

Highlighting research from the groups of Takeshi Ueki and Ryota Tamate.

Controlling mechanical properties of ultrahigh molecular weight ion gels by chemical structure of ionic liquids and monomers

A new class of ion gels formed by physical entanglement of ultrahigh molecular weight polymers in ionic liquids is synthesised *via* a facile one-step process. Ultrahigh molecular weight gels exhibit superior mechanical characteristics such as stretchability, recyclability, and self-healing ability. The effects of the chemical structures of the ionic liquids and monomers on the mechanical properties are investigated in detail.

### As featured in:



See Ryota Tamate,  
Takeshi Ueki *et al.*,  
*Soft Matter*, 2022, **18**, 8582.



Cite this: *Soft Matter*, 2022, 18, 8582

## Controlling mechanical properties of ultrahigh molecular weight ion gels by chemical structure of ionic liquids and monomers†

Yuji Kamiyama,<sup>ab</sup> Ryota Tamate, <sup>cd</sup> Kenta Fujii <sup>e</sup> and Takeshi Ueki <sup>ab</sup>

A new class of ion gels, termed ultrahigh molecular weight (UHMW) gels, formed by physical entanglement of ultrahigh molecular weight polymers in ionic liquids, are synthesised using facile one step radical polymerisation with significantly low initiator conditions, and exhibit superior mechanical characteristics such as stretchability, recyclability, and room temperature self-healing ability. In this study, UHMW gels are synthesised using various combinations of monomer and IL structures, and the effect of their chemical structures on the physicochemical properties of UHMW gels are thoroughly investigated. UHMW polymers are prepared *in situ* for all combinations of ILs and monomers used in this study, indicating the wide applicability of this fabrication strategy. The structure–property relationships between chemical structures and mechanical properties of UHMW gels are investigated in detail. Furthermore, the differences in self-healing efficiency of UHMW gels depending on the chemical structure is discussed in terms of individual polymer conformation and polymer–polymer interaction based on molecular dynamics simulations.

Received 27th June 2022,  
Accepted 7th October 2022

DOI: 10.1039/d2sm00853j

[rsc.li/soft-matter-journal](http://rsc.li/soft-matter-journal)

## Introduction

With the advent of the Internet of things (IoT) era, wearable and flexible electronics have received increasing attention.<sup>1–3</sup> In this regard, conductive and stretchable polymeric materials have been intensively investigated in recent years.<sup>4–10</sup> Ion-conductive polymeric materials are important components in flexible and wearable devices such as secondary batteries, sensors, electric double layer transistors, supercapacitors, and soft actuators.<sup>11–15</sup> Dry ionic conductors such as poly(ethylene oxide) composed with inorganic lithium salts have good mechanical toughness and are free from solvent evaporation.<sup>16</sup> However, ionic conductivity is generally low as the ion transport is dominated by the segmental motion of polymer chains. Although hydrogels and organogels that dissolve inorganic salts in swollen polymer

networks show relatively high ionic conductivity, they are susceptible to solvent evaporation.<sup>17</sup> Whereas, ion gels, defined as a gel-type ionic conductor containing ionic liquids (ILs) as a solvent, are a new class of ionically conductive soft materials that satisfy high ionic conductivity as well as non-volatility.<sup>18–21</sup>

In general, the poor mechanical properties of polymeric gels due to high liquid content limit their applications in flexible and wearable devices, in which tolerance for repetitive mechanical loading is required.<sup>22</sup> So far, various strategies for toughening polymer gels have been proposed.<sup>23–28</sup> Recently, introduction of self-healing functionalities into polymer gels has been investigated to achieve significant improvement in material lifetime.<sup>29–34</sup> Several studies have reported the fabrication of self-healing and stretchable ion gels using intricately designed reversible bonds.<sup>35–41</sup> However, such materials often require time-consuming preparation/purification procedures and complicated synthesis of specially designed functional monomers and polymer networks.

In our previous study, we have reported a facile one-step strategy for fabrication of stretchable and self-healable ion gels based on physical entanglement of ultrahigh molecular weight (UHMW) polymers.<sup>42</sup> The radical polymerisation of methacrylate monomers in ILs, for example a typical IL 1-ethyl-3-methylimidazolium bis(trifluoromethanesulfonyl)imide ( $[C_2mIm][TFSI]$ ), yields a monomer conversion of nearly 100% even at significantly low initiator concentrations, which consequently produced ion gels solely by the physical entanglements of UHMW polymers over  $10^6$  Da, which were named as UHMW gels. However, the effect of

<sup>a</sup> Research Center for Functional Materials, National Institute for Materials Science, 1-1 Namiki, Tsukuba, Ibaraki 305-0044, Japan. E-mail: UEKI.Takeshi@nims.go.jp

<sup>b</sup> Graduate School of Life Science, Hokkaido University, Kita 10, Nishi 8, Kita-ku, Sapporo, Hokkaido, 060-0810, Japan

<sup>c</sup> Center for Green Research on Energy and Environmental Materials, National Institute for Materials Science, 1-1 Namiki, Tsukuba, Ibaraki 305-0044, Japan. E-mail: TAMATE.Ryota@nims.go.jp

<sup>d</sup> PRESTO, JST, 7 Gobancho, Chiyoda-ku, Tokyo, 102-0076, Japan

<sup>e</sup> Graduate School of Sciences and Technology for Innovation, Yamaguchi University, Ube, Yamaguchi 755-8611, Japan

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2sm00853j>



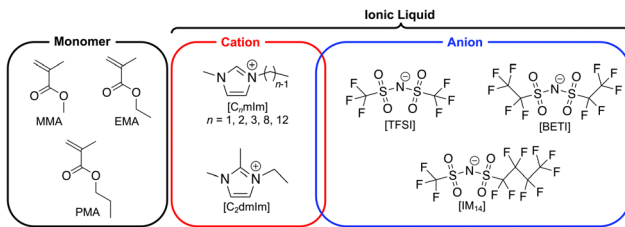


Fig. 1 The chemical structures of monomer, IL cation, and anions used in this study.

the chemical structures of ILs and methacrylate monomers on the polymerisation and mechanical properties of UHMW gels have not been understood. In particular, as the possibility to control the mechanical properties of gels through the solvent structure is a unique feature of ion gels, investigation of the effect of IL cation and anion structures has a great importance. Herein, UHMW gels were synthesised from various IL and monomer structures to investigate the effect of their chemical structures on the *in situ* formed UHMW polymers. The chemical structure of monomers and ILs are shown in Fig. 1. The IL solvents for ion gels were selected based on the compatibility of the methacrylate polymers. Next, the rheological, mechanical properties, and self-healing ability of UHMW gels composed of various IL and polymer structures were investigated in detail to elucidate the structure–property relationship of UHMW gels. Furthermore, molecular dynamics (MD) simulations were performed to decipher the influence of IL and polymer structures on self-healing ability of UHMW gels at microscopic level. It was implied that the individual polymer conformation and interpolymer interactions were adjusted depending on IL/polymer chemical structures, which resulted in the difference in the self-healing ability of UHMW gels.

