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Introduction

Electrochemical production of hydrogen has been carried out at an industrial scale from the 1920s,1 and has attracted great and growing interest in the last few years. Electrical energy, in the best case from renewable sources, is converted into both an efficient fuel and a feedstock for the chemical industry. Nowadays, hydrogen is mainly used in the synthesis of ammonia, methanol and many other bulk chemicals.² Coal gasification, natural gas reforming and partial oxidation of oil cuts are the most common processes for hydrogen production, which cause an emission of 500 Mt CO2 per year.3 In order to increase hydrogen production by water electrolysis, the cost of electrolysers must be reduced.⁴ However, when it comes to both alkaline water electrolysis (AWE) and proton exchange membrane water electrolysis (PEMWE), the capital costs are expected to decrease, mainly due to production scale-up, rather than from R&D investments, because of their high readiness level.5 Nevertheless, AWE cannot be operated at high current densities and has a long response time. PEMWE allows a high production rate and dynamic operation, but depends on both scarce and precious iridium catalysts and porous transport

High performance water electrolysis using a poly(fluorene phenylpropylammonium) anionexchange membrane with 2 M aqueous KOH

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Anion exchange membrane water electrolysis (AEMWE) has great potential to be established as a highperformance and low-capital cost technology for hydrogen production. High current densities can be achieved with a non-platinum group metal (non-PGM) catalyst. However, the harsh operation conditions require stable cell components. Here, we report on the use of a highly stable and ion conductive poly(fluorene alkylene) membrane (PdF–TMA) tethered with trimethylammonium cations *via* phenylpropyl side chains for AEMWE cells operating with 2 M aqueous KOH. The ether-free PdF–TMA polymer is efficiently prepared by polyhydroxyalkylation to reach a molecular weight of 236 kDa, a high thermal stability, and an ion-exchange capacity of 2.14 mequiv. g^{-1} (OH⁻ form). Using commercial electrodes of NiFe₂O₄ (anode) and RANEY® nickel (cathode) and PdF–TMA as an AEM, the output current reached 1 A cm⁻² at voltages below 1.9 V at 60 °C. Also, PdF–TMA outperformed AEMIONTM in terms of membrane resistance by almost 30% and, after 100 h at 0.5 A cm⁻², did not reveal any loss of conductivity, in contrast to AEMIONTM. Furthermore, both membranes were analysed by ¹H NMR spectroscopy after AEMWE tests and the PdF–TMA proved very stable even at 80 °C.

> layers coated with more platinum group metals (PGMs).6 On the other hand, the inherent limitations of these technologies can be overcome by a hybrid device that brings together the benefits of AWE and PEMWE.7 In anion exchange membrane water electrolysis (AEMWE), an alkaline environment is enabled by the use of an anion exchange membrane (AEM). Hence, a non-PGM catalyst and low-cost nickel or stainless-steel plates can be used, just as in AWE.8 Furthermore, a low ohmic cell resistance and a high current density can be achieved in a zero-gap design with a thin and highly ion conductive membrane, just as in PEMWE.9 In recent years, the conductivity and the stability of AEMs have been significantly enhanced and further developed.10 Additionally, AEMWE has the advantage that AEMs are less permeable to hydrogen than PEMs.^{10,11} This reduces the risk of forming an explosive atmosphere due to the lower solubility of hydrogen in solutions with high OH⁻ concentration^{12,13} and the low hydrogen permeability of hydrocarbonbased polymer membranes.

> Nonetheless, despite significant recent progress in the alkaline stability of AEMs, a high degradation rate still limits the lifetime under real conditions.⁷ And, according to Du *et al.*, only a limited number of studies have reported stability over more than 100 h and the degradation rate is seldom in the order of 100–200 μ V h⁻¹.¹⁴ For instance, piperidinium-based AEMs have been shown to be stable for 100 h or more operating in 1 M KOH (or 1 M NaOH)^{15–17} while commercially available imidazolium (SustanionTM)^{17–19} and benzimidazolium (AEMIONTM or

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AEMION+TM)²⁰⁻²² based AEMs have also been tested for 100 h or more under similar conditions. Notably, AEMION+TM has been shown to withstand 1 year of operation in a recent study.²³ Also, trimethyl-ammonium based membranes are promising candidates for AEMWE, as the commercially available ORIONTM.²⁴ Jiang *et al.* have recently reported a membrane that could withstand 3500 h of operation (1 M KOH, 60 °C, 1 A cm⁻²)²⁵ and Jang *et al.* assembled a 3-cell-stack which operated for 2000 h under milder conditions.²⁶ AEMWE durability can also reach several hundred hours with carbonate electrolytes,^{27,28} and more advanced ionomer chemistries have pushed the boundaries of pure water AEMWE to 100–200 h.^{11,28–30}

