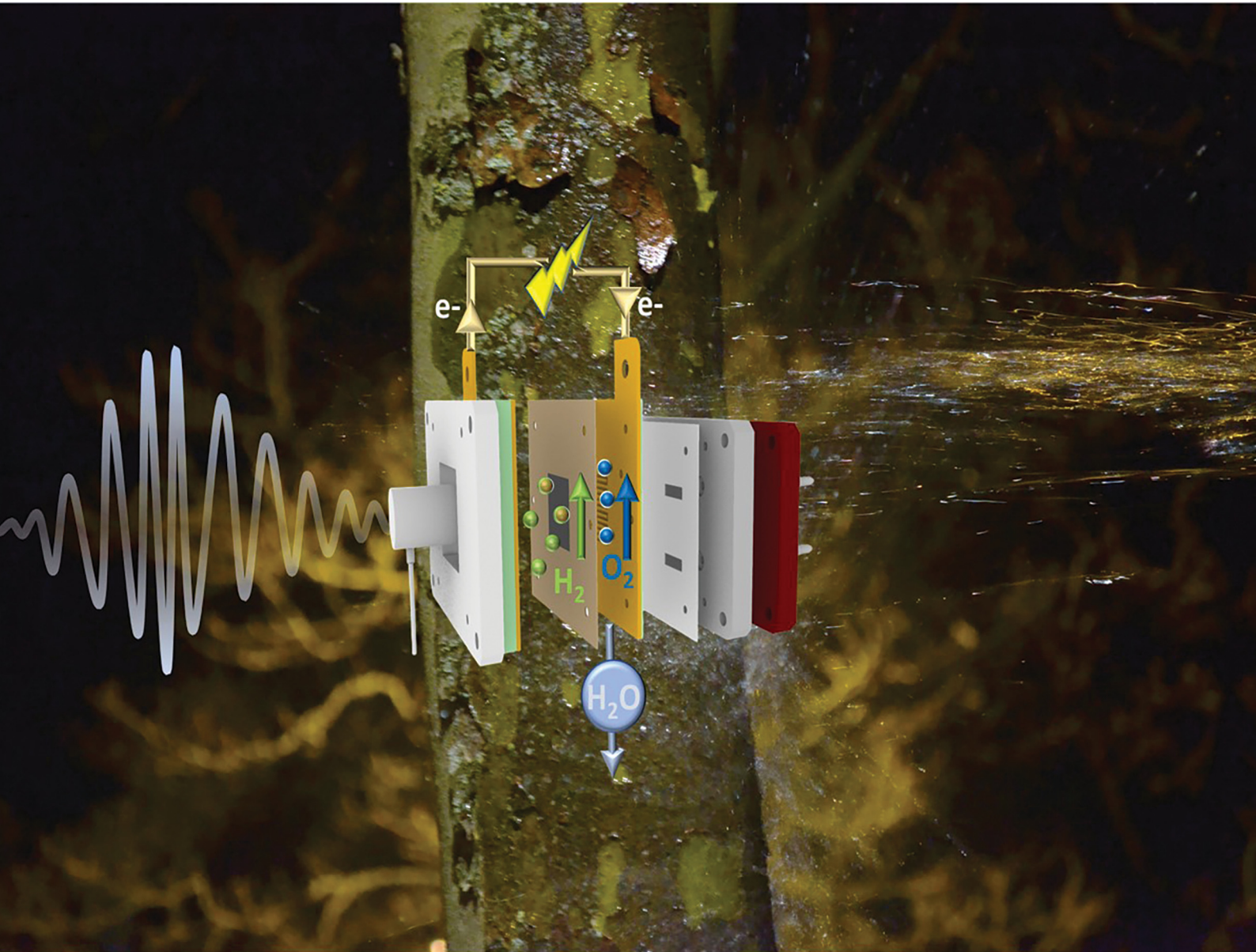


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Dynamic acoustic emission analysis of polymer electrolyte membrane fuel cells[†]

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The acoustic emission (AE) technique has been demonstrated as a non-invasive and non-destructive water management diagnostic tool for polymer electrolyte membrane fuel cells (PEMFCs). AE probes the dynamics of water generation and removal at the flow-field of a PEMFC to establish the hydration state inside the cell and has been utilised to electro-acoustically characterise the performance of a PEMFC under different operating conditions. In this study, the dynamic relationship between the acoustic activity and the rate of electrochemical reaction inside a PEMFC is explored by correlating AE from PEMFCs with their performance using different time-based characterisations (polarisation scans at 10 s, 60 s, and 120 s voltage stabilisation durations). Flooding resulted in ~16% decrease in maximum current density generated at 60 s and 120 s conditions compared to that at 10 s. Besides, flooding at longer durations is confirmed by acoustic emission as a function of polarisation (AEfP) and electrochemical impedance spectroscopy measurements. The effectiveness of the AE technique as a direct water diagnostic tool for PEMFCs is established through forward-reverse polarisation scans. Here, the AE energy generated during cell polarisations is utilised in understanding the water uptake and release mechanism inside the fuel cell. Furthermore, cell durability testing is performed through galvanostatic and potentiostatic measurements, where a synchronous relationship between the cell performance and the measured AE is identified.

