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# Effect of H-bonding on network junction and macroscopic elastomer properties in photocured polyacrylate films<sup>†</sup>

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Two series of polyacrylate films with different H-bonding capable monoacrylate copolymers were synthesized by UV-initiated photo-polymerization. Detailed IR analyses on cured samples show that networks made from structurally distinguishable mono-acrylate have significantly different extent of H-bonded monoacrylate present. We found that this is induced by differences in the relative reaction rate between homo-polymerization and copolymerization of the relevant monoacrylates and crosslinker (PEGDA). Although 'pre-organization' of H-bond capable monoacrylates was observed for those tested formulations prior to crosslinking, which should significantly increase the homo-polymerization rate, it was found that the co-polymerization rate was also significantly increased. Furthermore, DQ NMR analyses on these networks pinpoints the presence of H-bonding clusters, which decreases mobility of chain segments near the H-bonding functional groups in the monoacrylate sidechains (dangling chain segments of monoacrylate). However, the overall network topology does not alter significantly upon forming H-bonding clusters in the monoacrylate sidechains. Finally it was also found that larger H-bonding clusters increase the apparent network junction functionality. These findings are discussed in the context of utilizing the H-bonding capable monoacrylates to tailor the microscopic topological properties as well as macroscopic physical properties of photocured polyacrylate polymer films.

## 1. Introduction

Polyacrylate-based films have been widely used for biomedical applications ranging from tissue engineering to drug delivery due to their excellent biocompatibility and substantial durability. Among the available methods for producing polyacrylates, photo-initiated curing is the most popular approach implemented in industrial manufacture due to its ease of application and fast production.<sup>1–4</sup> Modification of photo-cured crosslinked networks can be easily achieved by copolymerizing functionalized (*e.g.* hydrogen bonding capable) mono-vinyl monomers with divinyl

crosslinkers, enabling tailoring of their macroscopic properties. However, as a consequence of the uncontrollable nature of free radical polymerization, the resulting materials often have complicated microscopic structures and network topologies which in turn strongly affects their characteristics in bulk.<sup>5,6</sup> Hence, insights into the effects of including functionalized monoacrylates on the local structure and chain dynamics within photo-cured polyacrylate networks are of fundamental interest for designing new functional materials.

H-bond capable monomers have been frequently introduced into polymer networks to alter their apparent network density in order to change their macroscopic physical properties. Burattini *et al.* constructed a polyimide based tough elastomeric material with self-healable properties arising from H-bonding and  $\pi$ - $\pi$  stacking interactions between di-imide and pyrenyl groups.<sup>7</sup> Bhattarai *et al.* created a chitosan-based tunable polymer network for localized drug delivery where the release mechanism was controlled by the size of domains arising due to H-bonding groups.<sup>8</sup> In general, the glass transition temperature ( $T_g$ ),<sup>9,10</sup> network morphology,<sup>11,12</sup> viscoelastic property<sup>13</sup> as well as the mechanical strength<sup>14</sup> of modified networks have been shown to be strongly influenced by the presence of H-bonding, both directly by altering chain

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interactions and indirectly through cluster formation. UV-curable (meth)acrylate monomers with H-bonding capability typically involve hydroxyl (-OH) or amine (-NH<sub>2</sub>) groups. This seemingly simple modification has been found to induce significant changes in the bulk properties. Peppas et al. analyzed a series of poly(ethylene glycol) based polyacrylate networks and found H-bonding strongly affects solute diffusion through the networks.<sup>15</sup> Lewis et al. described a strong correlation between monomer H-bonding strength and the viscoelastic properties of loosely crosslinked H-bond capable polyacrylates.<sup>13</sup> We have previously described how H-bonding can induce changes in mesh size, as assessed by NMR diffusometry and DSC thermoporomety.<sup>16</sup> Hence, while H-bonding capable monomers can in principle be used to regulate the final properties, the potential to exploit this approach to provide materials with tunable properties for biomedical application is limited by the lack of understanding into how changes at a molecular level can induce local and intermediate-range structural modifications that determine the emergent physical properties of the cured polymer networks.

Multiple techniques are used in combination to probe the complex chemical structure and molecular dynamics in crosslinked polymeric network. Nuclear magnetic resonance (NMR) owing to its non-destructive and informative nature has been widely applied in analyzing these systems.<sup>17</sup> While the conventional NMR spectral analysis provides highly selective structure determination,18-20 NMR relaxometry provides further insights into polymer chain dynamics.<sup>21,22</sup>  $T_2$  relaxometry is more commonly used for polymeric materials due to greater sensitivity to small differences<sup>23,24</sup> in crosslink density and chain mobility. The ongoing developments in this important field have been reviewed on several occasions.<sup>25,26</sup> By implementing Hahn-echo-based  $T_2$  analysis, Litvinov et al. successfully evaluated crosslink density of UV-cured PEG-DA networks.6 Recently, application of high resolution multiple quantum, multi-quantum (MQ) NMR, techniques to polymeric materials

has been implemented, with chemical shift-specified residual dipolar coupling constants,  $D_{res}$ , now obtainable through accepted experimental and regularization procedure.<sup>27,28</sup> We have previously shown that  $T_2$  and MQ techniques can be particularly useful when combined for extracting information about network crosslink density and its relation to chain dynamics, as modulated by functional group type and relative composition.<sup>5</sup>

In this study two series of UV-polymerized poly(ethylene glycol)-based polyacrylate networks were synthesized by copolymerizing the diacrylate (PEGDA-700) with monoacrylates, bearing either terminal hydroxy (t-OH) or non-terminal amide (nt-NH) H-bond capable functional groups on their side chains (see Table 1) forming networks with multifunctional acrylic network junctions interconnected by PEG chains with a fraction of dangling monoacrylate sidechains. The weight percentage (w%) of the PEGDA-700 in the curing formulation was systematically varied to evaluate the effect of gradually increasing the H-bonding capability on the networks. After confirming high double bond conversion by FTIR and <sup>13</sup>C NMR spectroscopy, the networks were studied by a series of analytical techniques. The H-bonded fractions were estimated from the O-H and N-H IR stretching bands in detailed FTIR analyses, while the polymer chain dynamics were evaluated by dynamic mechanical analyses (DMA), low field <sup>1</sup>H NMR  $T_2$  relaxometry and high field DQ NMR. We demonstrate how strikingly different properties can emerge due to the formation of clusters of H-bonded monoarylate units, illustrated schematically in Fig. 1, associated with the sidechains, whose sizes are regulated by the relative ratio between homo- and co-polymerization of the H-bonding capable monomers. The effects of H-bonding clusters on network chain mobility were further examined using chemicalshift specified residual dipolar coupling constants, D<sub>res</sub>, extracted from high resolution DQ NMR. Finally, the impact of H-bonding on the curing kinetics, resulting in differences in cluster formation, and on the network chain dynamics of H-bonding capable side chains and polyacrylate network junctions are discussed.