Also, operating at higher temperature and higher KOH concentration has a positive effect on electrolyte conductivity and catalyst activity, but may further decrease the membrane stability. Along similar lines, some reports have pointed out that the performance of AWE can be enhanced by using a dense polymeric membrane, e.g., polybenzimidazole (PBI), as a separator to absorb electrolyte and conduct the hydroxide anions.13,31 However, in contrast to AEMs, PBI can achieve high conductivities ($\approx 100 \text{ mS cm}^{-1}$) only when placed in contact with 30 wt% KOH solutions,32,33 which decreases the corrosion resistance of plates and other plant components such as pumps, and poses a further challenge to the stability of polymeric materials. The continued development of new membrane polymers over the last decade has greatly improved the stability of AEMs in 1 M aq. KOH solution. Furthermore, some reports have demonstrated the potential application of 2 M KOH.^{34,35}

In the present work a newly synthesized membrane (PdF-TMA) (Scheme 1) and a commercial benchmark membrane (AEMIONTM) have been studied for application in AEMWE at relatively high concentration of KOH (1–2 M) and temperature (60–80 °C). The high-molecular-weight PdF–TMA membrane polymer was tailored for high conductivity and alkaline stability. Hence, the water uptake was tuned to reach high ion conductivity and alkaline stability. The high molecular weight facilitates mechanical stability, which is generally required at high water uptake. As previously demonstrated, attaching the anion exchange groups to the backbone polymer *via* flexible spacer units increases the ionic clustering and the conductivity, as well as the alkaline stability.^{36,37} The membranes were evaluated with commercial electrodes using 1 and 2 M KOH at 60 ° C. The PdF–TMA membrane was also evaluated at 80 °C. The

a) $F_{3}C_{+}$ $F_{4}C_{+}$ $F_{4}C_{+}$

Scheme 1 (a) Synthetic route to the poly(fluorene phenyl-propylammonium) (PdF–TMA) and (b) photograph of a PdF–TMA membrane.

resistivity of the membranes was studied *in situ* by galvanostatic electrochemical impedance spectroscopy (GEIS) before and after a 100 h test. The chemical stability of the AEMs was finally investigated in a post mortem ¹H NMR study.

Experimental

Membrane synthesis

PdF-TMA was synthesized in two steps following a recently reported method.³⁶ First, a precursor polymer (PdF-Br) was prepared via a superacid-mediated Friedel-Crafts polycondensation using 9,9-H,H-dimethylfluorene (dF) and 1bromo-3-(trifluoroacetylphenyl)-propane (TFAp-Br). In a 25 mL round-bottom flask equipped with an ice/water bath, TFAp-Br (1.68 g) and dF (1 g) were dissolved in dichloromethane (DCM, 6 mL) under stirring. Next, trifluoromethanesulfonic acid (1 mL) was dropwise added to the solution. The reaction proceeded at room temperature for 3 h, before the viscous solution was diluted with chloroform followed by precipitation of the product in methanol (500 mL). A white and fibrous precipitate was collected, dissolved in chloroform and re-precipitated in methanol to obtain PdF-Br in a quantitative yield. Functionalization of PdF-Br using trimethylamine was performed in dimethylacetamide (DMAc) at 40 °C. PdF-Br (0.82 g) was dissolved in 20 mL DMAc, before adding a trimethylamine solution (3 mL, 45 wt% in H_2O). The Menshutkin reaction was kept for 7 h, before precipitation of PdF-TMA in diethyl ether. The white powder was washed repeatedly with diethyl ether, before being collected and dried.

AEMs were prepared from a 5 wt% PdF–TMA solution in dimethyl sulfoxide (DMSO). The solution was passed through a syringe-driven PTFE filter ($\Phi = 5 \ \mu m$) onto Petri dishes ($\Phi = 5 \ cm$). Membrane casting was performed in an air-ventilated oven at 80 °C for 2 days. The AEMs were then gently peeled off after hydration with water, and stored in deionized water before characterization.