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1. Introduction

Polymer electrolyte membrane fuel cells (PEMFCs) are devices that generate electrical energy from the electrochemical reaction between hydrogen fuel and air or oxygen. In addition to the electrical energy, the electrochemical reaction inside the fuel cell produces heat and water as by-products. The rate of electrochemical reaction in the cell defines the quantity of heat and amount of water generated – these, in turn, determine the cell temperature and the hydration conditions, respectively. Furthermore, temperature and hydration conditions influence the reaction rate, ionic conductivity, and the associated cell performance significantly.^{1,2}

Efficient water management inside a PEMFC is critical for the cell to deliver better operating performance. Improper water

regulation, resulting in either 'dry' or 'flooded' conditions, can deteriorate the performance of the cell considerably. Several methods are utilised for indirect and direct evaluation of water distribution inside a PEMFC.^{3,4} For instance, measurement techniques based on cell resistance,^{5,6} cell voltage,^{7,8} and pressure drop^{9,10} have been adopted to provide a qualitative understanding of the water dynamics inside the cell. To establish a quantitative understanding, direct water evaluation methods based on visualisation techniques like optical imaging,^{11–13} neutron imaging,^{14–16} X-ray,^{17–20} and magnetic resonance imaging^{21–23} have been used. However, these methods have certain operational and utilisation limitations, like material or component compatibility issues,¹³ measurement expense, and limited availability.^{24,25}

The acoustic activity of a system can be measured using the acoustic emission (AE) technique. An AE event results from an elastic wave generated within a material or component as a result of a mechanical perturbation. Easy accessibility of this technique allows a wide range of applications to utilise AE for monitoring, diagnostic, analytical, and other scientific studies in fields as diverse as corrosion science,²⁶ medical physics,²⁷ automotive industry,²⁸ and building and construction materials.²⁹ In the field of electrochemical science and engineering, AE has been utilised for diagnostic purposes in several

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applications as well, and a detailed study on this is given in previous work.³⁰ So far, AE has been utilised in fuel cell technology for fault analysis and diagnosis,³¹ thermo-mechanical characterisation of solid oxide fuel cells (SOFCs),³² damage evaluation of SOFCs,^{33,34} monitoring structural properties during drying and swelling of Nafion membrane,³⁵ and diagnosis of physiochemical changes occurring inside a PEMFC under a range of operating conditions.³⁶

Acoustic activity generated within a PEMFC can be utilised for the direct water evaluation inside the cell, irrespective of its material properties. Recently, Bethapudi *et al.*^{30,37} developed an acoustic emission as a function of polarisation (AEfP) technique for hydration diagnostics of PEMFCs. AEfP probes the water dynamics inside a PEMFC by analysing the AE energy released from the cell at discrete points of polarisation. The measured AE energy is instrumental in establishing correlations between the level of cell polarisation and the water generation and removal within flow-fields. Furthermore, the AEfP technique has successfully diagnosed flooding in fractal flow-field PEMFCs.³⁸

In this study, the AEfP technique is utilised to study the dynamic relationship between PEMFC performance and its corresponding acoustic activity under a time-dependent parametrisation. In addition, the cells are acoustically tested through consecutive forward and reverse polarisation scans to explore the electro-acoustic dynamic relationship. Galvanostatic and potentiostatic tests are performed to highlight the synchronous behaviour between the cell performances affected by hydration conditions and the generated acoustic activity. Furthermore, cell temperature profiles are utilised to better understand the correlation between electrochemical reactions or other processes within the fuel cell and acoustic activity.

2. Experimental setup and methods

2.1. PEMFC flow-field plates

The PEMFC used in this study had single-serpentine cathode and anode flow-fields. The cathode flow-field was developed from planar printed circuit board (PCB) plates using a layer-wise assembly approach.³⁹ Each flow-field was assembled using layers

of PCB plates, an approach that has been identified as cost-effective, easy and scalable for manufacturing.^{39–41} Each plate had dimensions of 80 mm × 80 mm × 2 mm. One plate accommodated the flow-field features, while the other plate acted as the backing plate. The PCB plate accommodating the flow-field features had a 35 µm thick Cu coating on its surface. The corresponding flow-field features were milled and drilled using a Roland – 40 Computer Numeric Control (CNC) machine (Roland, USA).

The plate with flow-field features was Ni electroplated using 0.13 M $\text{Ni}(\text{SO}_3\text{NH}_2)_2$ solution at 4.3 mA cm^{−2} current density (voltage range 3.0–3.5 V) for 3 min. After this, the Ni-coated plate was Au electroplated using 0.02 M $\text{KAu}(\text{CN})_2$ solution at a current density of 2.4 mA cm^{−2} (voltage range 3.5–3.7 V) for 60 min. The flow-field backing plate was a plain PCB plate with no metallic coating. The above two PCB plates, along with a pre-impregnated thermosetting polymer as an adhesive between them, were hot-pressed, as shown in Fig. 1, under a pressure of 400 psig at 150 °C for 60 min, followed by a 120 min ambient cooling period.^{30,39}

Fig. 1 shows the final assembled cathode flow-field plate, which has a 1 mm² square channel single-serpentine flow-field path with a depth of 1 mm covering an active membrane electrode assembly (MEA) area of 25 cm². Vertical surface air outlet paths allow for the removal of excess cathode reactant gases and product water. The overall dimensions of the cathode flow-field plate after the hot press were 80 mm × 80 mm × 4 mm.

The anode single-serpentine flow-field was developed from a graphite plate of 80 mm × 80 mm × 8 mm in dimensions (Fig. S1, ESI†). The anode serpentine flow-field features had similar channel dimensions and depth to those of the cathode flow-field but with a closed hydrogen removal path.