## Experimental

### Materials

1-Alkyl-3-methylimidazolium bis(trifluoromethanesulfonyl)imide ( $[C_nIm][TFSI]$ ,  $n = 1, 2, 8, 12$ ) and 1-ethyl-2,3-dimethylimidazolium bis(trifluoromethanesulfonyl)imide ( $[C_2dmIm][TFSI]$ ) were purchased from Kanto Chemical (Japan).  $[C_3mIm][TFSI]$  was purchased from TCI (Japan).  $[C_2mIm][BETI]$  ( $BETI$ : bis(pentafluoroethanesulfonyl)imide) and  $[C_2mIm][IM_{14}]$  ( $IM_{14}$ : (trifluoromethanesulfonyl)(nonafluorobutanesulfonyl)imide) were synthesised according to a literature reference with slight modifications.<sup>43</sup> Lithium salts,  $Li[BETI]$  and  $Li[IM_{14}]$ , which were used for synthesis of these ILs, were purchased from Mitsubishi Materials Electronic Chemical Co., Ltd (Japan). All the ILs were dried under vacuum at 120 °C for 24 h prior to use. Methyl methacrylate (MMA) and ethyl methacrylate (EMA) were purchased from Kanto Chemical (Japan), and propyl methacrylate (PMA) was purchased from TCI (Japan). All monomers were purified by passing them through the basic alumina column to remove polymerisation inhibitor. 2,2'-Azobis(isobutyronitrile) (AIBN) were purchased from FUJIFILM Wako Pure Chemical Corporation (Japan) and used as received.

### Synthesis and characterisation of ultrahigh molecular weight gels

For the synthesis of all UHMW gels, the monomer concentration and the AIBN content were set at 40 vol% and 0.02 mol% *vs.* monomer content, respectively. A representative synthetic procedure for the PMMA/ $[C_2mIm][TFSI]$  UHMW gel is described as follows. MMA (1.6 mL, 15.0 mmol),  $[C_2mIm][TFSI]$  (2.4 mL, 9.32 mmol), and AIBN (0.49 mg,  $3.0 \times 10^{-3}$  mmol) were charged in a glass cup and sealed with a rubber septum, through which argon was bubbled for 15 min at room temperature. Subsequently, polymerisation reaction was carried out at 80 °C for 24 h, resulting in the formation of the transparent ion gel. Monomer conversion was determined using  $^1H$  nuclear magnetic resonance (NMR) measurements for the as prepared UHMW gels dissolved in deuterated chloroform ( $CDCl_3$ ). For the determination of molecular weight of *in situ* formed UHMW polymers, UHMW gels were purified by reprecipitation with acetone as good solvent and methanol as poor solvent twice and then vacuum dried at 60 °C, resulting in white powder of UHMW polymers. The molecular weight and polydispersity index (PDI) of the polymers were determined using gel permeation chromatography (GPC), in which 10 mM lithium bromide (LiBr) in *N,N*-dimethylformamide (DMF) solution was used as the eluent. The GPC columns (Showa Denko, Japan) were calibrated using PMMA standards. Prior to rheological and mechanical measurements, UHMW gels were dried *in vacuo* at 80 °C overnight to remove residual monomers.

### Rheological measurements

Oscillatory shear measurements were performed with an Anton Paar MCR 102 rheometer (Anton Paar, Austria) using the parallel plate geometry with a 12 mm diameter plate. Gap spacing was kept at 0.5 mm for all the measurements. Frequency sweep measurements were performed over a frequency range of 0.1–100 rad s<sup>-1</sup> with a strain amplitude of 1% at constant temperatures from 10 to 120 °C at intervals of 10 °C. The time-temperature superposition (tTS) master curves for UHMW gels were constructed from the frequency sweep data at different temperatures.

### Tensile tests

Tensile tests of UHMW gels were carried out using a Shimadzu AGS-X tester (Shimadzu, Japan). The gels cut into a dumbbell-shape (JIS K 6251-7 size, width: 2 mm, gauge length: 12 mm) were stretched at a speed of 10 cm min<sup>-1</sup> with a 100 N load cell. For the self-healing test, the dumbbell-shaped gels were cut at the centre, subsequently the two cut surfaces were contacted and allowed to heal at room temperature for 24 h, after which tensile tests were performed for the healed samples.

### Thermal analyses

Differential scanning calorimetry (DSC) measurements were performed using a DSC250 instrument (TA Instruments, USA) under a nitrogen atmosphere. The samples were placed on an open aluminum pan and heated from 40–200 °C, followed by



cooling to  $-120\text{ }^{\circ}\text{C}$ , and then reheated to  $200\text{ }^{\circ}\text{C}$  at a cooling and heating rate of  $10\text{ }^{\circ}\text{C min}^{-1}$ . The glass transition temperature ( $T_g$ ) (the temperature at the peak top of derivative heat flow) were determined from the DSC thermograms during the 2nd heating step.

### Molecular dynamics simulations

All-atom molecular dynamics (MD) simulations were carried out using the GROMACS 2020.1 program under the isobaric-isothermal (NTP) ensemble (298 K and 1 atm) in a cubic cell. Procedures similar to that reported in our previous study were followed in this case.<sup>42</sup> Number of ILs and polymers in the simulation boxes are listed in Table S1 (ESI<sup>†</sup>). A 20 mer (approximately 2 kDa) was used as the polymer due to computing performance issues. The total simulation time was set to be 25 ns. From the last 500 ps, the data was collected with 0.1 ps intervals, and analysed to determine the radial distribution function  $G(r)$ . The density values calculated from the present MD simulations are consistent with the density values estimated from the respective densities of the IL and the polymer, which are also listed in Table S1 (ESI<sup>†</sup>). CLaP and OPLS-AA force fields, including intermolecular Lennard-Jones (LJ), coulombic interactions, and intramolecular interactions (bond stretching, angle bending, and torsion of dihedral angles) were used for the  $\text{C}_2\text{mIm}$  and  $\text{C}_{12}\text{mIm}$  cations, TFSI anion, and PMMA and PEMA polymers.<sup>44–46</sup> The radius of gyration ( $R_g$ ) for the polymer was calculated from the intramolecular component  $G_{\text{intra}}(r)$ , according to the following equation, where  $r_{\text{max}}$  represents the upper limit of the integration, and here we assumed a sufficiently large  $r$  (35 Å) that there is no intramolecular correlation of the polymers:

$$R_g^2 = \frac{\int_0^{r_{\text{max}}} r^2 4\pi r^2 G(r) dr}{\int_0^{r_{\text{max}}} 4\pi r^2 G(r) dr}$$