Membrane characterization

The chemical structures of PdF-Br and PdF-TMA were determined by ¹H Nuclear Magnetic Resonance (NMR) spectroscopy on a Bruker DRX 400 spectrometer at 400 MHz using DMSO-d₆ as solvent. The molecular weight of PdF-Br was characterized by size exclusion chromatography (SEC) using chloroform as eluent, and the glass transition temperature (T_g) of PdF-Br was determined using differential scanning calorimetry (DSC). The ion exchange capacity (IEC) of the PdF-TMA AEM was determined by Mohr titrations. Samples in the Br⁻ form were dried and immersed in aq. NaNO₃ (1 M, 25 mL) for 48 h to ensure complete ion exchange. Next, the solution was titrated with 0.1 M AgNO₃ using K₂CrO₄ as colour indicator. The water uptake of the PdF TMA in the OH⁻ form was measured gravimetrically between 20 and 80 °C. Ion-exchange from the Br⁻ to the OH⁻ was done using 1 M aq. NaOH in a N₂-flushed desiccator. After ion-exchange, the sample was quickly washed in degassed deionized (DI) water, and equilibrated in DI water for 24 h at 20 °C (or 8 h at higher temperatures), before the weight and dimensions were measured. The KOH solution uptake of PdF TMA was measured similarly. After equilibrations in 1 or 2 M aq. KOH solutions, the changes in the weight and dimensions of PdF–TMA were directly recorded. The OH⁻ conductivity of PdF TMA in water was measured using electrochemical impedance spectroscopy (EIS) between 20 and 80 °C.

Electrolysis set-up

The as-prepared PdF-TMA membranes and commercial AF1-HNN8-50-X (AEMION[™], purchased from IonomrTM) membranes with a thickness of 50 µm were evaluated in an AEMWE setup. Both membrane types were soaked in 1 M KOH for at least 36 h prior to the electrolysis, and the KOH solution was changed twice to fully exchange the I⁻ ions (AEMIONTM) or Br⁻ ions (PdF-TMA) to OH⁻. For the electrolysis cell, anodes (NiFe2O4 on stainless-steel felt with Nafion) and cathodes (RANEY® nickel on nickel felt with Nafion) were purchased from Dioxide Material®. Nickel serpentine flow fields with a 5 cm² area were chosen to withstand the high KOH concentration. The active area was isolated to 1 cm² with PTFE gaskets ensuring 80% compression of the electrodes. The thickness of the gaskets was 250 μ m on the cathode and 500 μ m on the anode. A torque of 9 Nm was applied to seal the cell. The flow fields were heated with two heat cartridges, and the temperature controlled by a PID controller with a thermocouple. 1 M or 2 M KOH electrolyte was fed to both the anode and cathode with a peristaltic pump at a flowrate of 3 mL min⁻¹. The electrolyte was pre-heated using two heat exchangers and recirculated after passing through the cell, allowing the gases to disengage in two tanks. An XP4105 potentiostat (Ivium, the Netherlands) with 20 A booster was used for AEMWE trials.

Electrochemical characterization

The electrolysis cell was operated at 60 and 80 °C. Before recording a polarization curve, the cell was subjected to 10 cycles between 1.5 V and 1.8 V with a scan rate of 50 mV s⁻¹ in order to stabilize cell performances. Polarization curves were recorded holding the current constant for 120 s at each current level, where the average value of the cell voltage was calculated during the last 30 s. After recording a polarization curve, GEIS was performed at each current density to evaluate the values of high frequency resistance (HFR). Then, the stability of the membranes was evaluated by applying a current density of 500 mA cm⁻² for 100 h. A second polarization curve was recorded after the stability test, followed by GEIS measurements.

Results and discussion

Ex situ membrane analyses

The molecular weight of the PdF–Br membrane is 236 kDa and was obtained by SEC, while the IEC of the PdF–TMA membrane is 2.14 mequiv. g^{-1} (OH⁻ form). The commercial benchmark (AEMIONTM) was chosen to have a similar IEC (2.1–2.5 mequiv. g^{-1}).³⁸