Table 1         Details of chemicals used for network formation								
Compounds	Structure	$M_{ m w} ({ m g \ mol}^{-1})$						
Poly(ethylene glycol) diacrylate (PEGDA)	H <sub>2</sub> C H <sub>2</sub> C H <sub>2</sub>	700						
4-Hydroxyethyl acrylate (HEA)	H <sub>2</sub> C OH	116.1						
2-[[(Butylamino)carbonyl]oxy]ethyl acrylate (BAC)		215.5						
Hexyl Acrylate (HA) <sup><i>a</i></sup>	H <sub>2</sub> C OCH <sub>2</sub> (CH <sub>2</sub> ) <sub>4</sub> CH <sub>3</sub>	156.2						
2,2-Dimethoxy-2-phenylacetophenone (DMPA)	H <sub>3</sub> CO OCH <sub>3</sub>	256.3						

 $^{a}$  HA is included here as data<sup>5</sup> for networks formed using it are used in the comparisons analyses below.



Fig. 1 Illustration of the formation of microscopic H-bonding clusters during bulk UV curing of a polyacrylate networks.

That comparative discussion is facilitated by the inclusion of relevant data from our earlier NMR and DMA study<sup>5</sup> on similar networks that were formed using the hexyl acrylate (HA) monomer which contain no H-bond capable groups.

## 2. Experimental

#### 2.1. Sample preparation

Polyethylene glycol-700-diacrylate (PEGDA-700,  $M_{\rm n}$  700 g mol<sup>-1</sup>,  $n_{\rm av}$  16, PDI: 1.05), Hexyl acrylate (HA), hydroxyethyl acrylate (HEA), 2-[[(butylamino)carbonyl]oxy]ethyl acrylate (BAC); photoinitiator: 2,2-dimethoxy-2-phenylacetophenone (DMPA) were used for the syntheses of networks. All the chemicals were obtained from Sigma Aldrich and used without any further purification.

Networks were prepared by bulk UV-curing of the chemicals (typically 5 g of mixture) with the proportion described in Table 2 (Photo initiator: 10–20 mg 2,2-dimethoxy-2-phenylace-tophenone (DMPA)). The formulation was poured into a Petri dish, and then the Petri dish was passed through the UV-rig (Heraeus Noblelight Fusion UV Inc.'s F300S model, with I300MB irradiator which has a lamp power of 300 W in<sup>-2</sup> (120 W cm<sup>-2</sup>), 1800 W total) 30 times. Afterwards, the cured film was inverted, and was then passed through the UV-rig a further 30 times. The rim of the resulting film was cut off,

and the remaining part with a thickness around 1 mm was used for all the tests.

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The double bond conversions of the films (which describes the completeness of crosslinking reaction) was measured by FTIR, and molecular weight of the hydrolyzed acrylate backbone chains was determined by chromatographic analyses.<sup>29</sup> Results of network characterization are summarized in Table 2.

#### 2.2. ATR-FTIR analyses

All the attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR) analyses were performed on a Bruker Alpha-P FTIR spectrometer for the original uncured acrylate mixture and the cured polyacrylate films. The spectrometer consisted of a diamond crystal ATR with a KBr beamsplitter. The scanning range of the FT-IR used for this analysis was ranged from  $360-4000 \text{ cm}^{-1}$  with a resolution of 2 cm<sup>-1</sup>. 64 scans were collected for each sample in total. Examples of spectral deconvolution can be found in Fig. S10 (ESI†).

#### 2.3. Mechanical analyses

The glass transition temperature and storage modulus of all samples were measured (under elongation without pre-stretching) with a TA DMA Q800 Analyzer at a frequency of 1 Hz. Dog-bone test sample bars of 5 mm width along the working region, and 15 mm length were cut from cured films. The thickness of the test bars were measured with a calibrated micrometer. The DMA test were initiated at -90 °C and the temperature was gradually increased up to 150 °C with a ramp speed of 3 °C min<sup>-1</sup>. The oscillatory force was 13.28 N. The typical displacement (extension) was around 4–8 µm (depending on sample).

#### 2.4. Solid state NMR analyses of network structure

Solid-state magic angle spinning (MAS)  $^{13}$ C NMR spectra were recorded using a Bruker Avance III 600 MHz instrument equipped with a solid standard-bore Magic Angle Spinning (SBMAS) probe. A 4 mm MAS rotor (of 80  $\mu$ L sample volume)

 Table 2
 Sample formulations and characterization of PEG-HEA and PEG-BAC networks by FTIR, dynamic mechanical analyses (DMA), gel permeation chromatography (GPC), NMR relaxometry and DQ NMR analyses

No.	МА Туре	$w\%_{MA}{}^{a}(\%)$	Conversion <sup><math>b</math></sup> (%)	DMA <sup>c</sup>		$\operatorname{GPC}^d$			$T_2$ relaxometry <sup>e</sup>		DQ NMR <sup>f</sup>
				$T_{\rm g}$ (K)	E' (MPa)	$M_n$ (PAA) (kDa)	PDI (PAA)	n(PAA)	$T_{2,\mathrm{av}}^{\mathrm{dry}}\left(\mu s\right)$	$T_{2,\mathrm{av}}^{\mathrm{sw}}(\mathrm{\mu s})$	$D_{\rm avg}/2\pi$ (kHz)
1	HEA	10	$91\pm5$	246	28.3	9.5	2.2	131	154	160	2.26
2	HEA	30	$95\pm5$	254	20.8	10.5	2.1	146	174	188	1.83
3	HEA	50	$92\pm5$	263	13.5	11.8	2.5	164	212	237	1.51
4	HEA	70	$91\pm5$	274	7.5	12.7	2.7	177	261	319	0.99
5	HEA	90	$90\pm5$	289	2.8	11.3	2.6	157	324	565	0.43
6	BAC	10	$92\pm5$	247	27.1	11.2	2.2	155	150	164	2.20
7	BAC	30	$93 \pm 5$	252	19.5	10.2	2.7	142	172	188	1.71
8	BAC	50	$91\pm5$	260	12.5	12.8	2.0	178	226	247	1.43
9	BAC	70	$91\pm5$	268	7.0	11.5	2.6	160	332	418	0.89
10	BAC	90	$89 \pm 5$	274	2.5	9.9	2.1	138	574	697	0.42