2.2. Membrane electrode assembly (MEA)

The MEA utilised in this study had an active area of 25 cm². The membrane used was Nafion[®] 212 (DuPont, USA) and was ~50 µm in thickness, the gas diffusion layer (GDL) was a carbon fibre paper with PTFE treated microporous layer (MPL) (Freudenberg H23C9) and was ~210 µm in thickness, and the catalyst loading was 0.4 mg_{pt} cm^{−2} HyPlat Pt catalyst (HyPlat, South Africa).

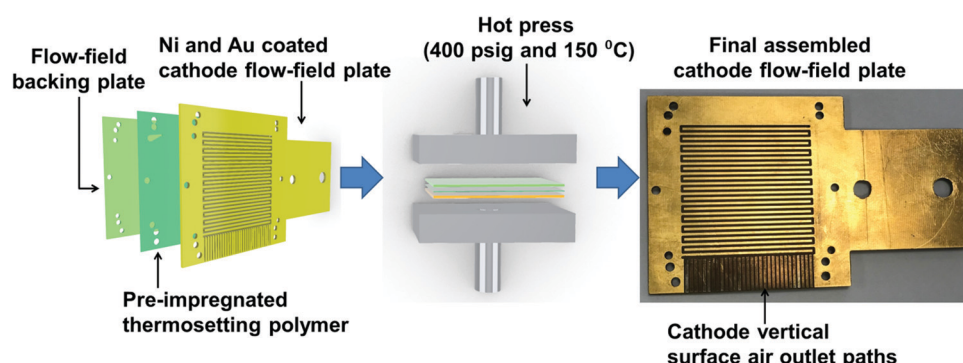


Fig. 1 Schematic outline of layer-wise PCB based PEMFC assembly.



These MEA components were hot press assembled for 3 min under a pressure of 400 psig and at a temperature of 150 °C. The overall thickness of the final assembled MEA was ~500 µm.

2.3. Fuel cell testing

PEMFC experimental testing was performed using a Scribner 850e fuel cell test station (Scribner Associates, NC, USA). The test station could supply reactants under controlled conditions of temperature, relative humidity, pressure, flow rates, and gas mixtures. In this study, high purity (99.995%) hydrogen at a flow rate of 200 cm³ min⁻¹ and bottled air at a flow rate of 1000 cm³ min⁻¹, under ambient conditions, were supplied to the PEMFC anode and cathode, respectively. Constant flow rates avoid acoustic disturbances generated from stoichiometric based flows.³⁰

The PEMFC was preheated to a temperature of 45 °C using cylindrical heaters; the corresponding cell temperatures generated were measured using a K-type thermocouple. The cell was operated under ambient cooling conditions.

A polarisation curve, potentiostatic and galvanostatic measurements were carried out as detailed in Tables 1 and 2, respectively. Before each experimental procedure, cell conditioning was performed by supplying the reactants under open circuit voltage (OCV) conditions for 2 min, which allowed the gas flow and the reactant humidity to stabilise.

2.4. Acoustic emission analysis

A schematic outline of the AE mechanism occurring inside the PEMFC is shown in Fig. 2. In this study, AE from the PEMFC was recorded using a piezoelectric transducer type acoustic sensor, as shown in Fig. 2. The acoustic sensor was externally

Table 1 Polarisation curve measurement parameters

Experimental condition	Voltage range	Hold time per voltage point	Voltage interval per point
Polarisation	OCV – 0.3 V	30 s	0.05 V
		60 s	
		120 s	
Forward and reverse polarisation	OCV – 0.3 V	60 s	0.05 V
	0.3 V – OCV	60 s	

Table 2 Potentiostatic and galvanostatic test parameters. Arrow indicates the sequence in which the experimental conditions are carried out

Experimental condition	Voltage/ Current	Hold duration	Duration per voltage acquisition point
Potentiostatic	0.8 V	900 s	10 s
	OCV	300 s	
	0.6 V	900 s	
	OCV	300 s	
	0.4 V	900 s	
	OCV	300 s	
Galvanostatic	20 A	600 s	
	OCV	300 s	
	25 A	600 s	
	OCV	300 s	
	30 A	600 s	
	OCV	300 s	

coupled to the cathode flow-field plate using silicone paste. The sensor was connected *via* a USB cable to a computer running AEwin™ data acquisition and replay software (Physical Acoustics Corporation, USA). The software allows for data logging, interpretation, and reproduction of the measured acoustic data. A detailed outline of the signal filters and pre-amplification devices used in the data acquisition is given in a previous study.^{30,38}

The acoustic sensor used was a single-channel acquisition system with an operating frequency of 200–1000 kHz and a resonant frequency of 500 kHz, respectively. The acquisition system had lower and upper analogue filters of 30 kHz and 1 MHz, respectively. Here, a threshold amplitude of 30 dB was considered based on the background noise, including the reactant flow through the channels, developed during the PEMFC testing. Signals with a magnitude beyond this threshold are considered as AE hits. The critical data points measured from the AE hits were threshold, peak amplitude (dB), counts, and absolute energy (aJ), as detailed in previous work.³⁰

2.5. Electrochemical impedance spectroscopy

Electrochemical impedance spectroscopy (EIS) measurements were performed using a Scribner 850e impedance analyser (Scribner Associates, NC, USA). The frequency range for analysis was 10 kHz to 0.1 Hz, with 10 points per decade, and the AC modulation amplitude was 5% of the DC input signal.

3. Results and discussion

3.1. Polarisation stabilisation based AE analysis

Fig. 3 shows the influence of the polarisation voltage stabilisation time on the performance of a PEMFC and its associated AE generated under three different stabilisation conditions – hold times of 10 s, 60 s and 120 s.

