A Raman spectro-microscopic investigation of ETFE-based radiation-grafted anion-exchange membranes†

Wai Hin Lee, Carol Crean, John R. Varcoe and Rachida Bance-Soualhi*†

This study used Raman spectro-microscopy to investigate the synthesis and degradation of radiation-grafted anion-exchange membranes (RG-AEM) made using 50 μm thick poly(ethylene-co-tetrafluoroethylene) (ETFE) films, vinylbenzyl chloride (VBC) monomer, and 1-methylpyrrolidine (MPY) amination agent. The data obtained confirmed the operation of the grafting-front mechanism. VBC grafting times of 1 and 4 h led to low degrees of grafting homogeneity, while 72 h led to extreme levels of grafting that resulted in mechanically weak RG-AEMs due to the excessive H2O contents. A grafting time of 16 h was optimal yielding a RG-AEM with an ion-exchange capacity (IEC) of 2.06 ± 0.02 mmol g⁻¹ (n = 3). An excess of grafting was detected at the surface of this RG-AEM (at least within the first few μm of the surface). This RG-AEM was then degraded in O2-purged aqueous KOH (1.0 mol dm⁻³) for 14 d at 80 °C. Degradation was detected throughout the RG-AEM cross-section, where the Raman data was quantitatively consistent with the loss of IEC. A slight excess of degradation was detected at the surface of the RG-AEM. Degradation involved the loss of whole benzyl-1-methylpyrrolidinium grafted units as well as the direct attack on the pendent (cationic) pyrrolidinium groups by the hydroxide anions.

Introduction

Radiation-grafting is a useful technique for the production of functional membranes (and other polymeric materials). This technique has been commonly used to produce solid electrolytes, e.g. ion-exchange membranes (IEM), including proton-exchange membranes (PEM) for use in technologies such as proton-exchange membrane fuel cells (PEMFCs) and PEM-based water electrolyzers. A more recent line of research is the development of anion-exchange membranes (AEM) for use in alkaline Anion-Exchange Membrane Fuel Cells (AEMFCs) and AEM-based Alkaline Water Electrolysers (AEM-AWE); this includes the use of radiation-grafted (RG) types. The use of high pH, hydroxide-conducting solid electrolytes allows for a broader range of electrocatalysts that do not involve scarce, expensive Pt. Scheme 1 outlines the steps involved in the synthesis of such a RG-AEM: pre-irradiation in air (peroxidation) of poly(ethylene-co-tetrafluoroethylene) (ETFE) precursor film, followed by grafting with vinylbenzyl chloride (VBC) monomer and amination with 1-methylpyrrolidine (MPY).

Incomplete grafting can lead to RG-IEMs with lower than ideal functionalisation, which are then too ionically resistive for application. Radiation-grafting often involves the grafting-front mechanism, especially with thicker precursor films; this is where grafting initiates on the surfaces of the precursor films.
with the gradual penetration of the grafts into the bulk of the films (Fig. 1). As radiation-grafting involves the modification
of solid polymer films (up to 150 μm thickness in the literature),
it is essential to assess the uniformity of the monomer grafting
and any subsequent functionalisation throughout the core of the
resulting RG-IEMs. Techniques that have been used to study the
uniformity of grafting include Scanning Electron Microscopy coupled with Energy-Dispersive X-ray Spectroscopy (SEM-EDX)\(^6\)\(^9\)\(^1\)\(^0\)\(^2\) and Raman spectro-microscopy.\(^6\)\(^9\)\(^1\)\(^0\)

Raman spectroscopy can achieve spectral resolutions down to ca. 0.3 cm\(^{-1}\) and is a powerful spectroscopy that both
complements and supports other spectroscopies (vibrational,
rotational, and electronic).\(^1\)\(^1\) As Raman spectrometers involve
the use of lasers, the spectrometers are commonly coupled to
(confocal) microscopic equipment that allows individual
spectra to be recorded on different parts of a sample with spatial
resolutions as low as ca. 1 μm diameter.\(^1\)\(^1\)\(^2\)\(^3\) As the cost of Raman
instrumentation has generally decreased in the last 10 years\(^4\)
and the familiarity with the technique has increased, the use of
Raman spectro-microscopy is becoming more widespread and
routine in the field of materials science. The use of Raman
spectro-microscopy often has the advantage of requiring
minimal levels of sample preparation. The primary disadvantage
is that the quality of spectra is often degraded due to the
presence of photoluminescence interferences such as fluorescence.
If Raman equipment is available that possesses multiple
laser wavelengths (λ), it is often possible to select a laser
wavelength that minimises such interferences (but, obviously,
this adds to the cost of the equipment).

Given the spatial resolution of Raman spectro-microscopy
it can be as low as 1 μm during routine measurements (and that
previous RG-IEMs reported in the literature are typically up to
ca. 150 μm in thickness),\(^1\) this technique can be a useful quality
assurance tool for the study of the radiation-grafting homogeneity
especially through the membrane cross-section. Raman
spectro-microscopic mapping has already been used to study
RG-AEMs, but these experiments have typically taken 8–16 h per
map (as Raman scattered light is of low intensity);\(^6\)\(^9\)\(^1\)\(^0\)\(^2\) this is non-
ideal considering the life-times of expensive (£10k+) lasers is
typically only a few thousand hours (of in use time).

Herein, the aim of this study is to investigate the optimum
laser wavelength and spectral collection parameters that will
allow for the more rapid and routine use of Raman spectro-
microscopy in the study of each stage (grafting, amination) of
the synthesis of RG-AEMs. This study will then use this
knowledge to conduct an initial investigation into the homo-

Experimental

Materials and chemicals

Commercial Nowoflon ET-6235Z ETFE film of 50 μm thickness
was supplied by Nowofol Kunststoffprodukte GmbH (Germany)
and was used as the precursor material. Vinylbenzyl chloride
(VBC, 97%), mixture of 3- and 4-isomers, used without removal
of 4-tert-butylicatechol and nitromethane inhibitors), 1-octy-2-
pyrrolidone dispersant, and 1-methylpyrrolidinone (MP, 97%)
were purchased from Sigma-Aldrich. Standardised analytical
aqueous solutions of AgNO\(_3\) (20.00 ± 0.06 mmol dm\(^{-3}\)) and
aqueous HNO\(_3\) (2.0 mol dm\(^{-3}\)), used for the IEC determina-
tions, were supplied from Fluka. All other chemicals, including
analytical grade aqueous solutions of NaCl (1.0 mol dm\(^{-3}\)) and
KOH (1.0 mol dm\(^{-3}\)), toluene, and NaNO\(_3\) (8) were used as
received. Ultrapure water (UPW, resistivity = 18.2 MΩ cm) was
used throughout this study.