Table 1 shows the water uptake, OH⁻ conductivity and swelling ratio (in-plane and through-plane) of the PdF-TMA membrane after being fully anion-exchanged and washed in DI water. The AEM showed a very high conductivity, up to 175 mS cm⁻¹ at 80 °C and a water uptake of 190%. When compared to the selected commercial benchmark AEMION™, the PdF-TMA membrane shows a much lower water uptake and swelling ratio. In fact, the in-plane swelling ratio of AEMION™ is reported to be around 140% at 25 °C,²² thereby indicating a much higher water uptake than that measured for PdF-TMA. In fact, excessive water uptake and dissolution in pure water is reported for non-crosslinked hexamethyl-p-terphenyl-poly(benzimidazolium) (HMT-PBI) membranes such as AEMIONTM.³⁹ In addition, the hydroxide conductivity is reported to be around 50 mS cm⁻¹ at 40 °C for AEMIONTM,⁴⁰ while the conductivity of PdF-TMA was measured to be twice as high. Considering the high conductivity and the anticipated high stability of PdF-TMA, the two membrane types were compared in an electrolysis cell operating with 1 M and 2 M KOH, respectively.

Electrolysis using 1 M KOH at 60 °C

Fig. 1a shows the polarization curve of AEMWE obtained before (solid symbols) and after the 100 h stability test (open symbols) at 60 °C using 1 M KOH and two different membranes: PdF-TMA (blue circles) and AEMION™ (AF1-HNN8-50-X, black squares). At low current densities (<0.1 mA cm^{-2}), similar cell voltages were obtained with both AEMs as the electrode kinetics limited the cell performance. However, at higher current densities, lower cell voltages were obtained when the AEMION[™] membrane was used. Furthermore, the HFR for the cell with AEMION™ was 30% lower than for the one with PdF-TMA (Table 2), indicating a higher conductivity of AEMION[™] in 1 M KOH at 60 °C. These data agree well with the HFR data reported by Pushkareva et al.,⁴¹ who used AEMION™ with PGMfree catalysts for AEMWE in 1 M KOH at 60 °C. Also, both AEMs showed almost no variation in HFR at varying current densities, indicating the absence of dehydration of the membrane at

Table 1	Membrane	properties	of	PdF-TMA
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Temperature (°C)	Water uptake (wt%)	OH ⁻ conductivity in water (mS cm ⁻¹)	Swelling ratio (%, in-plane)	Swelling ratio (%, through-plane)
20	113	78.5 ± 2.5	25.6 ± 0.2	20.1 ± 1.6
40	135	104 ± 3	30.7 ± 0.3	23.4 ± 2.7
60	163	136.5 ± 4.5	33.3 ± 1.7	28.4 ± 2.3
80	190	171 ± 4	42.1 ± 0.9	32.3 ± 1.9

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Fig. 1 (a) Polarization curves obtained in 1 M KOH at 60 °C with PdF–TMA (black squares) and AEMIONTM (AF1-HNN8-50-X) (blue circles), before (solid symbols) and after the stability test (open symbols). Commercial electrodes with NiFe₂O₄ (anode) and RANEY® nickel (cathode) were used. (b) Stability evaluations at 500 mA cm⁻² in 1 M KOH at 60 °C showing the degradation rate of the PdF–TMA (black) and AEMIONTM (AF1-HNN8-50-X), blue circles), before used.

Table 2 HFR data of the electrolysis cells under different conditions

	AEMIONTM	4 (AF1-HNN8-50-X)	PdF-TMA		
KOH concentration (M)	1	2		1	2	
Temperature (°C)	60	60	80	60	60	80
HFR ^{<i>a</i>} (m Ω cm ²) before the stability test	110	128	105	162	86	75
HFR^{a} (m Ω cm ²) after the stability test	106	169	180	165	95	77
^{<i>a</i>} Measured at 100 mA cm ^{-2} .						

higher current densities, since enough water is provided from the KOH electrolyte.⁴²

After the stability evaluation at 500 mA cm⁻² for 100 h, both cells showed a significant decrease in performance. Fig. 1b shows the long-term results with degradation rates of 0.19 mV h⁻¹ and 0.26 mV h⁻¹ for the cells with AEMIONTM and PdF-TMA, respectively. The HFR values recorded before and after the long-term tests (Fig. 1a) are close to their initial values, suggesting stable membrane operation as the membrane resistance is the main contributor to HFR. The loss of performance at both low and high current densities is likely related to the electrodes rather than to the membranes and is possibly caused by detachment of catalyst particles from the electrodes. Actually, a colour change was observed at the outlet of the cell and the electrode appeared discoloured. We can conclude that the degradation rates are similar for the two AEMs in 1 M KOH at 60 °C as the major cause of loss of performance was due to the electrodes.