It can be observed from Fig. 3(a) that, with increased current density and especially between OCV and 0.6 V (Ohmic region of operation), the performance of the cell remained similar, irrespective of the stabilisation conditions. However, decreasing the voltage further resulted in a more significant decrease of the cell performance for 60 s and 120 s hold times compared to 10 s. The longer the voltage hold, the longer the electrochemical reaction will occur, with associated water generation.⁴² Thus, the poorer performance at lower cell voltages for longer hold times of 60 s and 120 s can be attributed to mass transport limitations that can be induced by flooding occurring inside the cell, as a result of excess water generation and retention with time.^{43,44} Furthermore, the overall cell polarisation for the 60 s and 120 s hold times remained similar, as seen in Fig. 3(a), indicating that the impact of the level of cell flooding on the cell performance did not become worse after 60 s.

Cell flooding can be identified from the corresponding AEfP curves, which reflect the AE energy measured from the impacts of water generation and release into the flow-fields, as shown in Fig. 3(b). It can be observed from Fig. 3(b) that the cumulative absolute AE energy (CAEE) measured from the cells increased



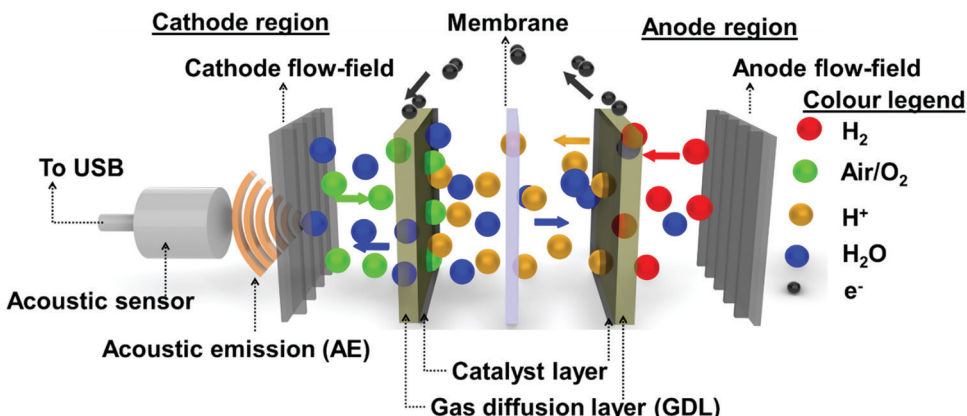


Fig. 2 Schematic outline of acoustic emission (AE) generation mechanism inside a PEMFC.

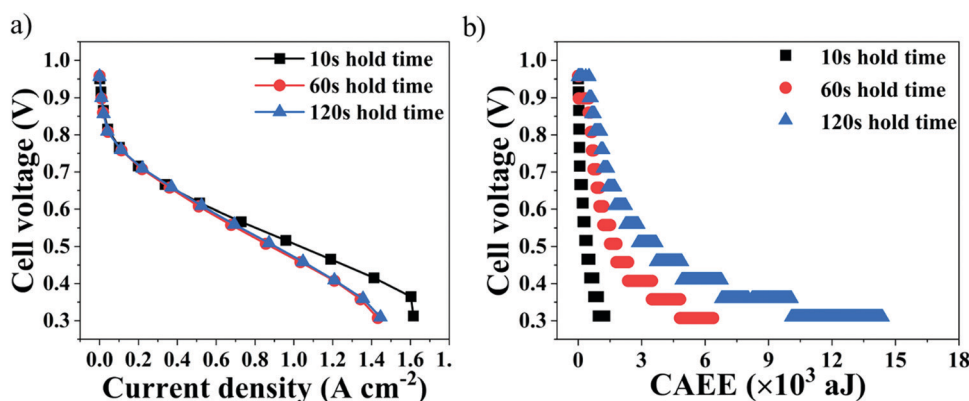


Fig. 3 (a) Polarisation curves and (b) simultaneous acoustic emission as a function of polarisation (AEfP) curves: for voltage stabilisation durations (hold time) of 10 s, 60 s, and 120 s. Note: Sampling period for measuring the CAEE and the voltage is same.

as the cell voltage decreased, which can be attributed to the increased extent of the electrochemical reactions and associated water generation inside the cell.^{30,36,45}

The overall CAEE measured was lowest at 10 s hold time, namely ~ 1500 aJ, which also delivered superior performance compared to the 60 s and 120 s hold times. However, the overall CAEE generated for the 60 s and 120 s hold times was ~ 6500 aJ and ~ 14500 aJ, respectively, and such very high acoustic activity can be attributed to excess water accumulation with time occurring inside the cell. Furthermore, the CAEE measured at each voltage point on the polarisation curves increased considerably when moving from a hold time of 60 s to 120 s. For instance, at 0.4 V, the corresponding CAEE was ~ 1150 aJ and ~ 1900 aJ, respectively.^{30,44}

3.2. Electrochemical impedance spectroscopy (EIS) measurements

EIS measurements on the same PEMFC were performed to characterise cell performance in the high current density operating region. Here, EIS was performed at 0.8 A cm^{-2} (20 A), 1 A cm^{-2} (25 A) and 1.2 A cm^{-2} (30 A). The corresponding EIS spectra and resistances developed in the PEMFC

are given in Fig. 4 and Table 3, respectively (equivalent circuit use for modelling the EIS data is provided in Fig. S2, ESI†).