<sup>*a*</sup> The remaining *w*% in each case is PEGDA, summing to 100%. <sup>*b*</sup> Determined by FTIR analyses of the C—C–H stretching band (810 cm<sup>-1</sup>). <sup>*c*</sup> E' is determined at 50 °C by DMA. <sup>*d*</sup> Polyacrylic acid (PAA) derived from the backbone component following network hydrolysis, was characterized by gel permeation chromatography;  $M_n$ , number average molecular weight; PD, polydispersity; *n*(PAA), degree of polymerization. The error in  $M_n$  is estimated at *c*. 10%. <sup>*e*</sup>  $T_{2,av}^{diy}$ ,  $T_{2,av}^{sw}$ , average  $T_2$  relaxation for dry and swollen samples at 70 °C, respectively; the standard deviation is *c*. 10%. <sup>*f*</sup>  $D_{avg}$ , average residual dipolar coupling constant recorded at 70 °C, the standard deviation is again estimated at *c*. 10%.

was used with a spinning rate of 12 kHz. Inverse-gated decoupling and cross-polarization (CP) pulse sequences were applied. The 90° pulse length was calibrated to be 2  $\mu s$ . The spectra were recorded with two different cross-polarization times ( $\tau_{cp}$ ) of 0.5 and 10 ms, which were selected to distinguish between rigid and soft parts of the material. The data recorded using 10 ms is shown in ESI.†

#### 2.5. <sup>1</sup>H NMR T<sub>2</sub> relaxation test

The <sup>1</sup>H magnetization spin–spin relaxation ( $T_2$ ) times of the networks were measured using a Bruker Minispec MQ20 spectrometer operating at 20 MHz. The  $T_2$  relaxation decay was measured with the Hahn-echo (HE) pulse sequence,  $90_x^{\circ}-\tau_{\rm HE}-180_y^{\circ}-\tau_{\rm HE}-$ (acquisition), where  $\tau_{\rm HE} \ge 35 \ \mu s$ . In the HE experiment, an echo signal is formed after the second pulse in the HE with a maximum at time  $t = 2\tau_{\rm HE}$  after the first  $90^{\circ}$  pulse. By varying  $\tau_{\rm HE}$ , the amplitude of the transverse magnetization, A(t), is measured as a function of time t. The <sup>1</sup>H NMR  $T_2$  relaxation experiments in this study were performed at 70 °C, which is >60 °C above the  $T_g$  value of the networks. For experiments on swollen samples, samples were swollen in the deuterated solvent  $C_2D_2Cl_4$  overnight before the measurement. Detailed description about the  $T_2$  fitting can be found in ESI<sup>†</sup> as well as our previous work.<sup>5</sup>

## 2.6. Homonuclear <sup>1</sup>H-<sup>1</sup>H double quantum-single quantum (DQ-SQ) correlation NMR

The 2D <sup>1</sup>H double-quantum (DQ) correlation experiments were carried out on a Varian VNMRS 850 MHz operating at a <sup>1</sup>H Larmor frequency of 849.71 MHz equipped with a 1.6 mm triple resonance probe. The MAS frequency was set to 40 kHz and the sample temperature to 70  $^{\circ}$ C. The 90 $^{\circ}$  pulse length was calibrated to 2 µs. The data for all network samples were acquired using a recently published improved version of the Back-to-Back (BaBa) sequence, BaBa-xy16, for broadband homonuclear DQ recoupling.<sup>28</sup> The corresponding coupling constant distributions are obtained by the evaluation of the normalized build-up curves with *ftikreg* as describe in our previous work.<sup>30</sup> The extracted residual dipolar coupling constants (D<sub>res</sub> values) depend on the amplitude of the segmental dynamics, e.g. bond vector fluctuations in a large flexible molecule regarding to the angle between internuclear vector and the symmetry axis of motion.<sup>28</sup>

#### 3. Results

#### 3.1. Network chemical structure characterization

Two series of networks containing either *t*-OH (PEG-HEA) or *nt*-NH (PEG-BAC) groups with increasing weight fraction of monoacrylate (MA),  $w\%_{MA}$  ( $w\%_{MA} = 100 \times wt_{MA}/(wt_{MA} + wt_{PEG})$ ,  $wt_{MA}$  is the weight of monoacrylate, and  $wt_{PEG}$  the weight of PEGDA), were prepared as described in Section 2. To confirm complete crosslinking polymerization of the formulations, <sup>13</sup>C solid state NMR spectra and IR spectra were recorded for all the samples. <sup>1</sup>H solution-state NMR was used for analysis of

the sol fractions. The formulations and primary characterization for all samples in the two series are shown in Table 2, spectra and details of the analyses are included in the ESI† as well as in our previous publication.<sup>5</sup> The double bond conversions for all the samples are found to be very high (>90%), and the amount of sol fraction determined by extraction was below 1 *w*%. No significant side reactions occurred during the curing procedure for either series, as confirmed by <sup>13</sup>C solid state NMR analysis.

According to the procedure described in our previous publication, 5,29  $M_n$  and PDI values of the polyacrylic acid (PAA) present in the hydrolyzed networks were obtained by size exclusion chromatography (SEC) and listed in Table 2 (details of the analyses are in ESI<sup>†</sup>). PEG fraction was measured with mass spectrometry detection, assuming equal MS sensitivity, PEG has a  $M_n$  value of 488 Da. The molecular weight  $(M_n)$  of the acrylate backbone chains, a characteristic parameter for UV-cured PEGDA-based networks,<sup>29</sup> was also measured. It was found there is no statistically significant change across the network series, given the estimated error of  $\sim 10\%$ , and the values are similar to those we reported when using a non H-bonding monomer.<sup>5</sup> In general, these values are found to depend on the polymerization rate and on the bimolecular termination rate,<sup>31</sup> which is apparently similar for the different samples.