Du *et al.* identified more than 100 h of operation as the benchmark for AEMWE.¹⁴ In the present case, we exceeded 100 h of operation with a degradation rate of the AEMIONTM membrane below 200 μ V h⁻¹, which is quite low when compared with data reported by Du *et al.* after surveying the performances of AEMWE cells disclosed in the literature.¹⁴ This confirms that AEMIONTM can be efficiently used for AEMWE

due to the high conductivity and stability, as previously reported by Khataee *et al.*⁴³ Pavel *et al.*²⁷ have reported 200 μ V h⁻¹ after using an A201 membrane (Tokuyama) in a much less aggressive environment (1 wt% K₂CO₃ at 43 °C) with a higher HFR. With the same materials, Vincent *et al.*⁸ have reported 500 μ V h⁻¹ (1 wt% K₂CO₃ at 60 °C). Only Motealleh *et al.*¹⁸ have reported a much lower degradation rate using the SUSTANIONTM membrane in 1 M KOH. On the other hand, Park *et al.* obtained a higher degradation rate with the same membrane.¹⁹

Electrolysis using 2 M KOH at 60 °C

Given the high stability of the membranes and to improve the electrode performance, evaluations were also carried out in 2 M KOH. Fig. 2a shows the polarization curves obtained before (open symbols) and after the stability evaluation (solid symbols) in 2 M KOH. Once again, the performance of the two membranes was found to overlap at low current densities. Both cells showed improvements at low current density when 2 M KOH was used instead of 1 M KOH, which agrees with previous reports.^{41,44,45} This is likely due to the beneficial effects of the higher hydroxide concentration on both the OER and HER kinetics, and on the ion transport. However, the performance of the cell with AEMION[™] was limited by an increased membrane resistance at higher current densities. In fact, its HFR increased



Fig. 2 (a) Polarization curves obtained in 2 M KOH and at 60 °C with PdF–TMA (black squares) and AEMIONTM (AF1-HNN8-50-X, blue circles) before (solid symbols) and after stability evaluation (open symbols). Commercial electrodes with NiFe₂O₄ (anode) and RANEY® nickel (cathode) were used. (b) Stability evaluations at 500 mA cm⁻² in 2 M KOH at 60 °C showing the degradation rates of the PdF–TMA (black) and AEMIONTM (AF1-HNN8-50-X, blue) cells.

from 110 m Ω cm² to 128 m Ω cm² after shifting to 2 M KOH (Table 2). In contrast, the PdF-TMA membrane showed a remarkable increase in conductivity as the HFR decreased from 162 m Ω cm² to 86 m Ω cm² (Table 2). This reduction in HFR by almost 50% allowed the cell with the PdF-TMA membrane in 2 M KOH at 60 °C to reach a lower cell voltage at a higher current density than AEMIONTM in 1 M KOH at the same temperature.

The degradation of the cell using PdF–TMA was lower in 2 M KOH (0.16 mV h⁻¹) than in 1 M KOH (0.26 mV h⁻¹). Furthermore, in 2 M KOH the stability evaluations revealed a more precise difference between the two membranes. The cell using PdF–TMA showed a higher stability (0.16 mV h⁻¹) than the one using AEMIONTM (0.34 mV h⁻¹). The HFR remained almost constant after the 100 h stability evaluation of PdF–TMA,

whereas a significant increase ($\approx 40 \text{ m}\Omega \text{ cm}^2$) was observed for AEMIONTM. This dramatic loss in performance is clear from the polarization curves (Fig. 2a) and the stability results (Fig. 2b). The degradation of the cell with AEMIONTM in 2 M KOH may be caused by the combination of electrode and membrane degradation, while the cell with PdF–TMA seemed to only suffer from electrode degradation.

Electrolysis using 2 M KOH at 80 °C

To explore the possibility of further improving the electrolyser performance, both AEMs were evaluated at 80 °C in 2 M KOH (Fig. 3). The increased temperature enhanced the performances at all current densities, favoring electrode kinetics and ion transport. The HFR was found to decrease to 105 and 75 m Ω cm² for AEMIONTM and PdF–TMA, respectively (Table 2).



Fig. 3 (a) Polarization curves obtained in 2 M KOH and at 80 °C with PdF–TMA (blue circles) and AF1-HNN8-50-X (AEMION^M) (black squares) before (solid symbols) and after (open symbols) the stability test. Commercial electrodes with NiFe₂O₄ (anode) and RANEY® nickel (cathode) were used. (b) Stability evaluations at 500 mA cm⁻² in 2 M KOH and at 80 °C indicating the degradation rate of the PdF–TMA (black) and AEMION^M (AF1-HNN8-50-X, blue) cells.