It can be observed from Table 3 that with an increase in the current density from 0.8 A cm^{-2} to 1 A cm^{-2} , R_{mt} increases significantly with current density, while the high-frequency resistance (HFR) and R_{ct} are relatively unchanged. For this PEMFC, where the R_{mt} increases in the high current density region, accompanied by stable HFR and R_{ct} , any increase in the voltage hold duration will result in excess accumulation of water with time and exacerbates the mass transfer related issues like flooding, as described by the polarisation curves and the AEfP results shown in Fig. 3.⁴⁶

3.3. Forward and reverse polarisations

To evaluate the dynamic relationship between electrochemical activity and acoustic activity generated in a PEMFC, forward and reverse polarisation scans were performed, as shown in Fig. 5(a), at 60 s hold time. The corresponding acoustic activity during the scans is shown in Fig. 5(b).

The forward scan was run between OCV and 0.3 V, while the reverse scan was run between 0.3 V and OCV. The HFR during the forward scan, as in Fig. 5(c), initially decreases with an



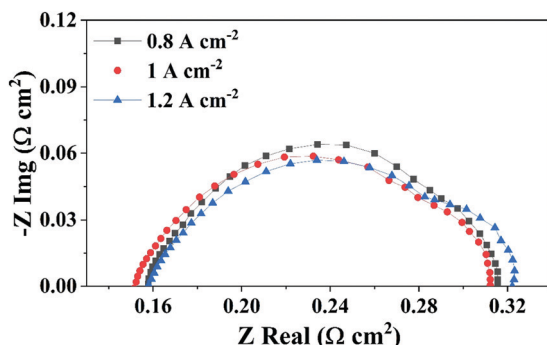


Fig. 4 Electrochemical impedance spectroscopy measurements at 20 A (0.8 A cm^{-2}), 25 A (1 A cm^{-2}) and 30 A (1.2 A cm^{-2}) current conditions, respectively.

Table 3 Resistances in the PEMFC derived from EIS measurements at current densities of 0.8 A cm^{-2} , 1 A cm^{-2} and 1.2 A cm^{-2} : high-frequency resistance – HFR; charge transfer resistance – R_{ct} ; mass transfer resistance – R_{mt}

Load	Resistance $\Omega \text{ cm}^2$		
	HFR	R_{ct}	R_{mt}
20 A/ 0.8 A cm^{-2}	0.158	0.285	0.025
20 A/ 1 A cm^{-2}	0.154	0.283	0.033
20 A/ 1.2 A cm^{-2}	0.158	0.281	0.040

increase in the cell operating current. This can be due to the improved membrane hydration, as a result of increased electrochemical reaction rates generating water, and its associated conductivity. However, beyond 0.45 V cell voltage the HFR increases; this is attributed to the increasing cell temperature at higher current, leading to membrane dehydration.^{15,47} The opposite can be attributed to the HFR measured during the reverse scan. At 0.3 V, the initial higher currents may have resulted in higher cell temperatures, resulting in some membrane dehydration. However, decreasing the cell current led to a lower cell temperature that resulted in increasing HFR until 0.65 V. Below 0.65 V, an increasing HFR is observed as a result of insufficient hydration occurring from lower cell currents.

It can also be observed from Fig. 5(a) that the polarisation performance during forward and reverse scans was similar, with only marginal differences observed in the high current density region. Such similarity in forward and reverse polarisation curves indicates optimal stabilisation conditions for the PEMFC to reach steady-state operation at each voltage point and avoid a hysteresis effect.⁴⁸ Furthermore, the similarity in polarisation performance was established by the near-identical HFR values generated, which establish the level of membrane hydration and its associated conductivity, as shown in Fig. 5(c).⁴⁹

The corresponding CAEE measured during the polarisation curves is shown in Fig. 5(b). It can be observed that, for both forward and reverse scans, similar acoustic activity was measured between OCV and 0.6 V, which establishes the measured AE as a dynamic response to the level of polarisation (electrochemical reaction) in the cell. Between 0.6 V and 0.3 V,

considerable acoustic activity was measured from the PEMFC, because of the increased water build-up and ineffective removal at higher current densities.

It can be observed from Fig. 5(b) that the net CAEE measured between 0.6 V and 0.3 V for the reverse scan was $\sim 18\text{--}25\%$ less than that measured for the forward scan. During a reverse scan, the initially high currents require a greater uptake of water by the membrane to maintain sufficient hydration and its associated conductivity. With progressive decreases in current density, the level of water in the GDL gradually reduces and this promotes re-absorption of the existing water in the cell into the GDL.^{50,51} This additional utilisation of water generated in the cell by the GDL during the reverse scan may have resulted in comparatively lower levels of water being released into the flow-field and associated AE measured compared to the forward scan, where the continuous electrochemical activity resulted in saturation of water in the GDL. Furthermore, the additional utilisation of water during the reverse scan can be established from the HFR, as given in Fig. 5(c).⁵² HFR is an index for the level of water uptake by a membrane and its associated conductivity.⁴⁹ The HFR initially decreases from $\sim 270 \Omega \text{ cm}^2$ at 0.85 V to $\sim 150 \Omega \text{ cm}^2$ at 0.55 V due to an increase in the hydration with increasing current. Later, HFR increases from $\sim 150 \Omega \text{ cm}^2$ at 0.55 V to $\sim 175 \Omega \text{ cm}^2$ at 0.3 V as a result of membrane dehydration due to increased temperatures at higher currents, which is a known trend from our previous work.^{39,41} The observed similar hydration conditions (HFR values) for both scans imply that during the reverse scan the utilisation of generated water may have offset the additional hydration requirements, resulting in less water released into the flow-field.