Preparation of the radiation-grafted anion-exchange
membranes

**Grafting stage.** The pre-aminated ETFE-g-poly(VBC)
membranes were prepared by grafting VBC onto peroxidated
ETFE films as described previously.\(^6\) The ETFE films (13 ×
13 cm) were irradiated in air to 70 kGy total absorbed dose using
a 4.5 MeV Dynamatron Continuous DC Electron Beam Unit
(Synergy Health, South Marston, UK, see Fig. S1 in the in the
ESI†). The peroxidated films were immersed into glass tubes
containing a mixture (250 cm\(^3\)) of 5% vol. VBC, 1% vol. 1-octy-2-
pyrrolidone, and 94% vol. UPW, which was then purged with N\(_2\)
for 2 h. The sealed tubes were subsequently placed in a pre-
heated water bath at 70 °C for the desired duration of grafting
reaction (1–72 h). Finally, the grafted membranes were thor-
oughly washed with toluene and then dried in a vacuum oven at

---

\(\dagger\) Sub-micron resolution can be less routinely obtained with the use of high cost
equipment and low wavelength lasers, especially with the use of tip-enhanced
Raman spectroscopy (TERS), which couples atomic force microscopes with
Raman instrumentation.

\(\S\) Raman microscopes are still generally more expensive than for IR
instrumentation (>£100k vs. <£50k).
70 °C. The degree of grafting (DoG) of each pre-aminated membrane was calculated as follows:

$$\text{DoG} (\%) = \frac{m_g - m_i}{m_i} \times 100$$  (1)

where \( m_g \) is the mass of the grafted membrane and \( m_i \) is the initial mass of the peroxidated ETFE film used.

**Amination stage.** A previous study revealed that amination with MPY yielded RG-AEMs that were more stable towards attack by OH\(^-\) anions (more alkali stable) than RG-AEMs synthesised using the more commonly encountered trimethylamine (TMA) reagent. Therefore, MPY was used to convert the pre-aminated membranes into the desired RG-AEM(Cl\(^-\))s. The pre-aminated ETFE-g-poly(VBC) membranes were submerged in aqueous MPY solution (50% vol.) and heated at 70 °C for 24 h. The crude RG-AEMs were then washed with UPW before being heated in water for 6 h at 70 °C. Subsequently, the membranes were soaked in aqueous NaCl (1.0 mol dm\(^{-3}\)) solution for 12 h (with several changes of solution during this period to ensure the Cl\(^-\)–anion forms of the RG-AEMs). The final RG-AEM(Cl\(^-\))s were obtained after thorough washing with UPW (over 16 h with at least 5 changes in UPW to remove all excess counter- and co-ions), and stored in UPW until required for use in experiments.

**Membrane designations**

The following nomenclature is used to unambiguously identify the different samples (see Scheme 1):

- **Ex** a final RG-AEM(Cl\(^-\)) synthesised with \( x \) h grafting time;
- **Intx** the pre-aminated ETFE-g-poly(VBC) membranes used to fabricate Ex;
- **AE16** the E16 RG-AEM(Cl\(^-\)) that has been aged for 14 d at 80 °C in aqueous KOH (1.0 mol dm\(^{-3}\)) solution that has been purged with \( O_2 \) (see below).

**Water uptakes (WU)**

RG-AEM(Cl\(^-\)) samples were removed from the storage water and excess surface water was removed by dabbing with filter paper. The hydrated masses \( m_{hyd} \) were speedily recorded (to avoid dehydration on prolonged exposure to the atmosphere). The AEM samples were subsequently dried in a vacuum oven at 50 °C for 24 h before the dry masses \( m_{dry} \) were speedily recorded (dry AEMs are generally hygroscopic). All measurements were repeated on \( n = 3 \) samples of each RG-AEM(Cl\(^-\)). The WU value for each sample was then calculated:

$$\text{WU} (\%) = \frac{m_{hyd} - m_{dry}}{m_{dry}} \times 100$$  (2)

**Ion-exchange capacities (IEC)**

The dehydrated RG-AEM(Cl\(^-\))s samples \( m_{dry}/g \) collected straight from the WU measurements above were individually immersed into excess aqueous NaNO\(_3\) (2.4 mol dm\(^{-3}\), 20 cm\(^3\)) solutions for 24 h. The solutions, still containing the AEM samples, were then acidified with aqueous HNO\(_3\) (2.0 mol dm\(^{-3}\), 2.0 cm\(^3\)) and titrated using the aqueous AgNO\(_3\) (20.00 ± 0.06 mmol dm\(^{-3}\)) solution. A Metrohm 848 TitrinoPlus autotitrator equipped with a Ag-Titrode was used for the titrations. The IEC for each sample was calculated from the end-point \( (E_p) \):

$$\text{IEC} \ (\text{mmol g}^{-1}) = \frac{E_p \ (\text{cm}^3) \times 0.02 \ (\text{mmol dm}^{-3})}{m_{dry} \ (g)}$$  (3)

**Alkali stability testing**

A select RG-AEM(Cl\(^-\)), synthesised with 16 h of grafting reaction duration \( (E_{16}) \), was subjected to alkali aging tests. A sample of \( E_{16} \) \((7 \times 7 \text{ cm})\) was immersed in an excess of aqueous KOH (1.0 mol dm\(^{-3}\)) solution that was then purged with \( O_2 \) for 1 h. Previous studies have indicated that alkali degradation is more extreme in \( O_2\)-containing aqueous alkali compared to inert-gas purged aqueous alkali. The polypropylene bottle was then immediately sealed (after purging) and placed in oven at 80 °C for 14 d. After this alkali ageing process, the sealed bottle was cooled to room temperature and the aged sample (designated AE16) was then re-exchanged back to the Cl\(^-\)–anion form using ion-exchange with aqueous solution NaCl (1.0 mol dm\(^{-3}\)) followed by thorough washing with UPW for 16 h (with at least 5 changes of fresh UPW during this period). As well as the Raman experiments, the IEC of AE16 was also determined.