## Results and discussion

### Synthesis of UHMW gels from various IL and monomer structures

To prepare UHMW gels with different IL structures, methyl methacrylate (MMA) monomer was polymerised in a wide variety of ILs consisting of combinations of different imidazolium-based cations ( $\text{C}_n\text{mIm}$ :  $n = 2\text{--}12$  and  $\text{C}_2\text{dmIm}$ ) and sulfonylimide anions with different lengths of perfluoroalkyl groups (TFSI, BETI, and  $\text{IM}_{14}$ ). In addition, ethyl methacrylate (EMA) and propyl methacrylate (PMA) were polymerised in  $[\text{C}_2\text{mIm}]\text{[TFSI]}$  and  $[\text{C}_2\text{mIm}]\text{[BETI]}$ , respectively to investigate the effect of monomer structures. Note that when PMA was polymerised in  $[\text{C}_2\text{mIm}]\text{[TFSI]}$ , it was too soft to be moulded for mechanical tests probably due to its very low glass transition temperature ( $T_g$ ). Therefore, polymerisation of PMA monomer was performed in  $[\text{C}_2\text{mIm}]\text{[BETI]}$ , which has a higher  $T_g$  than that of  $[\text{C}_2\text{mIm}]\text{[TFSI]}$ . The chemical structures of ILs and monomers used in this study are shown in Fig. 1. For the synthesis of UHMW gels, the monomer concentration was fixed at 40 vol% and the initiator (2,2'-azobis(isobutyronitrile), AIBN) content was set at as low as 0.02 mol% vs. monomer content. After argon

bubbling of precursor solutions containing ILs, monomers and initiator, polymerisation was allowed to proceed at  $80\text{ }^{\circ}\text{C}$  for 24 h. For all combinations of monomers and ILs, transparent ion gels without any macroscopic polymer-IL phase separation were formed *in situ* as polymerisation proceeded, indicating that the robust physical entanglement of UHMW polymers were formed in ILs. The results of polymerisation (molecular weight, polydispersity index (PDI), and monomer conversion) are summarised in Table 1. GPC traces of synthesised polymers extracted from UHMW gels and  $^1\text{H}$  NMR spectra of as prepared UHMW gels after polymerisation are shown in Fig. S1 and S2 (ESI<sup>†</sup>). For all monomer-IL combinations used in this study, UHMW polymers with molecular weights near or above  $10^6$  Da were obtained with high monomer conversion of over 95%. These results indicate that the fabrication strategy of UHMW gels can be applicable for wide variety of methacrylate polymer and IL structures, which would render tunability of physicochemical properties of UHMW gels. Previous experimental and computational investigations for the radical polymerisation in ILs suggested that the rate constant of propagation ( $k_p$ ) was enhanced with decrease in activation energy due to the IL-radical interaction.<sup>47,48</sup> In addition, our previous study that compared viscosity-tuned toluene and ILs as polymerization media revealed that the effect of solvent viscosity that potentially slow down the termination reaction of growing polymer on the molecular weight was not significant, thus enhancement of  $k_p$  would be a dominant factor for the formation of UHMW polymers in ILs.<sup>42</sup> In the present study, consistent with the previous results, the molecular weight of the synthesised PMMA in various ILs tended to increase with increasing ionic concentration in the precursor solutions (Fig. S3, ESI<sup>†</sup>), suggesting that the more the ionic moieties around the propagation radicals, the higher the  $k_p$  due to the IL-radical interaction. Furthermore, the PDIs of obtained UHMW polymers were slightly below 2.0, which were significantly lower than those of high molecular weight polymers produced as a result of decreased termination rate due to high viscosity, which was well-known as the Trommsdorff effect.<sup>49</sup> The relatively low PDI in this study could be attributed to the enhanced  $k_p$  and

Table 1 Characterisation results of UHMW polymers synthesised in ILs

Monomer	Solvent	$M_n^a/\text{kDa}$	$M_w^a/\text{kDa}$	PDI <sup>a</sup>	Conversion <sup>b</sup>
MMA	$[\text{C}_1\text{mIm}]\text{[TFSI]}$	1587	2896	1.83	98.0
	$[\text{C}_2\text{mIm}]\text{[TFSI]}$	1280	2377	1.85	98.1
	$[\text{C}_3\text{mIm}]\text{[TFSI]}$	1216	2400	1.97	97.3
	$[\text{C}_4\text{mIm}]\text{[TFSI]}$	1274	2235	1.75	95.9
	$[\text{C}_{12}\text{mIm}]\text{[TFSI]}$	1169	2263	1.94	97.0
	$[\text{C}_2\text{dmIm}]\text{[TFSI]}$	1492	2773	1.86	96.8
	$[\text{C}_2\text{mIm}]\text{[BETI]}$	1473	2693	1.83	97.9
	$[\text{C}_2\text{mIm}]\text{[IM}_{14}\text{]}$	1162	2652	2.28	97.8
	$[\text{C}_2\text{mIm}]\text{[TFSI]}$	988	1700	1.72	95.8
	$[\text{C}_2\text{mIm}]\text{[BETI]}$	1019	1820	1.79	95.9
EMA					
PMA					

<sup>a</sup> Calculated by GPC using PMMA standards (eluent: 10 mM LiBr/DMF). For the characterisation by GPC, polymers were extracted from ILs as follows: UHMW gels were dissolved in acetone, subsequently UHMW polymers were reprecipitated into excess amount of methanol. <sup>b</sup> Calculated from  $^1\text{H}$  NMR. As prepared UHMW gels were dissolved in  $\text{CDCl}_3$ , and the monomer conversion was calculated by comparing the residual vinyl peaks of monomers with polymer peaks.



significantly low concentration of radicals in this polymerization condition that might lower the influence of the termination reaction.

### Rheological properties of UHMW gels

Rheological properties of UHMW gels with different polymers and IL structures were evaluated by frequency sweep measurements at different temperatures ranging from 10 to 120 °C. Next, time-temperature superposition (tTS) master curves for storage modulus ( $G'$ ), loss modulus ( $G''$ ), and  $\tan \delta$  for each UHMW gel was obtained at a reference temperature of 10 °C (Fig. 2 and Fig. S4, ESI†). Note that, although the tTS principle was well applicable, there was a deviation for  $G''$  and  $\tan \delta$  at high frequency region for all UHMW gels. This phenomenon was observed for other ion gel systems also, and it was originated from the relaxation related to solvent viscosity.<sup>50–52</sup>

In all systems, a rubbery plateau region was observed over a wide frequency range; crossover of  $G'$  and  $G''$  and subsequent terminal relaxation ( $G' \propto \omega^2$  and  $G'' \propto \omega$ ) did not appear at least within this measurement condition, which indicates that UHMW gels behave as viscoelastic solids over very long timescales. This is due to the significantly slow relaxation of physical crosslinking owing to the extremely high molecular weight of the polymers, as the reptation theory suggests that the relaxation time of polymer entanglement correlates with the 3rd power of molecular weight. Even at frequency sweep measurement at 120 °C,  $G'$  was higher than  $G''$  for all frequency region, indicating excellent mechanical robustness against temperature for UHMW gels (Fig. S5, ESI†).

### Tensile properties of UHMW gels

Fig. 3 shows the uniaxial tensile stress–strain curves for UHMW gels obtained with various polymer/IL combinations.