Overall, there is a natural hysteresis associated with the time it takes for liquid water to be cleared from the system, *i.e.* increasing cell currents constantly increase the amount of water in the system, leading to a build-up of water, especially in the forward scan, while in the reverse scan increasingly less water is generated, so build-up decreases. This explains the observed higher and lower CAEE observed in the forward and backward scans, respectively (assuming that the rate of water removal is constant).

3.4. Potentiostatic tests

Potentiostatic tests (voltage hold) were performed to probe the dynamics of current density development on the acoustic energy released inside the cell. The AE response to dynamic electrochemical conditions inside the PEMFC were studied under potentiostatic conditions, as shown in Fig. 6. Potentiostatic tests were carried out under three regions of PEMFC operation, namely, ‘activation’ (0.8 V), ‘Ohmic’ (0.6 V) and ‘concentration polarisation’ (0.4 V) regions. Under each voltage condition, the cell was held for 900 s at a particular voltage (gradient profile) followed by an OCV hold for 300 s (flat profile).

It can be observed from Fig. 6(b) that, at 0.8 V hold, the current density decreased from 0.035 A cm^{-2} to 0.028 A cm^{-2} with periodic ‘spikes’ observed in the current density profile,



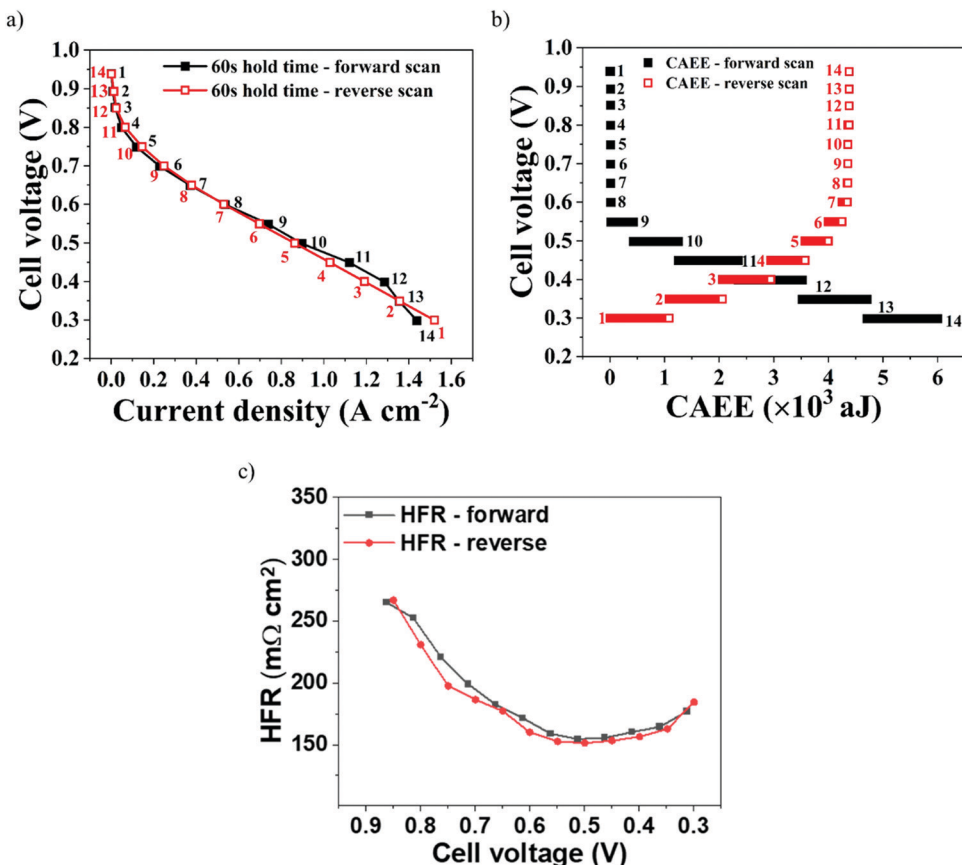


Fig. 5 (a) Forward (black solid square) and reverse (red open square) polarisation curves and (b) CAEE measured during forward and reverse polarisation scans, and (c) HFR for forward and reverse polarisation scans in potentiostatic mode at 60 s voltage hold conditions.

which are commonly referred to as 'flushing events' in a PEMFC.³⁰ Flushing events occur when the water in a flow-field builds up continuously, reducing the apparent cell performance, and eventually resulting in the sudden ejection of water through the flow-field out of the cell followed by a spontaneous increase in the cell performance.⁵¹ Here, such spurts of water inside the cell resulted in correlated AE responses, as presented in Fig. 6(b-d).^{30,36} It can be observed from Fig. 6(b) that the AE response is synchronous with the current density distribution, with step increase and plateau regions in AE corresponding to spikes and stable regions in current density distribution, respectively.³⁶ The CAEE's sequence of steps with a broad distribution – many small steps, and some very large ones, when a lot of water is suddenly released – is remarkably similar to a randomised Cantor function or fractal Devil's Staircase, discussed by Mandelbrot.⁵³ This reflects the stochasticity of water release, with many trickles and occasional spurts that, however, dramatically dominate the overall behaviour. Whilst the quantity of data are insufficient to ascertain a power law distribution of water releases, the qualitative features of the Devil's Staircase are clear.