Fig. 4 ¹H NMR spectra of the AEMION™ (AF1-HNN8-50-X) membrane before (a) and after long-term stability evaluation in the electrolysis cell (b-d).

However, the stability evaluation showed a substantial performance loss of the cell with AEMIONTM over time. The degradation rate was higher at 80 °C (0.44 mV s⁻¹) than at 60 °C (0.34 mV s⁻¹). Notably, the HFR increased to 180 m Ω cm². In contrast, the resistance of PdF-TMA did not show any appreciable increase and the HFR of the cell did not exceed 77 m Ω cm². Also, the performance decay was less evident at 80 °C than at 60 °C (Fig. 3b). In fact, the degradation rate of the cell with PdF-TMA was lower at 80 °C (0.10 mV h⁻¹) than at 60 °C (0.16 mV h⁻¹). This can be explained by a higher activity of the electrodes and a higher conductivity of the electrolyte at 80 °C, which may compensate for any degradation of the electrodes.

Even though the cell using the PdF–TMA membrane showed a similar, or higher, degradation rate in 2 M KOH compared to what Schauer *et al.* and Hnát *et al.* achieved with a quaternized qPPO membrane (0.1 mV h⁻¹)³⁴ and a DABCO functionalized PSEBS membrane (0.02 mV h⁻¹),³⁵ respectively, both these AEMs show a lower conductivity than PdF–TMA. In addition, considering that the cell with PdF–TMA in 2 M KOH at 60 °C and 80 °C surpassed the good performance of AEMION[™] in 1 M KOH, the PdF–TMA can be considered to be an excellent AEM material for AEMWE applications at high KOH concentration.

NMR spectroscopy of AEMs after stability evaluation

To further investigate the stability of the AEMs, ¹H NMR analysis was performed on the membranes before and after the cell evaluation. Fig. 4a shows the spectrum of the pristine AEMIONTM membrane (AF1-HNN8-50-X), while Fig. 4b shows the NMR spectrum of the same membrane after evaluation in the electrolysis cell for 100 h at 0.5 mA cm⁻² in 1 M KOH at 60 ° C. As seen, after these conditions some structural changes were detected (Fig. 4c). This further confirms that the loss of conductivity was due to polymer degradation caused by the high OH⁻ concentration. A similar chemical degradation was also detected after operation at 80 °C (Fig. 4d).



Fig. 5 ¹H NMR spectra of the PdF–TMA membrane before (a) and after long-term stability evaluation in the electrolysis cell (b–d).

In contrast, the NMR spectra of PdF–TMA remained unchanged (Fig. 5), indicating an intact molecular structure after 100 h of operation in 1 M KOH at 60 °C, and even in 2 M KOH at 80 °C. This supports the finding that the PdF–TMA membrane is highly stable at higher OH⁻ concentrations, as the electrolysis data did not indicate a decrease in conductivity (Table 2).

Electrolyte uptake

As previously reported, the concentration of the KOH electrolyte highly influences the OH[−] conductivity. A high KOH concentration promotes absorption of hydroxide ions within the membrane and provides extra charge carriers.¹⁰ On the other hand, this environment reduces the membrane water content due to a higher osmotic pressure difference, hindering OH[−] transport through the ionic channels.⁴⁶ From the results in Table 2, AEMIONTM shows a good trade-off between these effects in 1 M KOH, but suffers from a decreased conductivity in 2 M KOH. In contrast, the PdF–TMA membrane shows improved conductivity in 2 M KOH, which may be due to increased KOH absorption. Given these premises, the electrolyte uptake of the two membranes was quantified to understand their different behaviours in the electrolyzer cells at 1 M KOH and 2 M KOH.