**Raman spectro-microscopic procedures**

Raman spectra and cross-sectional maps were recorded using an InVia Reflex Raman Microscope (Renishaw, UK) using 4 laser excitation wavelengths (Table 1). The Raman microscope was fitted with a cooled charged coupled detector (CCD) along with holographic notch filters and gratings tailored for each laser wavelength. The attached Leica DMLM optical microscope was equipped with different objective lenses and a trinocular viewer that accommodates a video camera, allowing direct viewing of the sample. Daily calibration of the instrument was conducted by recording the Raman spectrum of silicon in static mode. If necessary, an offset correction was performed to ensure that the position of the silicon peak to be 520 ± 1 cm\(^{-1}\).

| Wavelength/nm | 785 | 633 | 532 | 457 |
|---------------|-----|-----|-----|-----|
| Max. power/mW | 300 | 20  | 50  | 20  |
| Colour        | Line| Spot| Spot| Spot|
| Grating/lines mm\(^{-1}\) | 1200 | 1800 | 2400 | 2400 |
| Laser spot size/μm | 1.28 | 1.03 | 0.87 | 0.74 |
| Filter cut-off/cm\(^{-1}\) | 50  | 100 | 100 | 100 |

\( ^{\dagger} \) The Renishaw Raman instrument used contains a 5th laser (244 nm deep UV) but this is not appropriate for this study: the laser damages the polymer samples, even at the lowest power setting.
Table 2 The parameters used to obtain the maps for the pre-aminated ETFE-g-poly(VBC) membranes (Intx where x = the grafting reaction time/h) with par-sampling (sample-stage step size = theoretical minimum laser spot diameter)

| Parameter          | Wavelength/nm | Power at sample/mW | Step size/µm | Exposure time/s | Total mapping time/min |
|--------------------|----------------|--------------------|--------------|-----------------|------------------------|
| 633                | 20             | 1.03               | 1            | 1.5             | 40                     |
| 457                | 2              | 0.74               | 1.5          | 40              | 110                    |

* Calculated for 30 × 65 µm cross-sections with 1 spectral accumulation.

In this study, the laser beam was focused on the sample using a 50× (NA = 0.75) objective to collect the backscattered light so the resulting laser spot diameter was ca. 1 µm (see Table 1). Other spectral collection parameters are presented in the figure captions (as appropriate). The raw data was processed using Renishaw’s Wire 4.3 software: processing included baseline correction, smoothing, normalization, interpolation, and curve fitting of the peaks with the Gaussian–Lorentzian function to obtain accurate peak areas (where appropriate).

For the cross-sectional (through-plane) mapping experiments (Fig. 1), the membrane samples were held and pressed between two metal holders and fresh edges were obtained by sectioning using a scalpel. The membrane samples were mounted perpendicular to the sample-stage so that the exposed cross-sections are parallel to the stage. The stage was moved in the x- and y-direction under computer control. Cross-sectional maps were collected with point mapping using the instrument’s static scan mode over the spectral range 600–1700 cm⁻¹. Maps were recorded over 30 × 65 µm cross-sectional areas (all the same so that the experiments could be timed).

For the pre-aminated ETFE-g-poly(VBC) membranes, the optimised mapping parameters used are presented in Table 2 (for par-sampling where sample step size = theoretical min. laser spot diameter). Maps with different sample step sizes were also performed on a sample of Int4 to determine an appropriate step size to be used for mapping experiments (Fig. 2): both under-sampling (step size > theoretical min. laser spot diameter) and over-sampling (step size < theoretical min. laser spot diameter) were studied.

For the final E16 and the alkali-aged AE16 samples, cross-sectional maps were collected using the 785 nm line laser (pinhole applied, 300 mW power) with a step size of 1.28 µm (par-sampling) in both the x- and y-directions. These spectral maps were collected using a 50× objective, 5 s exposure times, and averaging over 2 accumulations (mapping time of 220 min).

Results and discussion

Membrane characterisation

A series of pyrrolidinium-based RG-AEM(Cl⁻)s with various degrees of grafting (DoG) were prepared via the peroxidation radiation-grafting of VBC monomer onto ETFE films followed by amination with 1-methylpyrrolidine (MPY) (Scheme 1). The IECs and WUs of the AEMs are presented in Table 3 (Fig. S2 in the ESI† shows the variation in IEC vs. DoG for the RG-AEM(Cl⁻) s). The IECs of E16 and E72 were 80% of the theoretical IECs (calculated from DoG) due to side-reactions as explained previously (e.g. conversion of –CH₂Cl to –CH₂OH).† The WU is related to the number of ion-exchange sites. As expected, the WUs of the AEM(Cl⁻)s increased with IEC (Fig. S2 in the ESI†), due to the higher hydrophilic content levels. The IEC and WU values obtained for RG-AEM(Cl⁻)s are bulk averages and do not give any information on the spatial distribution of the grafted quaternary ammonium groups. This is why techniques, such as Raman microscopy, are required to evaluate such micro-phase segregation in IEMs, e.g. the RG-AEMs in this study.||

Raman spectroscopy to elucidate optimal laser wavelength

A vital consideration for Raman spectro-microscopy is the selection of laser excitation wavelength (λ).14 This choice is not always straightforward and must balance the following requirements:

- Maximised intensity of the Raman scattered radiation to allow for more rapid collection of spectral data: this is dictated (at each wavelength) by the Raman efficiency and the highest level of laser power that can be applied that does not cause sample damage;
- Minimised level of fluorescence to facilitate spectral data processing and to ensure all relevant peaks are observed;
- Minimised spectral resolution to ensure adequate resolution of overlapping peaks;
- Ability to obtain the spatial resolution required, which is partially controlled by the laser spot diameter obtainable (Table 1): the minimum Airy disk diameter = 1.22 × λ/NA (where NA is the aperture of the objective being used).15

The intensity of Raman scattering is proportional to λ⁻⁴,16 hence, a shorter wavelength will yield a stronger Raman signal: the intensity of the scattered radiation with a 633 nm laser is only 27% of the intensity obtained with a 457 nm laser. For the

Fig. 2 A diagram summarising the different sampling modes used in the Raman mapping experiments.
higher wavelength laser, this intrinsic lower intensity can often be offset by increasing the laser power as long as sample damage does not occur (higher wavelength radiation yields less $J$ of energy per photon). Fluorescence is commonly a decisive factor and arises from the coincidence of the excitation energy with electronic transitions. A way of avoiding this problem is to choose a longer wavelength (e.g. near-IR) with the compromise of lower spatial resolutions, or to move to UV wavelengths with the compromise of lower spectral resolutions and with the increased risk of sample damage occurring. Finally, the spatial resolution is especially important for point mapping, where a small laser spot diameter is often required.

Fig. 3 displays the Raman spectra of the Int16 membrane at the four different wavelengths both with and without baseline correction and normalisation. The baseline-corrected spectra were normalised to the intensity of the CF$_2$ stretch peak at 835 cm$^{-1}$ peak to aid visual comparison (the peak indicated by arrow). The enhancement in the 1612 cm$^{-1}$ peak when using the 457 nm laser is shown by the dashed box.

Fig. 4 The Raman spectra of (from bottom to top): the pristine ETFE, Int16, E1, E4, E16 and E72. Spectra were recorded with a 785 nm laser at 300 mW power (the only laser that could be used for all samples). The spectra were normalised to the intensity of ETFE peak at 835 cm$^{-1}$ to aid visual comparison. The arrows indicate the key peaks used for the Raman cross-section mapping experiments.

**Table 3** A summary of the pyrrolidinium-based RG-AEM(Cl$^-$)s synthesised

| RG-AEM(Cl$^-$)s | E1 | E4 | E16 | E72 |
|----------------|----|----|-----|-----|
| Grafting time/h | 1  | 4  | 16  | 72  |
| DoG (%)        | 18 | 44 | 99  | 155 |
| IEC (Cl$^-$ anion)/mmol g$^{-1}$ | 0.91 ± 0.05 | 1.70 ± 0.06 | 2.06 ± 0.02 | 2.36 ± 0.07 |
| WU (%)         | 8 ± 3 | 79 ± 5 | 104 ± 7 | 145 ± 6 |

Water$^a$ Number of H$_2$O molecules per Cl$^-$ anion [not to be confused with laser wavelength $\lambda$], calculated as: $\lambda_{water} = \text{WU} (%) / (100 \times 18.02 \times \text{IEC})$, where IEC is in mol g$^{-1}$.

---

The Raman spectra recorded using 457 and 532 nm lasers showed evidence (increase in opacity) of a high level of grafting. The use of 457 and 532 nm lasers led to moderate fluorescence, whereas the 633 and 785 nm lasers yielded spectra with minimal fluorescence. After baseline correction and normalisation, a number of Raman peaks were obtained with enhanced intensities with the 457 nm laser (especially the aromatic peak at 1612 cm$^{-1}$) indicating an intriguing but small Raman enhancement effect. Therefore, the 457 and 633 nm spot lasers were down-selected for the characterisation of pre-aminated ETFE-g-poly(VBC) membranes using Raman cross-sectional mapping. The 457 nm laser could be used up to 10% maximum power before sample damage occurs, while the 633 nm laser could be used at maximum power.

Regarding the (orange-brown coloured) RG-AEM(Cl$^-$)s, the Raman spectra recorded using 457 and 532 nm lasers showed
excessively strong fluorescence backgrounds that could not be reliably subtracted (see Fig. S3 in the ESI†). In contrast, the 785 nm laser yielded spectra with lower levels of baseline noise. Furthermore, it was impossible to record adequate spectra with the dark brown AE16 sample with all but the 785 nm laser due to the very high levels of fluorescence observed. The 785 nm laser proved to be the only available option that was suitable for recording spectral maps of both the pre- and post-alkali-treated RG-AEMs. On the downside, the use of this longer wavelength laser (with pinhole applied to make it a spot laser) gave weak signals and so longer acquisition times were required (even when using maximum power).

Chemical characterisation using Raman spectroscopy

The chemical structures of the pristine ETFE along with Int16, E1, E4, E16, and E72 were evaluated using Raman spectroscopy on the surfaces of the membranes (Fig. 4). The ETFE spectrum is dominated by the presence of two sharp peaks: CF$_2$ stretches at 835 cm$^{-1}$ (ref. 17) and CH$_2$ bends at 1444 cm$^{-1}$. The grafting of the VBC onto the peroxidated ETFE film introduced new peaks at 1612, 1266, and 1000 cm$^{-1}$, which correspond to the aromatic ring breathing, CH$_2$Cl deformations, and meta- and disubstituted benzene rings, respectively. The spectra of RG-AEM(Cl) reveals that the CH$_2$Cl peak at 1266 cm$^{-1}$ (Fig. 4) disappears indicating successful amination with the MPY. In addition, a new band at 900 cm$^{-1}$ was observed, which derives from the presence of cationic pyrrolidinium groups. All this data corroborates the Raman data previously reported for ETFE-MPY-based RG-AEMs.

Raman mapping of the pre-aminated ETFE-g-poly(VBC) membranes

Raman mapping was first performed on cross-sectional samples of the pre-aminated ETFE-g-poly(VBC) membranes to measure the through-plane distributions of the poly(VBC) grafts (Fig. 1). This was assessed using the peak area ratio $A_{1612}/A_{835}$: this is the area of the 1612 cm$^{-1}$ aromatic peak (deriving from the grafted poly(VBC) chains) vs. the area of the 835 cm$^{-1}$ peak (deriving from the backbone of the ETFE films). Peak areas were calculated by curve fitting using the Gaussian–Lorentzian function (after the spectra had been baseline corrected).