#### 3.2. Spectral analysis of hydrogen bonding in the networks

Several methods can demonstrate the presence of H-bonding in polymeric networks, with FT-IR most commonly used owing to its convenience.<sup>32,33</sup> H-bonding usually significantly reduces the O-H (or N-H) bond strength, lowering the force constant and shifting the stretching modes to lower wavenumbers. For the PEG-HEA network series, it is observed that the O-H stretching band energy decreases progressively with increase in  $w\%_{HEA}$  (Fig. 2a). Typically, the stretching band of free hydroxyl groups connected to an aliphatic chain is found well above 3520 cm<sup>-1</sup> within the blue area indicated.<sup>34</sup> As illustrated in Fig. 2d, we assign the gradual shift to lower wavenumber and broadening of this band with increasing  $w_{\text{HEA}}$  to a greater fraction of OH groups residing within increasingly large H-bonding clusters, as opposed to those at their interfaces.<sup>35</sup> We will return to the nature of these clusters in Section 4. Furthermore, across this series of samples the effect of curing on this region of the spectra is minimal as shown by the IR analyses of pre-curing formulation (Fig. S9, ESI†), demonstrating that most, if not all, of the HEA monomers participate in H-bonding before curing. This observation suggests that there is significant spatial re-arrangement of HEA monomers in the formulation, and curing does not alter this re-arrangement.

A different situation is observed for the PEG-BAC network series. The free N–H stretching band of secondary amide groups is normally observed above 3450 cm<sup>-1</sup>. The amide groups in the network series were found to gradually shift with increasing  $w%_{BAC}$  from free to H-bonded forms.<sup>36,37</sup> For the pre-curing samples the N–H stretching feature corresponds to the H-bonded form only (see Fig. S9, ESI<sup>†</sup>) and the spectra are



**Fig. 2** Selected regions of the IR spectra of; (a) PEG-HEA networks; (b) PEG-BAC networks; (c) extent of H-bonding as a function of  $w_{MA}^{*}$ , the molar ratio of H-bonding capable monomer HEA or BAC in the formulations; (d) schematic representation of H-bonded monoacrylate clusters (*t*-OH case); (i) on the left a larger cluster with lower fraction of interfacial H-bonding groups; (ii) right, a smaller block with a higher of fraction of interfacial groups. Interfacial H-bonded OH groups are shown in light green, this inside cluster are in blue, and a 'free' OH group is shown in brown. Red line indicates PEG chain, orange line the polyacrylate main chain, and dark green lines the monoacrylate side chain.

identical for all PEG-BAC pre-curing formulations, while the cured PEG-BAC samples have an increasing contribution of stretching features in the characteristic range for H-bonded amides with increasing w%BAC. This suggests that curing induces different spatial arrangements for the different H-bonding-capable monoacrylates, perhaps due to the greater number of H-bonding possibilities for PEG-HEA, resulting in larger H-bonding cluster size, as compared to PEG-BAC (see below). Finally, for the two network series IR spectral deconvolution indicates that while >95 mol% of HEA sidechains are in their H-bonded form throughout this series, the fraction of H-bonded BAC side chains rises progressively with increasing  $w\%_{BAC}$ , and even at the highest fraction some BAC retain non H-bonded characteristics (Fig. 2c). These two rather different H-bonding clustering behaviors inferred following curing they may result in different morphological, topological and mechanical properties. These effects that have not been previously reported for H-bond capable acrylate networks. Control of this clustering behavior over the emergent properties through the H-bond capable mono-acrylate content and its impact are described in the later sections.

Conventional <sup>1</sup>H NMR spectroscopy is also routinely used for evaluating the presence of H-bonding,<sup>16</sup> in particular the temperature dependence of characteristic resonance shifts can provide information on H-bonding induced changes in local electron density. As shown in Fig. S5 and S6 (ESI<sup>†</sup>), changes in linewidth are apparent, but even more clearly a significant change in the chemical shift can be observed in both series for resonances assigned to the H-bonding groups (3.4 ppm for  $\delta$ (OH) in HEA, 6.1 ppm for  $\delta$ (NH) in BAC), confirming the presence of H-bonding in the fully cured samples.

#### 3.3. Mechanical analysis of the networks

Dynamic mechanical analysis (DMA) provides viscoelastic parameters (*e.g.* storage and loss moduli) which can give insights into the macroscopic properties of polymer networks. The presence of a high temperature plateau and a single clean inflection for all the samples (Fig. 3), demonstrates that there is a single glass transition temperature,  $T_g$ , for all networks, indicating good mesoscopic mixing (on a length scale of smaller than several tens of nanometers) between PEGDA and the monoacrylate in all cases. The  $T_g$  values of the networks were determined from the temperature at the maximum of tan  $\delta$ .<sup>4</sup> The storage moduli (*E'*) of the two series show similar decreases with increasing monoacrylate content. According to classical rubber elasticity theory, the mean molar mass of



**Fig. 3** Storage modulus, E', of cured; (a) PEG-HEA; (b) PEG-BAC networks measured by DMA with increasing w<sub>HEA</sub> or w<sub>BAC</sub>; (c)  $T_g$  of the cured all samples, determined from DMA analysis, as a function of the weight % monoacrylate ( $\omega$ %<sub>MA</sub>). The PEG-HA data is extracted from our previous study.<sup>5</sup> The data for 100% monoacrylate samples were obtained from analysis of samples polymerized from pure BAC, HEA and HA.

network chains between chemical crosslinks and trapped chain entanglements,  $M_{c+e}$ , can be calculated using rubber elasticity theory (see ESI† for details). This relation is applicable to affine networks of the same chain length and without network defects. The  $M_{c+e}$  values are likely suffer from a large error for heterogeneous networks,<sup>38</sup> so the extracted values are used for qualitative comparisons of network heterogeneity within the series in the discussion below.