The mechanical strength of all the UHMW gels was much better than that of a conventional PMMA ion gel using a chemical crosslinker, which was previously reported as a comparison of the tetra-branched PEO-based stretchable ion gels.<sup>53</sup> Tensile properties and glass transition temperatures ( $T_g$ s) of UHMW gels are summarised in Table S2 (ESI†). Since the polymer/IL combinations used in this study are all compatible systems, the  $T_g$  of the ion gel can be determined by DSC measurements as a single point (Fig. S6, ESI†). When the cation structures of UHMW gels were changed (Fig. 3a), Young's modulus ( $E$ ) varied from 94 to 167 kPa, and the fracture stress and strain varied from 182 to 353 kPa and 578 to 853%, respectively. On the other hand, when the anion and polymer structures were changed (Fig. 3b), the stress–strain curves changed much significantly compared to changes in the UHMW gels with different cation structures. In particular, UHMW gels from PMMA/[C<sub>2</sub>mIm]–[BETI] and PMMA/[C<sub>2</sub>mIm][IM<sub>14</sub>] exhibited significantly high fracture stress. One of the possible reasons for high mechanical strength in the PMMA/[C<sub>2</sub>mIm]–[BETI] UHMW gel could be attributed to its relatively high  $T_g$  of –57.2 °C (Table S2, ESI†), which possibly contributed to stresses associated with micro-Brownian motion of polymer segment. Note that the tensile speed in this study is 10 cm min<sup>–1</sup>, where the correlation between  $T_g$  and  $E$  was observed (Fig. S7, ESI†). This could be attributed to the influence of the transition from rubbery plateau to glassy region at this timescale. In contrast, the PMMA/[C<sub>2</sub>mIm][IM<sub>14</sub>] UHMW gel showed the upward deviation from the correlation between  $T_g$  and  $E$ . Furthermore, it exhibited the highest fracture stress among UHMW gels in this study despite possessing relatively low  $T_g$  of –69.7 °C. A recent study suggested that nanophase separation occurred in [C<sub>2</sub>mIm][IM<sub>14</sub>] due to the aggregation of the long perfluoroalkyl chains of [IM<sub>14</sub>] anions.<sup>54</sup> Therefore, the nanophase separated structure in the

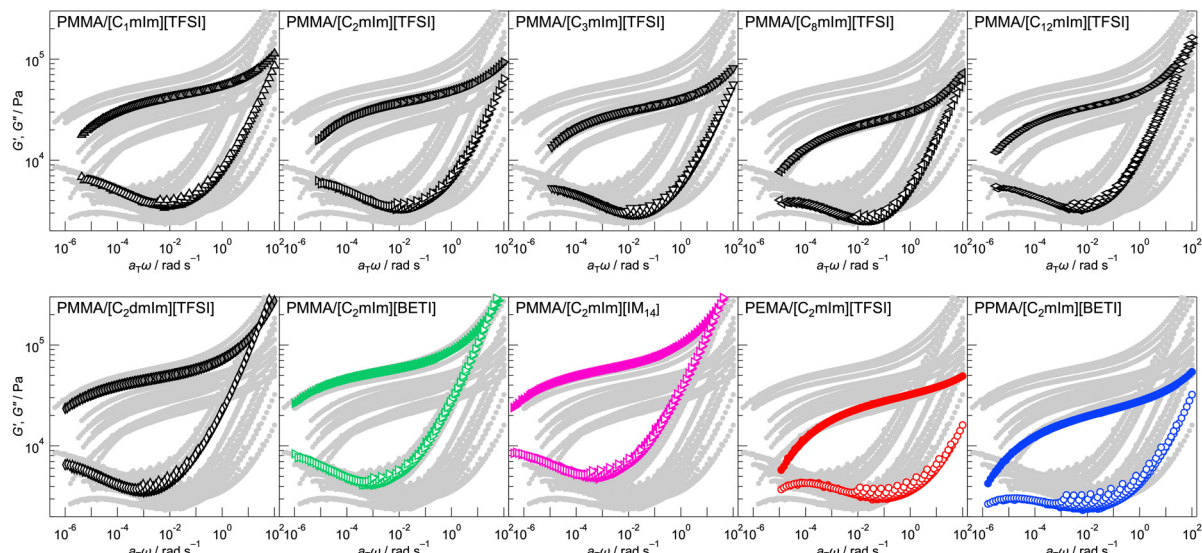


Fig. 2 Time-temperature superposition (tTS) master curves of  $G'$  (closed symbols) and  $G''$  (open symbols) for UHMW gels with different combinations of polymers and ILs constructed from the frequency sweep measurements at the temperature range from 10 to 120 °C. The reference temperature = 10 °C. In each graph, only the master curve of interest is highlighted, while the other curves are shown in grey to clarify their relative positions.



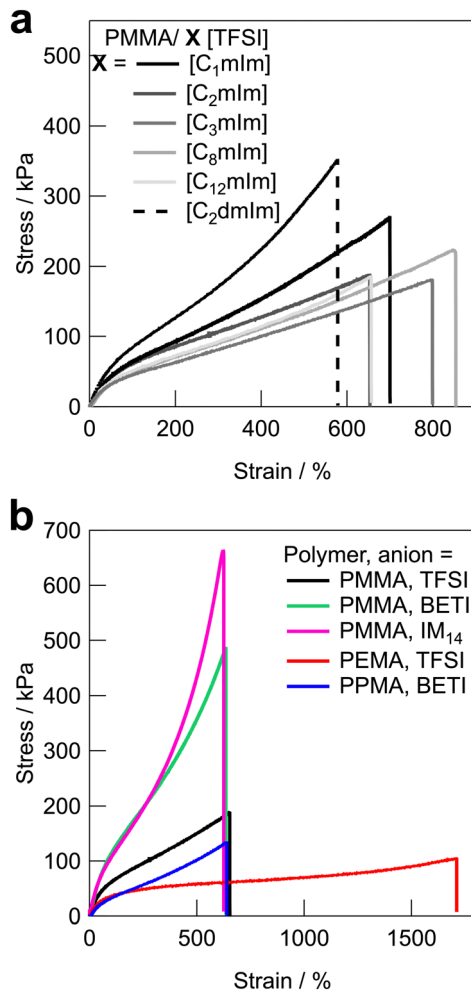


Fig. 3 Tensile stress–strain curves for UHMW gels with (a) different cation structures with the fixed polymer and anion structures (PMMA and [TFSI] anion), and (b) different polymer and anion structures with the fixed cation structure ( $[C_2mIm]$  cation).