Table 3 shows the electrolyte uptake and swelling ratio of the two AEMs when immersed in 1 M and 2 M KOH. Considering the electrolyte uptake and through-plane swelling ratio of AEMION[™], this membrane tends to absorb more electrolyte solution and swell more at all combinations of temperature and concentration. For this reason, the higher cell resistance recorded for PdF-TMA in 1 M KOH at 60 °C (Table 2) is unlikely due to a higher membrane thickness under the operation conditions. Even though the PdF-TMA membrane has a high conductivity in water (Table 1), AEMION™ becomes more conductive than this membrane in 1 M KOH due to a higher KOH absorption. On the other hand, AEMION[™] likely suffers from excessive loss of water in 2 M KOH, and the extra charge carriers cannot improve the conductivity. The PdF-TMA membrane performs better in this harsher environment due to its higher water retention. In fact, even though both membranes show a lower uptake of the electrolyte in 2 M KOH than in 1 M KOH, the decrease in percentage is lower for PdF-TMA than for AEMION[™]. Not only PdF-TMA can benefit from KOH absorption at this higher concentration, but also the water balance can play a role. As a matter of fact, an excessive water uptake can dilute the charge carriers in the membrane.46

The water uptake directly influences also the degradation of the membrane. When cationic groups are less solvated by water molecules, they become less protected from OH[−] attack.¹⁰ AEMIONTM suffered from a high degradation rate when exposed to 2 M KOH, which de-waters the AEM, as previously discussed for the conductivity. PdF–TMA did not reveal any signs of degradation even at 80 °C, while for AEMIONTM the degradation accelerated at this temperature. Besides the higher water retention, the improved stability of PdF–TMA is also due to the effect of the flexible phenylpropyl spacer chain (Fig. 4d), which protects the quaternary ammonium cations from hydroxide attack *via* steric hindrance.¹⁰ Notably, PdF–TMA and AEMIONTM showed different

Table 3 Electrolyte uptake a	nd in- and thr	ough-plane s	velling of the	AEMION TM ai	nd PdF-TMA	membrane	as a function c	of temperature	and KOH cor	ncentration		
	AF1-HNN8-5	0-X (AEMION	(M)				PdF-TMA					
KOH concentration (M)	1			2			1			2		
Temperature (°C)	20	60	80	20	60	80	20	60	80	20	60	80
Electrolyte uptake (%)	52	95	108	44	63	67	57	78	84	45	59	62
In-plane swelling (%)	11.6 ± 4.7	18.0 ± 3.7	16.7 ± 5.7	8.2 ± 4.1	9.8 ± 2.5	8.9 ± 3.3	14.3 ± 2.1	20.6 ± 0.9	23.5 ± 1.5	$\textbf{8.6}\pm\textbf{1.4}$	0.4 ± 1.6	13.9 ± 1.9
Through-plane swelling (%)	30	48	50	20	32	36	10	15	18	11	14	18

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extents of electrolyte uptake and water retention at similar ion exchange capacities (IEC). Even though the water uptake is highly dependent on this parameter, PdF–TMA may more readily absorb water due to its more flexible structure, which allows the ionic channels to "stretch" and host more water. Also, an improved phase separation may result in water channels with higher internal cation concentrations, which may explain the higher water uptake.

Conclusions

In this work, we have investigated the application of a poly(fluorene alkylene) membrane tethered with trimethylammonium cations via flexible phenylpropyl side chains for AEMWE. Since the membrane displayed a high alkaline stability at 1 M KOH, it was evaluated at higher KOH concentration and higher temperature (2 M KOH, 80 °C) than usual for AEMWE to boost electrolysis performances. Not only did the higher KOH concentration improve the kinetics and ion transport in the electrode, but the PdF-TMA membrane was also shown to be more conductive in 2 M KOH than in 1 M KOH at 60 °C. In comparison. the commercial benchmark **AEMION**TM membrane showed loss of conductivity and stability in 2 M KOH at 60 °C. Moreover, the conductivity of PdF-TMA increased without compromising its stability when operated at 80 °C in 2 M KOH, thus allowing a high electrolyzer performance. The chemical stability of the membrane was demonstrated by ¹H NMR analysis of AEMs after the evaluation. The PdF-TMA membrane shows a lower relative loss of electrolyte when KOH concentration is increased from 1 to 2 M. We speculate that the more flexible structure of PdF-TMA facilitates the retention of the electrolyte solution at high KOH concentrations.

Author contributions

Matteo Rossini: conceptualization, methodology, investigation, writing – original draft. Dong Pan: conceptualization, methodology, investigation, writing – review & editing. Burak Koyutürk: methodology, investigation, writing – review & editing. Si Chen: investigation. Amirreza Khataee: supervision, writing – review & editing. Göran Lindbergh: supervision, writing – review & editing. Patric Jannasch: supervision, writing – review & editing. Ann Cornell: supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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