Fig. 5 shows the interpolated Raman maps obtained using the 457 nm laser for cross-sections of Int4 with the following stage (x–y) step sizes: 0.5 μm (over-sampling), 0.74 μm (par-sampling) and 1.5 μm (under-sampling). Recall, Fig. 2 schematically highlights the difference between these sampling modes. Note, these maps were produced from different cross-sections of Int4, and so the grafting distributions show a natural level of variation. For Int4, the distribution of poly(VBC) grafts shows more intensity at the surface and the middle of the membranes with bands of slightly lower intensity separating these regions. It is hypothesised that this is an effect of the diffusion of monomer. The monomer fronts diffuse from both surfaces of the ETFE film to finally meet in the middle, which increases the concentration of monomer at this location:
this leads to more rapid grafting for a period of time in the centre of the membrane. This effect was previously seen in a study that used propan-2-ol in the grafting medium, rather than the water used in this current study.

The data recorded using over-sampling demonstrated a higher fidelity Raman image, at the expense of experiment duration: map time was 220 min (compared to 30 min for the under-sampled image). A step size of 0.74 μm (par-sampling) was chosen for this study as it balances image fidelity with measurement time (map time of 110 min). Despite this, under-sampling may still be useful if a future necessity arose that required regularly conducted quality assurance measurements on a large number of samples (e.g. routine checks for homogeneous grafting of multiple batches of RG-AEM) or if measurements on larger cross-sectional areas of individual samples were desired: the fidelity of the under-sampled maps is still adequate for such tasks.

The Raman maps collected with the 633 nm laser are presented in Fig. S5 in the ESI: with this laser, faster mapping times of 20 min, 40 min, and 76 min were obtained with under-(1.50 μm step-size), par- (1.03 μm), and over-sampling (0.74 μm), respectively. These mapping times were quicker as the 633 nm laser could be applied at higher powers (at the sample): 20 mW compared to the maximum 2 mW that could be used with the 457 nm laser (due to sample damage at higher powers). Par-sampling (stage step size of 1.03 μm) was again down-selected to map the distribution of the grafted poly(VBC) chains with the 633 nm laser.

Table 4  Statistical data extracted from the Raman maps presented in Fig. 6 (calculated for the 95% confidence level) for the two most homogeneously grafted pre-aminated ETFE-g-poly(VBC) membranes

| ETFE-g-poly(VBC) | Int16 | Int72 |
|------------------|-------|-------|
| Number of measurements | 3738  | 3738  |
| Mean | 2.83  | 3.13  |
| Standard deviation | 0.13  | 0.16  |
| Relative standard deviation RSD (%) | 4.6   | 5.1   |
| Confidence intervals | ±0.005| ±0.005|
| D’Agostino–Pearson omnibus test: $^a$ $p$ | <0.0001| <0.0001|
| Passed normality test $\{\alpha = 0.05\}$? | No    | No    |
| Mann–Whitney $U$ test: $^b$ $p$ | <0.0001| Yes   |
| Means significantly different? | Yes   |       |

$^a$ Test for normality. $^b$ Non-parametric two-tailed Mann–Whitney $U$ test.
Fig. 6 shows cross-sectional maps (457 nm, par-sampled) obtained with the pre-aminated ETFE-g-poly(VBC) membranes synthesised using increasing grafting reaction times (Int1, Int4, Int16, and Int72). Fig. S4 in the ESI† gives the box and whisker plots that summarises this data for all the pre-aminated ETFE-g-poly(VBC) membranes studied. For additional clarity, Fig. 7 presents line-map data (457 nm) taken from the centres of the cross-sectional maps.

From this Raman data, it was observed that Int1 (with the lowest DoG = 18%) contained poly(VBC) grafted chains at the surface and a relatively ungrafted core. Increasing the grafting time to 4 h resulted in a wider through-plane distribution of grafting in the resulting membrane (Int4), but the grafting distribution was still not fully homogeneous (discussed previously). As the DoG approaches a value of 99% (Int16), a more homogeneous grafting of the poly(VBC) chains was achieved. The 4-probe (in-plane) chloride-anion conductivities of fully hydrated E1, E4, and E16 (measured using the method described in ref. 9) were 14 ± 3, 39 ± 2, and 49 ± 2 mS cm⁻¹ (n = 3), respectively, which is consistent with this Raman data.

Prolonging the grafting time to 72 h (Int72, DoG = 155%) led to higher grafting levels but this has detrimental consequences: e.g. E72 was observed to possess an excessive water content with λwater > 30 (Table 3), where λwater is the average number of H₂O molecules per anion-exchange site. Prior studies have shown that not all of the H₂O molecules are directly associated with the

---

**Fig. 8** Raman cross-sectional interpolated maps of E16 (top row) and AE16 (bottom row) recorded using the 785 nm laser (300 mW power, pinhole applied) with par-sampling. The colour scale represents (from left to right columns): A₉₀₀/A₈₃₅ (pyrrolidinium vs. ETFE); A₁₆₁₂/A₈₃₅ (benzene vs. ETFE); A₉₀₀/A₁₆₁₂ (pyrrolidinium vs. benzene). Each map is 30 × 65 μm in size (y-axis is the through-plane direction).
mechanical properties. RG-AEM swelling, which results in a weakening of the ionic groups; many H$_2$O molecules are “non-useful” in that they form domains of water aggregates that lead to excessive swelling, which results in a weakening of the mechanical properties.\textsuperscript{9} Table 4 gives the results of a statistical analysis comparing the poly(VBC) grafting levels between Int16 and Int72 using all of the $A_{1612}/A_{235}$ peak area ratios recorded in the production of their Raman maps in Fig. 6. Int72 has a (statistically significant) higher amount of grafting than Int16.