At 0.6 V hold, as in Fig. 6(c), the current density decreased from 0.55 A cm^{-2} to 0.44 A cm^{-2} , which can be attributed to gradual water stagnation inside the cell that obstructs the

active sites of the catalyst layer, resulting in performance degradation.⁵⁴ However, the current density profile is less synchronous with CAEE, which is no longer a discrete staircase pattern, as shown in Fig. 6(b). This is due to the continuous water generation as a result of the increased rate of electrochemical reactions (current density) at 0.6 V that resulted in sustained AE generation. Further reducing the cell voltage to 0.4 V, Fig. 6(d), the current density decreased from 1.25 A cm^{-2} to 1.15 A cm^{-2} over the hold period and the corresponding increase in CAEE was much larger and more sustained compared to the previous two voltage holds, which can be attributed to greater amounts of water build-up, as a result of the higher current density in the cell.

In addition to the water generated, the cell temperature developed is another variable that depends on the rate of the electrochemical reactions occurring inside a cell. The cell temperatures during the potentiostatic tests are given in Fig. 7. At lower current densities (corresponding to a cell voltage of 0.8 V), the rate of electrochemical reaction is minimal, which results in a low and stable cell temperature.

With an increase in current density (corresponding to a lower cell voltage of 0.6 V or 0.4 V), the level of electrochemical reaction occurring in the cell increases significantly and this impacts the cell temperature too. Furthermore, the decreasing and increasing trends in cell temperatures observed at 0.6 V

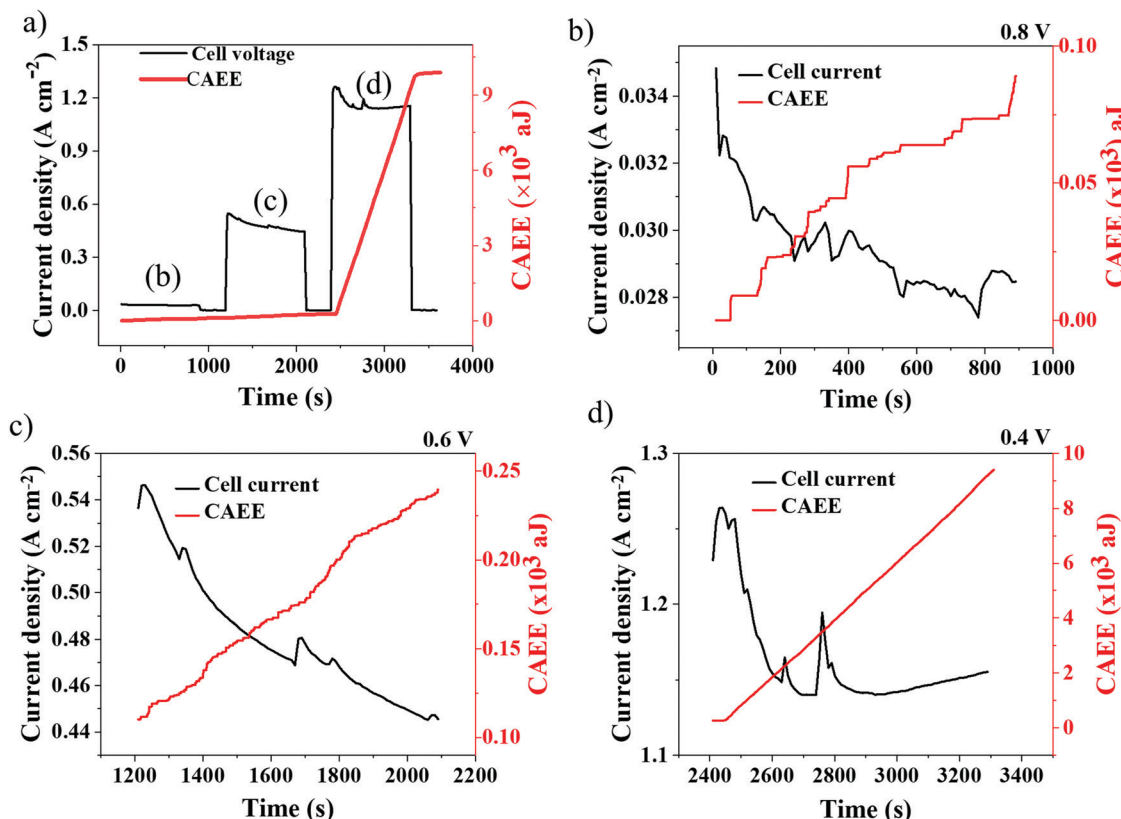


Fig. 6 (a) Overall current density developed during potentiostatic tests at 0.8 V, 0.6 V and 0.4 V; specific current density distribution during potentiostatic tests at (b) 0.8 V, (c) 0.6 V and (d) 0.4 V.

and 0.4 V can be correlated with the corresponding current density (electrochemical reaction rate) trends, as seen in Fig. 7. Besides, the decreasing trend in cell temperature at 0.6 V can be attributed to the cooling effect occurring from liquid water build-up at this condition, which is established by the decreasing trend in HFR at 0.6 V, as observed in Fig. 5(c). Similarly, the increasing trend in temperature at 0.4 V may result from reduced hydration conditions, which is established by an increasing trend in HFR at this voltage, as in Fig. 5(c). Overall, the observed correlation between cell temperature and CAEE

profiles confirms the measured AE to be a dynamic response to the electrochemical reaction rate, similar to the cell temperature.