UHMW gel might affect the mechanical properties of the UHMW gel. Although the detailed mechanism is still unclear, this result implies that the mechanical strength of UHMW gels could be further enhanced by properly controlling the nanophase separation structure in the gel. On the other hand, although the PMMA/[ $C_{12}mIm$ ][TFSI] UHMW gel had relatively high  $T_g$  of  $-56.1$  °C almost the same as that of PMMA/[ $C_2mIm$ ][BETI], it showed much lower Young's modulus and fracture stress than those of PMMA/[ $C_2mIm$ ][BETI]. The downward deviation from the correlation between  $T_g$  and  $E$  was also confirmed in Fig. S7 (ESI<sup>†</sup>). The long alkyl chain of  $[C_{12}mIm]$  cation is known to cause polar–apolar nanophase separation,<sup>55</sup> suggesting that in this case nanophase separation might have a negative effect on mechanical properties. It is also possible that, although the PMMA/[ $C_2mIm$ ][TFSI] UHMW gel is transparent and thus PMMA and  $[C_2mIm]$ [TFSI] are macroscopically compatible, the low affinity between the IL and the polymer on the microscopic scale may have an influence on the mechanical properties, which will be discussed later in the molecular dynamics (MD) simulations section. When the polymer structure was

changed from PMMA, it was observed that the PEPA/[ $C_2mIm$ ][TFSI] UHMW gel had the highest fracture strain and the lowest fracture stress among all UHMW gels. This can be attributed to the lowest  $T_g$  of  $-78.7$  °C and slightly lower molecular weight of PEPA/[ $C_2mIm$ ][TFSI] gel compared to other systems (Table 1 and Fig. S2, ESI<sup>†</sup>), which may lead to low mechanical strength and large elongation at break due to the disentanglement of polymer chains under the timescale of the tensile test.

### Self-healing ability of UHMW gels

Next, we evaluated the self-healing behaviour of UHMW gels from tensile tests. Our previous study revealed that the PMMA/[ $C_2mIm$ ][TFSI] UHMW gel showed novel self-healing ability at room temperature despite the fact that UHMW gel had a quite long relaxation time owing to the entanglement of UHMW polymers, whereas the PEPA/[ $C_2mIm$ ][TFSI] UHMW gel showed lower healing efficiency than that of PMMA/[ $C_2mIm$ ][TFSI].<sup>42</sup> This result implied that the self-healing ability was dependent on the chemical structure of the UHMW gel. However, detailed study to understand the relationship between polymer and IL chemical structures and self-healing ability of the UHMW gels has not been performed so far. Fig. 4 shows the tensile stress–strain curves for the pristine and healed ion gels for each UHMW gel with different polymer and IL structures. For the preparation of healed ion gel samples, dumbbell-shaped ion gels were cut at the centre, after that the two cut surfaces were brought into contact immediately and allowed to heal at room temperature for 24 h. In contrast to conventional chemically cross-linked ion gels with bifunctional chemical crosslinkers that had no self-healing ability, all UHMW gels exhibited self-healing properties, although there were differences in healing efficiencies. For UHMW gels with different cation structures, the healing efficiency reached to 100% for PMMA/[ $C_2mIm$ ][TFSI] and PMMA/[ $C_3mIm$ ][TFSI], whereas complete healing was not observed for other gels. When the anion structure was varied from TFSI to BETI and IM<sub>14</sub>, the healing efficiency was decreased to less than 40%. One of the possible reason of low healing efficiency for the PMMA/[ $C_2mIm$ ][BETI] and PMMA/[ $C_2mIm$ ][IM<sub>14</sub>] could be attributed to the trade-off relationship between self-healing ability and stiffness. The higher Young's modulus is related to lower dynamic properties of physical cross-linking by entanglement, which might lead to low self-healing properties. The trade-off relationship was also reported in other self-healable polymeric materials.<sup>32,56</sup>

Fig. 5 shows the relationship between Young's modulus and healing efficiency for UHMW gels. While several UHMW gels obeyed the trade-off relationship between Young's modulus and healing efficiency shown by the dotted straight line, some UHMW gels deviated upward and downward from the trade-off line. The PMMA/[ $C_2mIm$ ][TFSI] UHMW gel exhibited upward deviation from the trade-off line, indicating that there exists a driving force which accelerates the reformation of entanglement across the cut surfaces. On the other hand, PMMA/[ $C_{12}mIm$ ][TFSI], PEPA/[ $C_2mIm$ ][TFSI], and PPMA/[ $C_2mIm$ ][BETI] UHMW gels deviated significantly downward from the



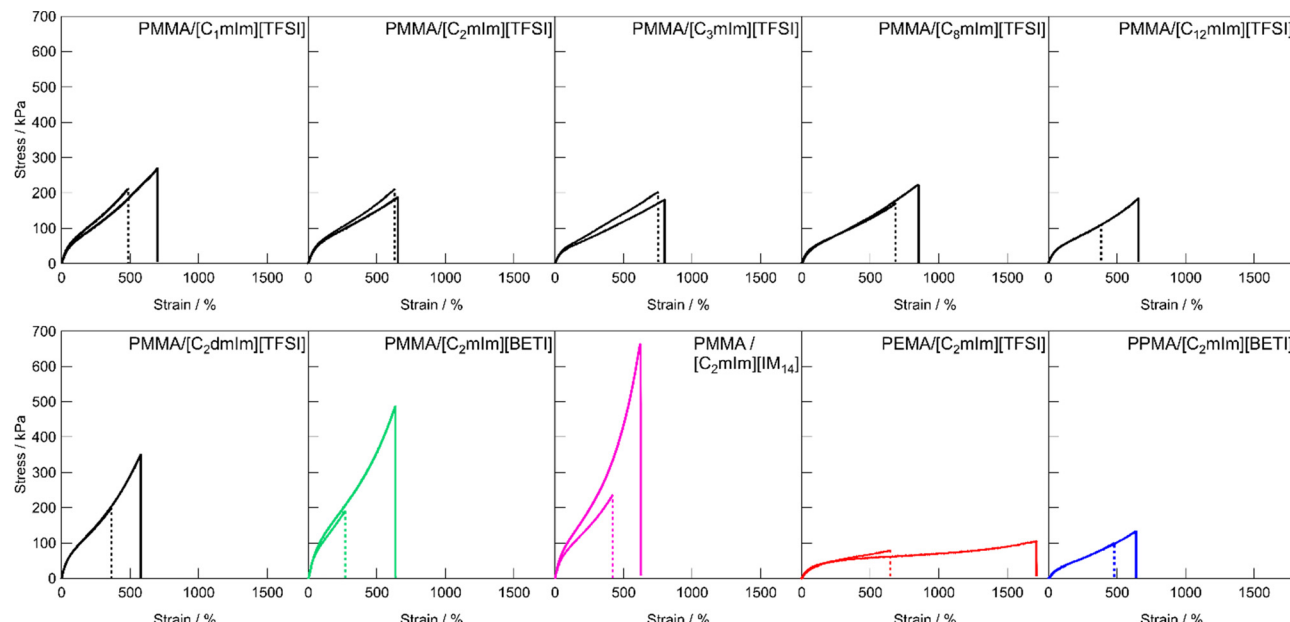


Fig. 4 Tensile stress–strain curves for pristine and healed UHMW gels with different polymer and IL structures. Solid and dashed lines indicate pristine and healed samples, respectively.

trade-off line. Notably, although the tensile stress–strain curves for pristine PMMA/[C<sub>12</sub>mIm][TFSI] and PMMA/[C<sub>2</sub>mIm][TFSI] UHMW gels were nearly identical, the healing efficiencies were quite different (Fig. S8, ESI<sup>†</sup>).