All of the above indicates that the grafting front mechanism is operating when VBC is grafted onto 50 $\mu$m thick ETFE films. Additional maps, acquired using the 633 nm laser on different cross-sections of the pre-aminated ETFE-g-poly(VBC) membranes (Fig. S6 in the ESI\textsuperscript{1}), confirm these observations. This Raman data is consistent with the bulk IEC values recorded for the final RG-AEM(Cl\textsuperscript{−})\textsuperscript{1} that were made by the amination of Int1, Int4, Int16, and Int72: IECs increase in the order E1 < E4 < E16 < E72. However, this Raman microscopy data provides additional evidence that the homogeneity of grafting is poor at low grafting reaction times.

**Raman studies on a RG-AEM(Cl\textsuperscript{−}): pre- and post-alkali-aged E16**

**Cross-sectional mapping.** The most important requirement for a high-performance RG-AEM is the homogeneous cross-sectional distribution of functional groups (Fig. 1).\textsuperscript{9} The following peak area ratios were used to characterise E16 (top row of Fig. 8), where $A_p$ is the area of the peak at $\gamma$/cm$^{-1}$ calculated by curve fitting using the Gaussian–Lorentzian function (after the spectra had been baseline corrected):

1. Ratio $= A_{900}/A_{835}$: pyrrolidinium groups vs. ETFE;
2. Ratio $= A_{1612}/A_{235}$: benzene rings vs. ETFE;
3. Ratio $= A_{900}/A_{1612}$: pyrrolidinium groups vs. benzene rings;

(1) and (2) represent the homogeneity of grafting and (3) represents the homogeneity of amination of the benzene rings.

For the pre-alkali-aged E16, it was evident that functionalisation was less homogeneous compared to the grafting studies on Int16. This may be due to segregation of the hydrophilic and hydrophobic phases in the RG-AEM sample that occurs after the amination process.\textsuperscript{††} The obvious “health warning” must also be kept in mind in that the apparent lower homogeneity may be down to sampling: the specific cross-sectional sample of E16 that was mapped could have been a low homogeneity section of the RG-AEM. Fig. 9 gives the box and whisker plots for data extracted from the top row of Fig. 8.

The data on the alkali degraded AE16 (bottom row of Fig. 8) clearly shows that degradation is evident when the E16 was aged in O$_2$-purged aqueous KOH (1.0 mol dm$^{-3}$) at 80 °C for 14 d. This data also shows that the degradation occurs throughout the cross-section of the RG-AEM. Fig. S7 in the ESI\textsuperscript{1} presents the box and whisker plots comparing the peak area ratios for E16 and AE16 (extracted from the Raman cross-sectional map data presented in Fig. 8). The larger variances seen in the AE16 data is due to the difficulty in reliably fitting all of the spectral data due to the significant fluorescence backgrounds encountered. Statistical analysis of the peak area ratio $A_{900}/A_{835}$ data (the ratio expected to change the most on RG-AEM degradation) is summarised in Table 5. This analysis shows that this peak area ratio data is significantly different for AE16 compared to E16 due to alkali degradation. The degradation observed was due to both: (1) loss of the positively-charged pyrrolidinium functionality (Scheme 2: nucleophilic hydroxide attack positions A, B, and C and Hofmann elimination attack position E); (2) loss of grafted benzene rings from the RG-AEM, complete with pyrrolidinium groups (Scheme 2: hydroxide attack position D). This is consistent with the observations reported in recent studies on both ETFE- and LDPE-based RG-AEMs.\textsuperscript{6,14}

\begin{table}
\caption{Statistical data for the peak area ratio $A_{900}/A_{835}$ data extracted from the Raman cross-sectional maps (785 nm laser) recorded on pre- and post-alkali-aged E16 (calculated for the 95% confidence level).}
\begin{tabular}{lccc}
\hline
 & E16 & AE16 \\
\hline
Number of measurements & 1248 & 1248 \\
Mean & 0.522 & 0.415 \\
Standard deviation & 0.081 & 0.076 \\
RSD (%) & 16 & 18 \\
Confidence intervals & ±0.005 & ±0.004 \\
D’Agostino–Pearson omnibus test$^a$ $p =$ & <0.0001 & <0.0001 \\
Passed normality test ($\alpha = 0.05$)? & No & No \\
Mann–Whitney U test$^b$ $p =$ & <0.0001 & \\
Means significantly different? & Yes & \\
\hline
\end{tabular}
\end{table}

$^a$ Test for normality. $^b$ Non-parametric two-tailed Mann–Whitney U test.

†† Note: with the use of the IR (785 nm) laser, the scale-bar for the $A_{1612}/A_{235}$ peak area ratio (middle column in Fig. 8) is compressed to the range 0–2, compared to the range 0–4 in Fig. 6, that was required to accommodate the higher ratios observed for Int72 with the use of the blue (457 nm) laser.
Surface Raman data. So far, this study has considered the cross-sectional (through-plane) distributions of functionality. However, the homogeneity of functionalisation across the surfaces of the RG-AEMs should also be evaluated (Fig. 10). Detailed mapping over large areas of RG-AEM surfaces is not viable for routine measurements. A more practical method is to record a number of spectra on different parts of the surface of a RG-AEM sample. In this study, Raman spectra (785 nm, 300 mW, pinhole applied) were recorded at 25 sampling points on the surfaces of E16 and AE16 (recorded as a 5 × 5 grid over ca. 1 cm² sized samples). Each spectrum was analysed for the three peak area ratios described above.

Fig. 11 shows the results recorded on the surface of a sample of E16. This data shows good homogeneity with RSDs < 10% for all three peak area ratios. However, this Raman data shows that the surface analysis of E16 yields larger area ratios A_900/A_835 and A_900/A_835 that are related to the level of grafting, compared to the cross-sectional analysis (Fig. 9). This suggests there is an excess of grafting directly at the surface of E16 (where the laser only penetrates a couple of µm into the membrane when focused on the surface). Therefore, for RG-AEMs, we strongly recommend that surface data should always be collected alongside cross-sectional mapping data (to obtain the fullest possible picture of the distribution of the grafted chains).