The Fox-Loshaek relation (a modification of Flory-Fox approach) predicts a linear dependence of  $T_{g}$  on monoacrylate content, see ESI.<sup>† 39</sup> As shown in Fig. 3c, a clear linear increase in  $T_{g}$  with  $w\%_{MA}$  is indeed observed for both series. Since a single sharp peak can be observed in the  $tan(\delta)$  temperature dependencies of all the samples (Fig. S7, ESI<sup>+</sup>), it can again be taken that there is good mixing and no mesoscopic phase separation of mono- and diacrylate. The increase in  $T_{\alpha}$  with increasing PEGDA content is the opposite to our previous observation for similar networks with the non H-bond capable monoacrylate HA.<sup>5</sup> We suggest that inter-chain H-bonding interactions act as temporary cross-links that restrict segmental motion and decrease the free volume of the network.<sup>40,41</sup> The slope of the w%<sub>MA</sub> dependence is greater for the HEA-than for the BAC-series, which, besides the significant larger molecular weight of BAC, is also a result of the larger 'spacer' (more atoms between the functional group and polyacrylate backbone) in the latter that is expected to increase molecular mobility and the available free volume, as previously reported for similar systems.13,42

#### 3.4. Low field (LF) <sup>1</sup>H NMR analysis of the networks

Polymer chain dynamics in crosslinked networks can be probed by <sup>1</sup>H NMR spin-spin relaxation time analysis, which is very sensitive to even small differences in the crosslink density due to the strong influence of anisotropic segmental motions on relaxation at temperatures well above  $T_{g}$  (more than 60 °C above  $T_{g}$ ).<sup>5</sup> As shown in Fig. 4, increasing  $w \%_{MA}$  (HEA or BAC) in the formulation raises the overall transverse relaxation times of the networks which we ascribe to decreasing crosslink density, as several previous studies have found that crosslink density of the polymeric network is inversely proportional to the  $T_2$  value at high temperature.<sup>5,25,43</sup> Swelling causes an increase in average  $T_2$ , as it releases temporary physical chain entanglements and a decrease in the strength of inter-chain spin-spin interactions. The shorter  $T_{2,av}^{sw}$  values across both series (Table 2), as compared to the non H-bonding capable (PEG-HA) networks we previously reported,<sup>5</sup> demonstrates reduced network chain mobility due to the presence of H-bonding clusters. We have shown<sup>5,6</sup> that it is also possible to estimate apparent  $M_{c+e}$ values from the  $T_2$  values (details in ESI<sup>+</sup>), the relative changes in the  $M_{c+e}$  values extracted will be discussed below. We note that the temperature of 70 °C, necessary for the LF NMR analysis, precludes direct comparison with the FT-IR data from Section 3.2.

#### 3.5. Network analyses by DQ NMR

<sup>1</sup>H DQ-NMR analysis offers more detailed insights into the effect of network structure on chain dynamics. The combination of fast sample spinning and high magnetic field strength provides spectral resolution permitting the acquisition of site-specific, chemical-shift resolved DQ build-up curves for the different chemical groups present, which is more informative than the averaged information provided by low field NMR (LF NMR) relaxometry, or indeed by the  $T_g$  analysis. The <sup>1</sup>H spectra for both sample series are shown in Fig. S5 (ESI<sup>†</sup>)



**Fig. 4** Normalized  $T_2$  relaxation decays, recorded at 70 °C and 20 MHz, of; (a and c) PEG-HEA networks; (b and d) PEG-BAC networks measured using Hahn echo pulse sequence ( $\tau_{CP}$ : 10 ms) in dry state (a and b), and swollen state (c and d). Volume swelling ratio in  $C_2D_2Cl_4$ , is  $\approx$  1.55). Vertical lines indicate the  $T_{2,av}$  values. The points represent the measured data and the lines are the corresponding fits. Details on normalization and fitting are provided in ESI† Section 5.

and representative normalized DQ build-up curves and fits are shown in Fig. S8 (ESI $\ddagger$ ).

By evaluating the DQ build-up curves for the <sup>1</sup>H resonance at 3.56 ppm, assigned to the CH<sub>2</sub> groups in PEG chains (except for the CH<sub>2</sub> next to the carbonyl group of the acrylate backbone), the conformational freedom of the PEGDA can be evaluated. As shown Fig. 5, the average residual dipolar coupling constant of the distributions ( $D_{avg}$ ), indicated by the vertical dotted lines (values in Table 2), also gradually decreases with  $w%_{MA}$  for both series due to greater motional freedom arising from decreasing crosslink density. This coincides with the increase in  $T_{2,av}^{sw}$  noted above which is also ascribed to increased motional freedom. These findings are similar to the observations for the PEG chain in the non H-bonding capable networks (PEG-HA),<sup>5</sup> indicating that H-bonding has very limited impact on the PEG chain motion, consistent with observations in similar systems.<sup>16</sup>

As shown in Fig. 6a and c, the  $D_{avg}$  values of all the chemical groups in the both series are found to decrease linearly with increasing  $w\%_{MA}$ , as observed for the non H-bonding networks.<sup>5</sup> As expected, resonances related to the polyacrylate backbone in both series have greater  $D_{avg}$  values than the PEG chains, with the highest values observed for PEG-HEA, indicating a stiffer polyacrylate backbone in this network series. The chain segments in the monoacrylate side chain (namely HEA

or BAC) behaved differently depending on their position in relation to the H-bonding moiety. Since the polymer chain (PA and PEG) segments comprise from distinct local spin systems, direct comparison of the absolute  $D_{avg}$  values is complicated. Therefore, the individual coupling constants have been normalized here with respect to those for  $w\%_{MA}$  (90%). The fact that the scaled values for the two resonances of the acrylic backbone are almost identical for both series demonstrates that this approach is at least internally consistent.