These results suggest that the self-healing behaviour of UHMW gels is not governed only by the trade-off relationship between the stiffness and the healing efficiency, other factors also contribute to the self-healing phenomena. In our previous study, MD simulations for PMMA/[C<sub>2</sub>mIm][TFSI] and PEMA/[C<sub>2</sub>mIm][TFSI] systems suggested that PMMA and PEMA showed different solvation structure in [C<sub>2</sub>mIm][TFSI].<sup>42</sup> Therefore, the deviation from the trade-off relationship might have originated from the difference in the microscopic interaction among polymers and IL cations and anions, which would lead to the acceleration or deceleration of reformation of entanglement across cut surfaces.

### MD simulations for different polymer/IL systems

To further understand the polymer–IL and polymer–polymer interactions at the microscopic scale, all-atom MD simulations and detailed statistical analysis were performed on the PMMA/[C<sub>2</sub>mIm][TFSI], PMMA/[C<sub>12</sub>mIm][TFSI], and PEMA/[C<sub>2</sub>mIm][TFSI] systems, in which healing efficiency was quite different. The simulations were carried out in an NTP ensemble. Density values derived from MD simulations are in good agreement with actual systems, indicating the validity of simulated polymer/IL structures (Table S1, ESI<sup>†</sup>). The radial distribution functions for all atoms ( $G_{\text{total}}(r)$  as an  $r$ -weighted difference form,  $r^2[G(r) - 1]$ , is shown in Fig. S9, ESI<sup>†</sup>) were calculated from the trajectory data obtained from the 500 ps calculations. The radial distribution function consists of intramolecular and intermolecular correlations for each component. The intramolecular radial distribution function for the polymer ( $G_{\text{intra}}(r)$ )

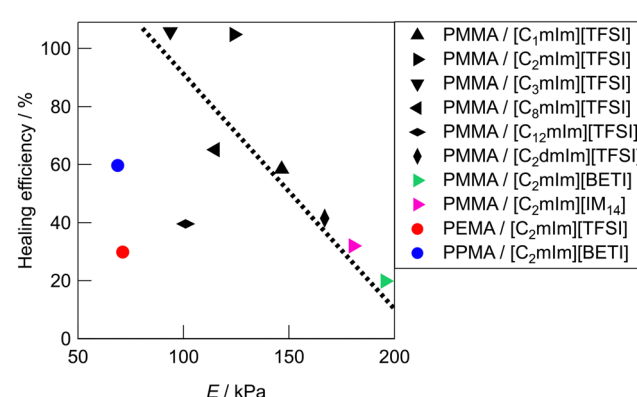


Fig. 5 Relationship between the Young's modulus ( $E$ ) and the healing efficiency of UHMW gels. The dotted line is a guide to the eye. The healing efficiency was defined as the ratios of mechanical toughness (the area under the stress–strain curve) between pristine and healed gels.

is shown in Fig. 6a. The jagged peaks appearing below 5 Å corresponded to correlations between directly bonded atoms or their neighbours in the polymer molecule. On the other hand, the broad peak with a peak top near 10 Å can be attributed to the summation of atomic correlations within non-bonded atoms. Therefore, the broadness of these peaks is an indicator for the degree of expansion of individual polymers solvated by ILs. It was observed that the PMMA/[C<sub>2</sub>mIm][TFSI] system had a broader peak than those of PMMA/[C<sub>12</sub>mIm][TFSI] and PEMA/[C<sub>2</sub>mIm][TFSI] systems, suggesting that the polymer conformation was most extended in the PMMA/[C<sub>2</sub>mIm][TFSI] system compared to PMMA/[C<sub>12</sub>mIm][TFSI] and PEMA/[C<sub>2</sub>mIm][TFSI] systems. In fact, the  $R_g$  calculated from the intramolecular radial distribution function was 14.2, 13.1, and 12.3 Å for



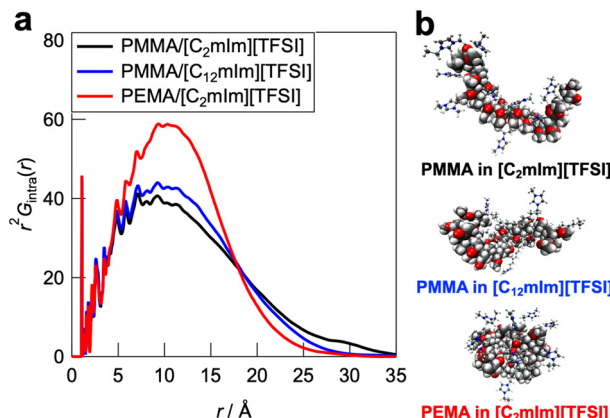


Fig. 6 (a) Intramolecular contribution of the radial distribution function ( $G_{\text{intra}}(r)$ ) for the PMMA/[C<sub>2</sub>mIm][TFSI], PMMA/[C<sub>12</sub>mIm][TFSI], and PEMA/[C<sub>2</sub>mIm][TFSI] systems. (b) Representative snapshot of individual polymer conformations in ILs extracted from the MD simulation boxes.

PMMA/[C<sub>2</sub>mIm][TFSI], PMMA/[C<sub>12</sub>mIm][TFSI], and PEMA/[C<sub>2</sub>mIm][TFSI], respectively. Fig. 6b shows representative polymer conformations for three systems.

Furthermore, to obtain insight into the polymer–polymer interactions in ILs, we evaluated the intermolecular radial distribution function ( $G_{\text{poly-poly}}(r)$ ) that represents the correlation between the different polymer chains (Fig. 7a). In the PMMA/[C<sub>2</sub>mIm][TFSI] system, the upturn of the intensity was observed around 6 Å, which indicates that the correlation between polymers starts to occur from this distance. On the other hand, in the PMMA/[C<sub>12</sub>mIm][TFSI] and the PEMA/[C<sub>2</sub>mIm][TFSI] systems, the upturns were observed from around 9 and 12 Å, respectively, which were farther than that of the PMMA/[C<sub>2</sub>mIm][TFSI] system. In addition, the depth of the convex curve for the PMMA/[C<sub>12</sub>mIm][TFSI] and the PEMA/[C<sub>2</sub>mIm][TFSI] systems in Fig. 7a was much deeper than that of the PMMA/[C<sub>2</sub>mIm][TFSI] system, suggesting the probability of having different polymer chains in the proximity is low. These results suggest that in the PMMA/[C<sub>2</sub>mIm][TFSI] system, individual polymer chain has a more