Fig. S8 in the ESI† presents the box and whisker plots comparing the peak area ratios from the surface analyses of E16 and AE16. Statistical data for the peak area ratio A_900/A_835 is summarised in Table 6. This analysis again shows that this diagnostic peak area ratio is significantly different for AE16 compared to E16 due to alkali degradation. There was a 25% loss of this peak area ratio on surface degradation, while only a 21% loss was observed in the analysis of the cross-sectional data. It is not unexpected that degradation would be more severe at the surface of RG-AEMs. AE16 possessed an IEC of 1.65 ± 0.17 mmol g⁻¹ (n = 3), which was 20% lower than the 2.06 ± 0.02 mmol g⁻¹ recorded for the pre-alkali treated E16. This (bulk) IEC data directly correlates with the loss of the peak area ratio A_900/A_835 in the Raman cross-sectional data.

Conclusions

This study highlights the usefulness of using Raman spectroscopy in the study of radiation-grafted membranes (anion-exchange membranes in this case). This technique is especially helpful for the study of the homogeneity of the cross-sectional grafting (on the micro-scale). This technique has the potential to be used as a quality assurance tool regarding the grafting process (with cross-sectional mapping times of <1 h being possible with under-sampling). This technique can also be useful for the study of the homogeneity of any post-grafting process (with cross-sectional mapping times of <1 h being possible with under-sampling). This technique can also be useful for the study of the homogeneity of any post-grafting process (with cross-sectional mapping times of <1 h being possible with under-sampling).
functionalisation process (e.g. amination) as well as for the (post-mortem) investigation of membrane degradation processes. For radiation-grafted membranes, the analysis of the surfaces must always be conducted in addition to any cross-sectional analysis, in order to obtain the whole picture regarding the distribution of membrane functionality (and membrane degradation).

For the radiation-grafted anion-exchange membranes (RG-AEM), made from 50 μm thick ETFE films in this investigation, a vinylbenzyl grafting time of 16 h appears to be optimal (with the specific synthesis conditions used). The Raman data is consistent with the operation of the grafting front mechanism. Amination with 1-methylpyrrolidine was homogeneous and yielded RG-AEMs with ion-exchange capacities (IEC) of >2.0 mmol g⁻¹. There was an excess of grafting at the surfaces of the RG-AEMs (at least in the first few μm). The level of degradation, measured using Raman spectro-microscopy of the RG-AEM cross-sections after ageing in hot aqueous alkali, matched the loss of IEC; however, the high levels of photolysis that occurs with the (dark-brown-coloured) alkali degraded membranes, while the 785 nm laser wavelength was the only option that could be used for the study of the pre- and post-alkali degraded RG-AEMs.

Author contributions

Dr Bance-Soualhi supervised the student and conducted a number of the experiments (surface maps and degradation data), as well as making a major contribution to authoring of this article. Dr Crean and Prof Varcoe are the principal investigators of the EPSRC grants who contributed to the formulation of this article and scientific interpretation of the data.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was funded by the UK’s Engineering and Physical Sciences Research Council (EPSRC) with resources provided by grants EP/M005933/1, EP/M014371/1, and EP/M022749/1 (the latter providing the capital funding that allowed for the purchase of the Renishaw Raman microscope system). Renishaw are thanked for their technical assistance related to our familiarisation with the instrument. Wai Hin Lee thanks the University of Surrey for providing additional funding: the bulk of the results in this paper was from his undergraduate final year project. All data is freely available with no restrictions: details on how to access this data can be found at DOI: 10.15126/surreydata.00842458.