3.5. Galvanostatic tests

Galvanostatic (current hold) measurements were performed to probe the effect of cell performance on the generated CAEE, specifically at higher current densities, where mass transport limitations are prominent (Table 3). Galvanostatic tests are presented in Fig. 8 for two different current densities, namely 1 A cm^{-2} (25 A) and 1.2 A cm^{-2} (30 A). At each current density, the cell was held for 600 s, followed by an OCV hold for 300 s.⁴⁸

It can be observed from Fig. 8(a) that, at each current density, there is a stabilisation phase, identified by the initial spike(s) in cell voltage, where an equilibrium operation state is achieved. This is known to depend on factors such as reactant concentration, current density, cell temperature, reactant humidity, membrane water content and GDL properties.⁴⁸

During OCV, no acoustic activity (CAEE) is detected (plateau region), implying that the measured CAEE during the cell operation is primarily associated with the water generated from electrochemical reactions. Under imposed current regions, there is a constant increase in the CAEE, which can be attributed to the continuous generation of water as a result of the greater extent of the electrochemical reactions occurring with increased current densities (water generation rates). At 1 A cm^{-2} (Fig. 8(b)) and 1.2 A cm^{-2} (Fig. 8(c)), a voltage peak is

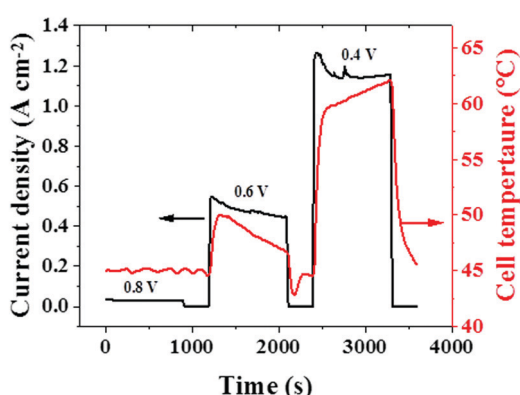


Fig. 7 Cell temperatures measured during potentiostatic tests at 0.8 V, 0.6 V, and 0.4 V.



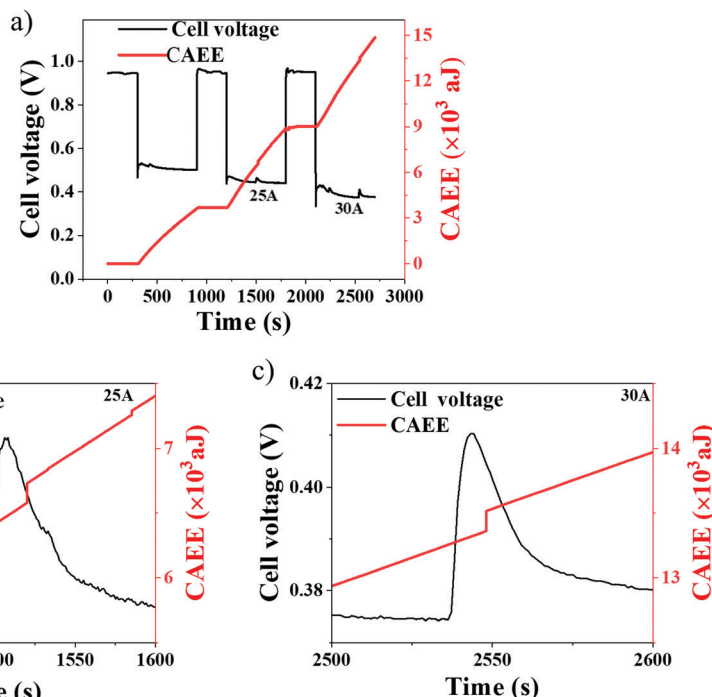


Fig. 8 (a) Voltage and CAEE variation during galvanostatic tests at 20 A (0.8 A cm^{-2}), 25 A (1 A cm^{-2}) and 30 A (1.2 A cm^{-2}); (b) and (c) show voltage and CAEE response for key features during the 25 A (1 A cm^{-2}) and 30 A (1.2 A cm^{-2}) hold, respectively.

observed at 1500 s and 2550 s, respectively. Such peaks occur due to the sudden release of built-up water, as a flushing event, from inside the cell.⁵⁵ Besides, the delay in CAEE edge point at 25 A compared to 30 A may be due to the lower voltage spike generated at 25 A (due to a lower current) compared to that at 30 A. Such flushing events provide improved reactant access for the electrochemical reactions, resulting in improved cell performance. Correlation is observed between the CAEE and voltage peaks in Fig. 8(b) and (c), and can be attributed to flushing events, as described above.³⁰ The corresponding cell temperature developed during the current hold conditions is given in Fig. S3 (ESI†).

4. Conclusions

This study measured acoustic activity from a PEMFC as a dynamic response during potentiostatic and galvanostatic operation. The measured AE and the electrochemical performance are identified to be related to each other at different electrochemical measurement conditions.

Increase in voltage stabilisation (hold) duration results in excess water build-up or flooding inside the cell that decrease its performance, especially in the high current density region. In addition, simultaneous acoustic activity measurements indicate higher levels of AE energy measured at higher stabilisation durations as a result of flooding.

It is shown that AE is an effective water management tool for diagnosing water build-up and removal through forward and reverse polarisation scans; the measured AE during these scans reflects the water uptake and release conditions inside the fuel cell.

Flushing events are observed in galvanostatic and potentiostatic tests. The impact of such events on the cell performance is correlated with the measured AE, with the spikes in current and voltage distributions synchronous with the peaks in CAEE developed. Furthermore, the measured AE as a dynamic response to the electrochemical reactions occurring inside the cell is established by the reached cell temperatures, where higher and lower cell temperatures coincide with larger and smaller levels of AE energy distribution, respectively.

Conflicts of interest

There are no conflicts to declare.

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