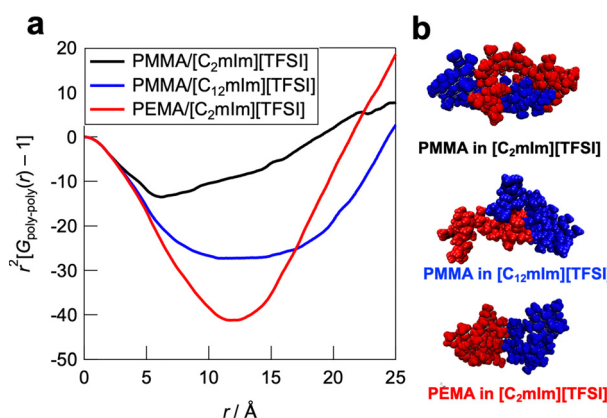


Fig. 7 (a) Intermolecular contribution of the radial distribution function ( $G_{\text{poly-poly}}(r)$ ) for the PMMA/[C<sub>2</sub>mIm][TFSI], PMMA/[C<sub>12</sub>mIm][TFSI], and PEMA/[C<sub>2</sub>mIm][TFSI] systems. (b) Representative snapshot of two adjacent polymers in ILs extracted from the MD simulation boxes.

extended conformation in the IL, and different polymer chains are more likely to exist in the vicinity (Fig. 7b). Therefore, even at the cut surfaces of the PMMA/[C<sub>2</sub>mIm][TFSI] UHMW gel, the polymer chains may take an expanded form and different polymer chains are more likely to be present in the vicinity, which facilitates the reformation of entanglement, resulting in the excellent self-healing ability. However, it should be noted that in this study, MD calculations were performed for polymers with a low polymerisation degree of 20 mers, which is quite different from the degree of polymerisation for UHMW polymers ( $>10\,000$ ). Therefore, further studies to clarify the polymer conformations and interpolymer interactions in UHMW gels more accurately by combining the large-scale MD calculations at polymerisation degree that exceeds the entanglement molecular weight is under progress along with high-energy X-ray total scattering experiments.<sup>57</sup> In addition, the influence of the difference in the polymer conformation calculated from MD simulations on the linear viscoelastic master curves is still elusive. Nonlinear rheological measurements are now underway to investigate the effect of the difference in the polymer conformation in ILs on physical properties.

## Conclusions

In conclusion, UHMW gels, which were formed solely by physical entanglements of *in situ* polymerised UHMW polymers in ILs, were fabricated from various IL/monomer structures. In the rheological measurements, all UHMW gels exhibited a wide rubbery plateau without crossover of  $G'$  and  $G''$  over wide frequency and temperature region, indicating that UHMW gels can maintain solid-like integrity over long timescales. Tensile tests suggested that stress–strain curves of UHMW gels were significantly dependent on the IL/polymer chemical structures. In particular, the PMMA/[C<sub>2</sub>mIm][IM<sub>14</sub>] UHMW gel showed the highest fracture stress despite relatively low  $T_g$ , implying that the nanophase separation of long perfluoroalkyl chain might affect the tensile property of UHMW gels. Self-healing properties of UHMW gels evaluated by tensile tests revealed the trade-off relationship between the stiffness and the healing efficiency, whereas some UHMW gels deviated from the trade-off line. MD simulations for different polymer/IL structures revealed that the intramolecular polymer conformations as well as interpolymer interactions were dependent on the polymer/IL structures. In the combination of polymer/IL for the UHMW gel that shows high self-healing efficiency, the polymer chain can take an extended conformation in the IL, and that different polymer chains are likely to be present in close proximity to each other. In the field of polymer gel science, “the style of cross-linking”, “polymer network”, and “solvent” are known as the three major components of gels. So far, efforts have been made to realise effects of the style of cross-linking and polymer network towards unprecedented functions and excellent mechanical properties of gels. However, solvents engineering of gels has never received any special attention. The ability to control the mechanical properties of gels by changing the



solvent structure is a unique strategy that is not found in other gels such as hydrogels and organogels. Further, our insights into the effects of the chemical structure on the mechanical and self-healing properties of UHMW gels provide a useful basis for the development of a new class of ion gels for wearable and flexible devices with desirable physicochemical properties.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by JST, PRESTO, Japan (Grant Number: JPMJPR2196 to R. T.), and JSPS KAKENHI (Grant Number: 20K15349 to R. T., 20H02804, and 20K21229 to T. U.).