For the PEG-HEA networks, as shown in Fig. 6b, apart from the PEG units discussed before, all the resonances can be divided into two groups: group I, those belonging to the polyacrylate backbone, and group II, those located on PEG and mono-acrylate chains near the polyacrylate backbone or H-bonding functional group (-OH or -NH-). It is apparent from Fig. 6b that the change in normalized  $D_{avg}$  with  $w\%_{MA}$  is larger for group I than for group II. Whereas stronger segmental motion restriction (higher  $D_{avg}$ ) in group II can be observed as compared to the equivalent sidechains in the non-H-bonding PEG-HA counterpart.<sup>5</sup> This is also apparent from the absolute  $D_{avg}$  values for H(f), which are two times larger than its counterpart (-(CH<sub>2</sub>)<sub> $\beta$ </sub>-) in the PEG-HA system.<sup>5</sup> For the PEG-BAC networks, the normalized  $D_{avg}$  values for resonances in group I show a smaller variation in PA backbone stiffness with increasing  $w\%_{MA}$  than for PEG-HEA or PEG-HA, which



**Fig. 5** Residual dipolar coupling constant,  $D_{\text{res}}$ , distributions of PEG chains ( $\delta$  3.56 ppm) for; (a) PEG-HEA; (b) PEG-BAC networks at different  $w_{\text{MA}}^{\text{MA}}$ . Data were recorded at 70 °C. Vertical doted lines indicate the (weighted) average coupling constant,  $D_{\text{avg}}$ , of each distribution.

may originate from the  $w\%_{BAC}$  dependence of the H-bonded fraction of BAC side chains (IR analysis, Fig. 2c). Moreover, a third group (*group III*) of resonances arising from methylene groups that are two bonds away from the H-bonding group and not adjacent to the polyacrylate backbone, can be further separated based on their less restrict segmental motions. The absolute and normalized  $D_{avg}$  values for *group III* can be found to be similar to those from the side chains of PEG-HA,<sup>5</sup> and have the weakest  $w\%_{MA}$  dependence.

#### 4. Discussion

#### 4.1. Effect of H-bonding on network structure

As proposed by Jansen *et al.* in their photo-induced polymerization kinetic studies, a so-called 'pre-organization' of monoacrylates containing H-bond capable groups in their side chains, typically occurs in the pre-curing formulations which can in turn increase the homo-polymerization rates.<sup>44,45</sup> In this study, IR analysis of all the pre-curing formulations (see Fig S10, ESI†) shows that for both series the monomers are in H-bonded forms, suggesting that pre-organization indeed occurred in both systems. However, further IR analysis on the cured samples shows that a significant fraction of BAC in PEG-BAC networks is not H-bonded, which is not the case for PEG-HEA. For polymer networks with preferable homopolymerization, a diagonal peak with stronger intensity than its corresponding cross peak would be expected in the DQ-SQ NMR analyses. Further DQ-SQ NMR analyses of PEG-BAC shows no preferable homo-polymerization of BAC, as the diagonal peak of BAC in 2D NMR spectra has similar intensity as the cross peak between BAC and PEGDA, again this is not the case for PEG-HEA (Fig. 7a and b).

This observation is unexpected, as Jansen pointed out in his work,<sup>44</sup> the homo-polymerization rates of H-bonding capable monomers are significantly (4- to 8-fold) enhanced, which should result in the formation of large clusters of H-bonded units. Hence, we propose a competing-reaction-rate model to explain this discrepancy for the two series. As illustrated in Fig. 7c using t-OH type mono-acrylate, upon mixing the diacrylate and H-bond capable monoacrylates, these monoacrylates spontaneously rearrange to form an ordered 'pre-organized' system. On initiation of photopolymerization, depending on the ratio of homo- $(k_{hp})$  and co-polymerization  $(k_{cp})$  propagation rates, the extent of retention of a H-bonded fraction can be significantly different. In scenario (i),  $k_{\rm hp} > k_{\rm cp}$  and we suggest this faster monoacrylate homo-polymerization can generate a network with large H-bonded clusters, as in the case of PEG-HEA. On the other hand, in scenario (ii) higher  $k_{cp}$  can result in a more homogeneous network with H-bonded clustering reduced, as in the case of PEG-BAC.

It is anticipated that the presence of H-bonding capable mono-acrylates should increase  $k_{\rm hp}$  as well as  $k_{\rm cp}$ , as suggested by previous work.<sup>44</sup> Hence, the above observation suggests that the induced increase in  $k_{cp}$  is comparable to that in  $k_{hp}$ , which in turn indicates that there is no direct connection between the formation of larger H-bonded mono-acrylate clusters and the pre-organization phenomenon. Cluster formation can be considered as similar to other acrylate systems, such as the non-Hbonding capable PEG-HA networks,<sup>5</sup> namely their formation is a kinetically driven process. To evaluate the differences in kinetic preference for homo-polymerization in HEA and BAC, a simple Alfrey-Price Q-e calculation of the relative reaction rate in these two networks was performed (details of these calculations are given in Section S7, ESI<sup>+</sup>). This analysis suggests a faster homo-polymerization rate for HEA than for its copolymerization with PEGDA, while the copolymerization reaction for PEG-BAC system is faster, which supports our interpretation.

In summary, as a result of faster homo-polymerization of HEA in PEG-HEA networks due to higher intrinsic homopolymerization rate, larger H-bonded HEA clusters are formed. On the other hand, despite pre-organization also being present for PEG-BAC, faster copolymerization between BAC and PEG largely negates any pre-organization and the reorganization of dynamic H-bonded network structure produces a more homogenous network with minimal retention of structures comprised of H-bonded BAC units. Hence, the difference in H-bonding cluster size between the two series is



**Fig. 6** Residual dipolar coupling,  $D_{res}$ , values of all <sup>1</sup>H chemical shift specified network segments for; (a) PEG-HEA, and; (c) PEG-BAC networks, as a function of  $w_{MA}^{*}$ . In (b) and (d) the  $D_{avg}$  values from (a) and (c) are normalized with respect to the appropriate  $D_{avg}$  values determined for  $w_{MA}^{*}$  90%.

mainly driven by the different copolymerization kinetics of the mono-acrylates.