References

1 T. R. Dargaville, G. A. George, D. J. T. Hill and A. K. Whittaker, Prog. Polym. Sci., 2003, 28, 1355; M. M. Nasef, S. A. Gürsel, D. Karabelli and O. Güven, Prog. Polym. Sci., 2016, 63, 1.
2 M. M. Nasef, Chem. Rev., 2014, 114, 12278; L. Gubler, Adv. Energy Mater., 2014, 4, 201300827; A. Albert, T. Lochner, T. J. Schmidt and L. Gubler, ACS Appl. Mater. Interfaces, 2016, 8, 15297.
3 J. Pan, C. Chen, L. Zhuang and J. Lu, Acc. Chem. Res., 2012, 45, 473; J. R. Varcoe, P. Atanassov, D. R. Dekel, A. M. Herring, M. A. Hickner, P. A. Kohl, A. R. Kucernak, W. E. Mustain, K. Nijmeijer, K. Scott, T. Xu and L. Zhuang, Energy Environ. Sci., 2014, 7, 3135; K. D. Kreurer, Chem. Mater., 2014, 26, 361; M. A. Hickner, Interface, 2017, 26, 69; T. J. Omasta, L. Wang, X. Peng, C. A. Lewis, J. R. Varcoe and W. E. Mustain, J. Power Sources, 2017, DOI: 10.1016/j.jpowsour.2017.05.006.
4 N. Walsby, M. Paronen, J. Juhanjoa and F. Sundholm, J. Polym. Sci., Part A: Polym. Chem., 2000, 38, 1512; J. R. Varcoe, R. C. T. Slade, E. L. H. Yee, S. D. Poynton, D. J. Driscoll and D. C. Apperley, Chem. Mater., 2007, 19, 2688; O. I. Deavin, S. Murphy, A. L. Ong, S. D. Poynton, R. Zeng, H. Herman and J. R. Varcoe, Energy Environ. Sci., 2012, 5, 8584; B.-M. Ko, J.-Y. Sohn and J. Shin, Polymer, 2012, 53, 4652; M. Mamlouk, J. A. Horsfall, C. Williams and K. Scott, Int. J. Hydrogen Energy, 2012, 37, 11912; T. Zhou, R. Shao, S. Chen, X. He, J. Qiao and J. Zhang, J. Power Sources, 2015, 293, 946; R. Espiritu, M. Mamlouk and K. Scott, Int. J. Hydrogen Energy, 2016, 41, 1120; Y. Zhao, K. Yoshimura, H. Shishitani, S. Yamaguchi, H. Tanaka, S. Koizumi, N. Szekely, A. Radulescu, D. Richtere and Y. Maekawa, Soft Matter, 2016, 12, 1567.
5 J. R. Varcoe, R. C. T. Slade, G. L. Wright and Y. Chen, J. Phys. Chem. B, 2006, 110, 21041; S. F. Lu, J. Pan, A. B. Huang, L. Zhuang and J. T. Lu, Proc. Natl. Acad. Sci. U. S. A., 2008, 105, 20611; C. A. Hancock, A. Ong, P. R. Slater and J. R. Varcoe, J. Mater. Chem. A, 2014, 2, 3047; H. A. Miller, A. Lavacchi, F. Vizza, M. Marelli, F. Di Benedetto, F. D. I. Acapito, Y. Paska, M. Page and D. R. Dekel, Angew. Chem., Int. Ed., 2016, 55, 6004; H. A. Miller, F. Vizza, M. Marelli, A. Zadick, L. Dubau, M. Chatenet, S. Geiger, S. Cherevko, H. Doan, R. K. Pavlicev, S. Mukerjee and D. R. Dekel, Nano Energy, 2017, 33, 293–305; H. Wang and H. Abruña, J. Am. Chem. Soc., 2017, 139, 6807; L. Wang, J. J. Brink, Y. Liu, A. M. Herring, J. Ponce-González, D. K. Whelligan and J. R. Varcoe, Energy Environ. Sci., 2017, DOI: 10.1039/C7EE02053H.
6 J. Ponce-González, D. K. Whelligan, L. Q. Wang, R. Bance-Soualhi, Y. Wang, Y. Q. Peng, H. Q. Peng, D. C. Apperley, H. N. Sarode, T. P. Pandey, A. G. Vivek, S. Seifert, A. M. Herring, L. Zhuang and J. R. Varcoe, Energy Environ. Sci., 2016, 9, 3724.
7 H.-P. Brack, H. G. Bührer, L. Bonorand and G. G. Scherer, J. Mater. Chem., 2000, 10, 1795; M. M. Nasef and E.-S.
A. Hegazy, Prog. Polym. Sci., 2004, 29, 499; Y. Buchmüller, A. Wokaun and L. Gubler, Fuel cells, 2013, 13, 1177.
8 V. Sproll, T. J. Schmidt and L. Gubler, Polym. Int., 2016, 65, 174.
9 L. Q. Wang, E. Magliocca, E. L. Cunningham, W. E. Mustain, S. D. Poynton, R. Escudero-Cid, M. M. Nasef, J. Ponce-González, R. Bance-Soualhi, R. C. T. Slade, D. K. Whellung and J. R. Varcoe, Green Chem., 2017, 19, 831.
10 S. Holmberg, J. H. Näsman and F. Sundholm, Polym. Adv. Technol., 1998, 9, 121; L. I. Şanli and S. A. Gürsel, J. Appl. Polym. Sci., 2011, 120, 2313.
11 F. Cardona, G. A. George, D. J. T. Hill, F. Rousoul and J. Maeji, Macromolecules, 2002, 35, 355; F. Cardona, G. A. George, D. J. T. Hill and S. Perera, J. Polym. Sci., Part A: Polym. Chem., 2002, 40, 3191; H. Herman, R. C. T. Slade and J. R. Varcoe, J. Membr. Sci., 2003, 218, 147.
12 J. E. Katon, Vib. Spectrosc., 1994, 7, 201; A. Chaplin, I. Hamerton, H. Herman, A. K. Mudhar and S. J. Shaw, Polymer, 2000, 41, 3945; E. Widjaja and M. Garland, J. Raman Spectrosc., 2012, 43, 828; R. Zeng, J. Handsel, S. D. Poynton, A. J. Roberts, R. C. T. Slade, H. Herman, D. C. Apperley and J. R. Varcoe, Energy Environ. Sci., 2011, 4, 4925; D. M. Hillman, S. H. Stephens, S. D. Poynton, S. Murphy, A. Ong and J. R. Varcoe, J. Mater. Chem. A, 2013, 1, 1018; R. Zeis, Beilstein J. Nanotechnol., 2015, 6, 68; S. Mukherjee and A. Gowen, Anal. Chim. Acta, 2015, 895, 12; C. Cardell and I. Guerra, Trends Anal. Chem., 2016, 77, 156; T. R. Pozegic, J. V. Anguita, I. Hamerton, K. D. G. I. Jayawardena, J.-S. Chen, V. Stolojan, P. Ballochi, R. Walsh and S. R. P. Silva, Sci. Rep., 2016, 6, 37334.
13 A. Zoubir, Raman Imaging: Techniques and Applications, Springer, Heidelberg NY (USA), 2012.
14 R. Espiritu, B. T. Golding, K. Scott and M. Mamlouk, J. Mater. Chem. A, 2017, 5, 1248; R. Espiritu, B. T. Golding, K. Scott and M. Mamlouk, J. Power Sources, 2017, DOI: 10.1016/j.jpowsour.2017.07.074.
15 E. Smith and G. Dent, Modern Raman Spectroscopy: A practical Approach, Wiley, Hoboken NJ (USA), 2005.
16 R. L. McCreery, Raman Spectroscopy for Chemical Analysis, John Wiley & Sons, New York (USA), 2005.
17 G. Galleja, A. Houdayer, S. Etienne-calas, D. Bourgogne, V. Flaud, G. Silly, S. Shibahara, A. Takahara, A. Jourdan, A. Hamwi and B. Ameduri, J. Polym. Sci., Part A: Polym. Chem., 2011, 49, 1517.
18 P. J. Larkin, Infrared and Raman spectroscopy: Principles and spectral interpretation, Elsevier, Amsterdam, 2011.
19 M. A. Vandiver, B. R. Caire, J. R. Carver, K. Waldrop, M. R. Hibbs, J. R. Varcoe, A. M. Herrin and M. W. Liberatore, J. Electrochem. Soc., 2014, 161, H677; T. P. Pandey, A. M. Maes, H. N. Sarode, B. D Peters, S. Lavinia, K. Vezzu, Y. Yang, S. Poynton, J. R. Varcoe, S. Seifert, M. Liberatore, V. Di Noto and A. Herrin, Phys. Chem. Chem. Phys., 2015, 17, 4367.