## References

- 1 A. Nathan, A. Ahnood, M. T. Cole, S. Lee, Y. Suzuki, P. Hirralal, F. Bonaccorso, T. Hasan, L. Garcia-Gancedo, A. Dyadyusha, S. Haque, P. Andrew, S. Hofmann, J. Moultrie, D. Chu, A. J. Flewitt, A. C. Ferrari, M. J. Kelly, J. Robertson, G. A. J. Amaralunga and W. I. Milne, *Proc. IEEE*, 2012, **100**, 1486–1517.
- 2 M. Stoppa and A. Chiolerio, *Sensors*, 2014, **14**, 11957–11992.
- 3 W. Liu, M.-S. Song, B. Kong and Y. Cui, *Adv. Mater.*, 2017, **29**, 1603436.
- 4 T. Sekitani, Y. Noguchi, K. Hata, T. Fukushima, T. Aida and T. Someya, *Science*, 2008, **321**, 1468–1472.
- 5 Y. Wang, C. Zhu, R. Pfattner, H. Yan, L. Jin, S. Chen, F. Molina-Lopez, F. Lissel, J. Liu, N. I. Rabiah, Z. Chen, J. W. Chung, C. Linder, M. F. Toney, B. Murmann and Z. Bao, *Sci. Adv.*, 2017, **3**, e1602076.
- 6 T. Q. Trung and N.-E. Lee, *Adv. Mater.*, 2017, **29**, 1603167.
- 7 H. Tran, V. R. Feig, K. Liu, Y. Zheng and Z. Bao, *Macromolecules*, 2019, **52**, 3965–3974.
- 8 K. Kim, Y. Park, B. G. Hyun, M. Choi and J. Park, *Adv. Mater.*, 2019, **31**, 1804690.
- 9 B. Wang and A. Facchetti, *Adv. Mater.*, 2019, **31**, 1901408.
- 10 M. L. Hammock, A. Chortos, B. C.-K. Tee, J. B.-H. Tok and Z. Bao, *Adv. Mater.*, 2013, **25**, 5997–6038.
- 11 A. Manuel Stephan, *Eur. Polym. J.*, 2006, **42**, 21–42.
- 12 M. Khatib, O. Zohar and H. Haick, *Adv. Mater.*, 2021, **33**, 2004190.
- 13 S. H. Kim, K. Hong, W. Xie, K. H. Lee, S. Zhang, T. P. Lodge and C. D. Frisbie, *Adv. Mater.*, 2013, **25**, 1822–1846.
- 14 G. A. Snook, P. Kao and A. S. Best, *J. Power Sources*, 2011, **196**, 1–12.
- 15 C. Jo, D. Pugal, I.-K. Oh, K. J. Kim and K. Asaka, *Prog. Polym. Sci.*, 2013, **38**, 1037–1066.
- 16 J. Mindemark, M. J. Lacey, T. Bowden and D. Brandell, *Prog. Polym. Sci.*, 2018, **81**, 114–143.
- 17 C. Yang and Z. Suo, *Nat. Rev. Mater.*, 2018, **3**, 125–142.
- 18 T. Ueki and M. Watanabe, *Macromolecules*, 2008, **41**, 3739–3749.
- 19 J. Le Bideau, L. Viau and A. Vioux, *Chem. Soc. Rev.*, 2011, **40**, 907–925.
- 20 T. P. Lodge and T. Ueki, *Acc. Chem. Res.*, 2016, **49**, 2107–2114.
- 21 Y. Kitazawa, K. Ueno and M. Watanabe, *Chem. Rec.*, 2018, **18**, 391–409.
- 22 C. Creton, *Macromolecules*, 2017, **50**, 8297–8316.
- 23 C. W. Peak, J. J. Wilker and G. Schmidt, *Colloid Polym. Sci.*, 2013, **291**, 2031–2047.
- 24 J.-Y. Sun, X. Zhao, W. R. K. Illeperuma, O. Chaudhuri, K. H. Oh, D. J. Mooney, J. J. Vlassak and Z. Suo, *Nature*, 2012, **489**, 133–136.
- 25 T. Matsuda, R. Kawakami, R. Namba, T. Nakajima and J. P. Gong, *Science*, 2019, **363**, 504–508.
- 26 C. Norioka, Y. Inamoto, C. Hajime, A. Kawamura and T. Miyata, *NPG Asia Mater.*, 2021, **13**, 34.
- 27 C. Liu, N. Morimoto, L. Jiang, S. Kawahara, T. Noritomi, H. Yokoyama, K. Mayumi and K. Ito, *Science*, 2021, **372**, 1078–1081.
- 28 T. Fujiyabu, N. Sakumichi, T. Katashima, C. Liu, K. Mayumi, U. Il Chung and T. Sakai, *Sci. Adv.*, 2022, **8**, 10.
- 29 A. Harada, Y. Takashima and M. Nakahata, *Acc. Chem. Res.*, 2014, **47**, 2128–2140.
- 30 K. Imato, T. Ohishi, M. Nishihara, A. Takahara and H. Otsuka, *J. Am. Chem. Soc.*, 2014, **136**, 11839–11845.
- 31 A. Campanella, D. Döhler and W. H. Binder, *Macromol. Rapid Commun.*, 2018, **39**, 1700739.
- 32 T. L. Sun, T. Kurokawa, S. Kuroda, A. Bin Ihsan, T. Akasaki, K. Sato, M. A. Haque, T. Nakajima and J. P. Gong, *Nat. Mater.*, 2013, **12**, 932–937.
- 33 D. L. Taylor and M. in het Panhuis, *Adv. Mater.*, 2016, **28**, 9060–9093.
- 34 Z. Deng, H. Wang, P. X. Ma and B. Guo, *Nanoscale*, 2020, **12**, 1224–1246.
- 35 Y. Cao, T. G. Morrissey, E. Acome, S. I. Allec, B. M. Wong, C. Keplinger and C. Wang, *Adv. Mater.*, 2017, **29**, 1605099.
- 36 E. Kamio, T. Yasui, Y. Iida, J. P. Gong and H. Matsuyama, *Adv. Mater.*, 2017, **29**, 1704118.
- 37 R. Tamate, K. Hashimoto, T. Horii, M. Hirasawa, X. Li, M. Shibayama and M. Watanabe, *Adv. Mater.*, 2018, **30**, 1802792.
- 38 R. Tamate and M. Watanabe, *Sci. Technol. Adv. Mater.*, 2020, **21**, 388–401.
- 39 R. Tamate, *Polym. J.*, 2021, **53**, 789–798.
- 40 L. Xu, Z. Huang, Z. Deng, Z. Du, T. L. Sun, Z.-H. Guo and K. Yue, *Adv. Mater.*, 2021, 2105306.
- 41 W. Li, L. Li, S. Zheng, Z. Liu, X. Zou, Z. Sun, J. Guo and F. Yan, *Adv. Mater.*, 2022, 2203049.
- 42 Y. Kamiyama, R. Tamate, T. Hiroi, S. Samitsu, K. Fujii and T. Ueki, *Sci. Adv.*, 2022, **8**, eadd0226.
- 43 A. Noda, K. Hayamizu and M. Watanabe, *J. Phys. Chem. B*, 2001, **105**, 4603–4610.
- 44 W. L. Jorgensen, D. S. Maxwell and J. Tirado-Rives, *J. Am. Chem. Soc.*, 1996, **118**, 11225–11236.
- 45 W. D. Cornell, P. Cieplak, C. I. Bayly, I. R. Gould, K. M. Merz, D. M. Ferguson, D. C. Spellmeyer, T. Fox, J. W. Caldwell and P. A. Kollman, *J. Am. Chem. Soc.*, 1995, **117**, 5179–5197.



46 J. N. Canongia Lopes and A. A. H. Pádua, *J. Phys. Chem. B*, 2006, **110**, 19586–19592.

47 S. Harrisson, S. R. Mackenzie and D. M. Haddleton, *Chem. Commun.*, 2002, 2850–2851.

48 K. Low, L. Wylie, D. L. A. Scarborough and E. I. Izgorodina, *Chem. Commun.*, 2018, **54**, 11226–11243.

49 Y. Suzuki, Y. Shinagawa, E. Kato, R. Mishima, K. Fukao and A. Matsumoto, *Macromolecules*, 2021, **54**, 3293–3303.

50 Y. He, P. G. Boswell, P. Bühlmann and T. P. Lodge, *J. Phys. Chem. B*, 2007, **111**, 4645–4652.

51 A. Noro, Y. Matsushita and T. P. Lodge, *Macromolecules*, 2008, **41**, 5839–5844.

52 X. Ma, R. Usui, Y. Kitazawa, R. Tamate, H. Kokubo and M. Watanabe, *Macromolecules*, 2017, **50**, 6788–6795.

53 S. Ishii, H. Kokubo, K. Hashimoto, S. Imaizumi and M. Watanabe, *Macromolecules*, 2017, **50**, 2906–2915.

54 F. Lo Celso, G. B. Appeteccchi, E. Simonetti, M. Zhao, E. W. Castner, U. Keiderling, L. Gontrani, A. Triolo and O. Russina, *Front. Chem.*, 2019, **7**, 285.

55 K. Fujii, R. Kanzaki, T. Takamuku, Y. Kameda, S. Kohara, M. Kanakubo, M. Shibayama, S. Ishiguro and Y. Umebayashi, *J. Chem. Phys.*, 2011, **135**, 244502.

56 Z. Wang, S. Gangarapu, J. Escorihuela, G. Fei, H. Zuilhof and H. Xia, *J. Mater. Chem. A*, 2019, **7**, 15933–15943.

57 K. Fujii, T. Ueki, K. Hashimoto, Y. Kobayashi, Y. Kitazawa, K. Hirosawa, M. Matsugami, K. Ohara, M. Watanabe and M. Shibayama, *Macromolecules*, 2017, **50**, 4780–4786.