#### 4.2. Effect of H-bonding on apparent $M_{c+e}$ of network chains

The crosslink/entanglement density, which is the inverse of  $M_{c+e}$ , is a key parameter linking microscopic network structures and macroscopic properties. As noted above, the apparent mean molecular mass,  $M_{cre}^{app}$ , can be calculated from the storge modulus E of the high temperature plateau in DMA, as well as from the  $T_{2,av}^{sw}$  values from LF NMR relaxometry, as described previously.<sup>44</sup> The data in Fig. 8 shows that the values from these two approaches agree well and are in the expected range, suggesting that they are at least sensitive to the same network density. Note that DQ NMR can in principle also be used to estimate  $M_{c+e}^{\text{app}}$ , however we have not included it here as that value would only reflect the individual moieties. Furthermore, it is difficult to extrapolate them to obtain the average value for the overall network due to the lack of  $d_{ref}$  values for different moieties other than PEG (see description below). As expected, the  $M_{c+e}^{app}$  value of the all the networks increases with increasing w%<sub>MA</sub>, an expected apparent decrease in crosslink density. Furthermore, as compared to the PEG-HA series, the values for PEG-BAC and PEG-HEA networks, are reduced across the  $w\%_{MA}$  range, due to the persistent effect of H-bonding induced reduction in the apparent network density. The data also suggests greater apparent crosslink/entanglement density in

PEG-HEA, as compared to PEG-BAC, which is again consistent with stronger H-bonded effects for the former. However, whether this effect is equally distributed across all the network segments cannot be assessed by these volume average techniques. Hence, we will use the chemical shift specified DQ NMR analyses to evaluate H-bonding effects on polymer chain mobility in more detail below.

#### 4.3. Effect of H-bonding on network polymer chain mobility

In photocured acrylate networks, the sidechain of these acrylates are known to form in a zipper-like topology in the final polymer, and are often referred to as zipper-like junctions.<sup>46,47</sup> This nano-scale network topology has far-reaching impacts on the macroscopic and in particular the mechanical properties.<sup>6</sup> We have previously shown that for PEG-HA networks, the weighted-averaged  $D_{res}$  value ( $D_{avg}$ ) is sensitive to differences in segmental polymer chain dynamics induced by changes in network topology.<sup>5</sup> As shown in Fig. 9a, the D<sub>avg</sub> of individual <sup>1</sup>H resonances in PEG-BAC network does not monotonically decrease with their position along the BAC side chain from the polyacrylate (PA) backbone, which is unlike the situation for non H-bonding PEG-HA network.<sup>5</sup> Here the units near the H-bonding moiety have a larger  $D_{avg}$  value as a result of reduced chain segment mobility. By plotting  $D_{avg}$  of the mono-acrylate side chain resonance groups assigned previously, see Fig. 9b, it can be seen that for PEG-BAC networks: (i) group II (<sup>1</sup>H in the



**Fig. 7** 2D DQ-SQ correlation spectrum of; (a) PEG-HEA  $w_{MEA}^{S}$  (50); (b) PEG-BAC  $w_{BAC}^{S}$  (50). The red dotted lines connect cross-peak pairs between BAC side chain and PEG resonances The red box highlights the cross-peak between PEG and HEA/BAC. (c) Schematic representation of copolymerization of diacrylate and monoacrylate containing H-bond capable moieties, the real situation is more nuanced, see Section 4.3. Pre-organization occurs spontaneously on mixing. After initiation of the free radical polymerization, depending on the relative monoacrylate co-polymerization ( $k_{CP}$ ) and homopolymerization ( $k_{hp}$ ) propagation rates, the final network may have H-bonding clusters with variable size, *i.e.* H-bonding clusters are dynamic domains that are fixed by crosslinking.

chemical group adjacent to the PA backbone or N-H/O-H group) show very similar changes with increasing  $w\%_{MA}$  as group II in PEG-HEA series; (ii) for group III (<sup>1</sup>H in the chemical group two bonds away from N-H/O-H and not adjacent to the PA backbone) in PEG-BAC networks, the w%<sub>MA</sub> dependence is very similarly to that for group III in non-H-bonded PEG-HA series. Note that here the extent of this increase in  $D_{avg}$  for <sup>1</sup>H in chemical group adjacent to N-H/O-H groups is also dependent on the distance between the backbone and H-bonding group, as suggested by a similar test for a PEG-HBA (4-hydroxybutyl acrylate) network, see Fig. S11 (ESI†). The data suggests that, as shown schematically in Fig. 9c, the H-bonding capable groups impose a strongly localized restraint (within 1-2 bonds) on side chain mobility, while subsequent units along these chains follow the free chain model (progressively decreasing Davg along the sidechain) as found for non H-bonding networks. In a previous mechanical study<sup>13</sup> it was suggested that side chains with multiple H-bonding groups significantly reduce chain mobility even in very loosely crosslinked networks. In this study we use monomers with single H-bonding sites. It is likely that H-bonding causes a strongly localized mobility restraint on nearby chain segments.

Furthermore, as suggested by Jansen's study on pre-organization effects,<sup>44</sup> though polyacrylates produced by a free radical polymerization normally yield predominantly syndiotactic materials (*i.e.* zipper-like network junction), the pre-organization of H-bonding capable mono-acrylates results in an increasingly favorable isotactic structure (*i.e.* brush- or comb-like network junctions) in the clusters (see Fig. 9c), which will in turn decreases the PA backbone chain mobility.<sup>48</sup> However, as shown in Fig. 6b, the normalized  $D_{avg}$  values of PA backbone in PEG-HEA network have very similar values and  $w%_{HEA}$  dependence as those in the PEG-HA series,<sup>5</sup> suggesting the overall network junction topology (zipper-like network) did not change significantly in presence of H-bonding induced pre-organization. Note that as mentioned above, the lower normalized  $D_{avg}$  values of



**Fig. 8** Mean molar mass between crosslinks and trapped chain entanglements,  $M_{C+e}^{app}$ , obtained from DMA (dry sample) and LF NMR analysis (swollen sample), as a function of monoacrylate weight fraction,  $w_{MA}$ . The PEG-HA data is extracted from our previous study.<sup>5</sup>

the PA backbone for PEG-BAC arose from an increased overall reduction of network chain mobility due to increased ratio of H-bonded BAC monoacrylates, hence we do not comment here on the change in its topology as compared to the other two network types.

#### 4.4. Effect of hydrogen bonding on network junction functionality

The network junction functionality, f, is defined as the number of elastically effective strands connected to a network junction, it is mainly determined by the chemistry, however, as free radical polymerization is uncontrollable, difference can also arise due to the change in the polymerization kinetics. For the networks in this study f is effectively the number of directly adjacent acrylate backbone monomer units bearing PEG chains,  $N_{\text{PEG}}$ . For a pure PEGDA network ( $w\%_{\text{MA}} = 0$ ), the functionality is theoretically equivalent to the polyacrylate backbone length which can be directly obtained from GPC analyses of the hydrolysates; n(PAA) and is c. 150, a similar value to that obtained for the PEG-HA networks in our previous study.5 The addition of mono-functional acrylates into the formulation generates topological defects in the zipper-like polyacrylate network junctions, which significantly reduces the network junction size (indicated in grey in Fig. 10c) and hence its functionality. Hence with increasing  $w %_{MA}$  it is expected that  $f_{eff}$  (the effective volume average functionality) will gradually decrease and finally reach a value no bigger than 3, since the network junction functionality is  $f = N_{\text{PEG}} + 2$  (or + 1, if the position of the crosslinking PEG is at the acrylic backbone terminal), and  $N_{\text{PEG}}$  = 1 for extremely diluted PEGDA networks. The functionality of acrylic network junctions in PEGDA-MA



**Fig. 9** (a) Average residual dipolar coupling constant  $D_{avg}$  values of chemical groups along the BAC side chains in PEG-BAC networks, for orientation within the network see panel (c); (b) The  $D_{avg}$  values for identified resonances of specified network segments in the monoacrylate side chains for PEG-HEA, PEG-BAC and PEG-HA as a function of  $w%_{MA}$ ; (c) Schematic representation of network dynamics for the PEG-BAC case, showing the polyacrylate backbone and sidechains and the PEG crosslinker with a H-bonding cluster highlighted in yellow.



**Fig. 10** (a) Effective functionality of acrylic crosslinks in PEG-HEA and PEG-BAC networks as well as PEG-HA (extracted from our previous data<sup>5</sup>) derived from DQ-NMR analysis; (b) comparison of phantom and affine model pre-factors of all these networks as a function of crosslink functionality; (c) illustration of network junction with H-bonding clusters; (d) illustration of network junction without H-bonding clusters.

networks (of any type) can be estimated by analyzing DQ-NMR results for PEG chains using a similar model (PEGDA-PETM) network with known functionality as a ref. 5. Based on these considerations, the  $D_{\text{avg}}$  and  $M_{c+e}$  values of the PEG chains can be related by:<sup>49,50</sup>

$$M_{c+e}^{\rm NMR} = \frac{f-2}{f} \frac{d_{\rm ref}}{D_{\rm avg}/2\pi}$$
(1)

Where  $d_{\rm ref}$  is the polymer specific reference coupling, a preaveraged dipolar coupling constant characterizing the spin arrangement and local motions within a statistical (Kuhn) segment.<sup>45,46</sup> As mentioned in our previous study,<sup>5</sup> for the network containing point-like junctions f can be calculated from its double bond conversion rate to be c. 3.64, assuming no significant amount of network defects, *e.g.* loops, are formed. Under the assumption that  $d_{\rm ref}$  does not change for the networks studied here, which is justified in part by the fact that all the networks comprise PEG chains of the same length, an apparent functionality can then be estimated for the two H-bonding capable networks. A deviation of  $f_{\rm eff}$  was observed from the expected f value for pure PEGDA of c. 150, which may be attributed to the residual mobility of the elongated polyacrylate chains. Hence f values obtained from eqn (1) using  $D_{\text{avg}}$ ,  $d_{\text{ref}}$  and  $M_{c+e}$  values from the experiments must be treated with some caution. The results are shown in Fig. 10a along with the values for PEG-HA. As expected the apparent functionality decreases strongly with increasing  $w \%_{MA}$  in all cases. It is found that PEG-HEA has consistently higher apparent  $f_{\text{eff}}$  than PEG-BAC and PEG-HA, which we suggest is due to the formation of larger H-bonding clusters, due to the kinetics considerations discussed above. This effect also pushes these networks towards a more affine-like (restricted chain motion) situation at all  $w \%_{\text{HEA}}$ , Fig. 10b. As illustrated in Fig. 10c and d, this interpretation makes intuitive sense; as the formation of such clusters should indeed drive the formation of larger network junctions and so higher functionality.

### 5. Conclusions

In this study, two series of polyacrylate networks with different H-bonding capable monoacrylate copolymers were synthesized *via* UV-initiated photo-polymerization. It was found that different patterns of H-bonding clusters associated with the side chains were produced in the two network series. IR analysis and reactivity calculations demonstrate that preferable formation of clusters during UV-curing is mainly affected by differences in homo- and co-polymerization kinetics of the monoacrylates and PEGDA, hence we propose a competing-reaction-rate model to explain preferred cluster formation for PEG-HEA. Hence the competing roles of H-bonding preorganization and copolymerization kinetics are supported by ex situ FTIR analyses and indirectly by the reactivity calculations. Additional in situ FTIR- and NMRbased kinetic analyses, which can help us to track the evolution of individual compound over time, are currently undergoing that will allow us to develop this interpretation. Nevertheless, in this view thermodynamically favorable pre-organization is present for both PEG-HEA and PEG-BAC, significantly increasing the homopolymerization rates, however the differences in H-bonding cluster size arise from differences in the co-polymerization rates. This is the first direct observation of H-bonding capable monoacrylate induced enhancement in co-polymerization between monoacrylate and crosslinker in UV-copolymerization.

Additionally, the presence of H-bonding reduces the overall chain mobility in both networks, while different chain segments are strongly influenced (as indicated by the change in  $D_{\rm res}$  along the side chain) by their position relative to the H-bonding group in the monoacrylate sidechain. Furthermore, though the preferable formation of H-bonding clusters does not change the overall polymer network junction (zipper-like) topology significantly, larger clusters indeed increase the size of the network junctions. This observation clearly demonstrates the important role of H-bonding type in the occurrence of monoacrylate pre-organization during UV initiated photo-polymerization processes. The microscopic picture that emerges is, H-bonding imposed dynamic restrictions during sample preparation and polymerization that coupled with the known monomer reactivities alter network organization, which also directly impact the emergent bulk properties.

## Conflicts of interest

There are no conflicts to declare.

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