Cross-Linking and Evaluation of the Thermo-Mechanical Behavior of Epoxy Based Poly(ionic Liquid) Thermosets

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Abstract: Poly(ionic liquids) (PILs) and ionenes are polymers containing ionic groups in their repeating units. The unique properties of these polymers render them as interesting candidates for a variety of applications, such as gas separation membranes and polyelectrolytes. Due to the vast number of possible structures, numerous synthesis protocols to produce monomers with different functional groups for task-specific PILs are reported in literature. A difunctional epoxy-IL resin was synthesized and cured with multifunctional amine and anhydride hardeners and the thermal and thermomechanical properties of the networks were assessed via differential scanning calorimetry and dynamic mechanical analysis. By the selection of suitable hardeners, the glass transition onset temperature ($T_{g,\text{onset}}$) of the resulting networks was varied between 18 °C and 99 °C. Copolymerization of epoxy-IL with diglycidyl ether of bisphenol A (DGEBA) led to a further increase of the $T_{g,\text{onset}}$. The results demonstrate the potential of epoxy chemistry for tailorable PIL networks, where the hardener takes the place of the ligands without requiring an additional synthesis step and can be chosen from a broad range of commercially available compounds.

Keywords: ionic liquid; epoxy; ionene; poly(ionic liquid); thermoset

1. Introduction

Ionic liquids (ILs), which are generally defined as salts with melting points below 100 °C, have attracted increased attention in recent years in research and industry [1–3]. They consist of a bulky organic cation and an organic or inorganic anion [4,5], decreasing the packing density of the molecules and reducing the melting temperature ($T_m$) compared to conventional inorganic salts [6,7]. Asymmetrical cations and weaker interactions between the ions usually shift the transition temperatures to lower values [8], in many cases even well below room temperature (RT) [9].

The unique properties of ILs, such as low flammability, negligible vapor pressure, moisture resistance, high thermal stability [10], reusability [11], ion conductivity [12], as well as catalytic activity [13–15], combined with their high tunability and wide range of available ILs with a vast number of possible anion-cation combinations give rise to many possible applications [10]. Examples are their use as solvents [16], latent curing agents [17], phase transfer catalysts [18], dispersing agents [19], and porogens [20,21], as well as their application in materials for CO$_2$ capture [22], magneto-rheological fluids [23], and self-assembly of polymers [24]. Other reports include the formation of ionogels produced from ionic liquids and a suitable gelator, exhibiting high ionic conductivity and tunable properties that can be used as quasi-solid electrolytes [25,26].

A further advantageous property of ILs is that they can also be functionalized with a variety of different reactive groups to act as non-volatile monomers for the realization of poly(ionic liquids) (PILs). Depending on their structure and position of the ionic moieties in the polymer chains, different terms are used to describe the resulting materials. An overview of the chemical structure of different poly(ionic liquids) is provided in Figure 1. The properties of the resulting PILs are greatly influenced by their respective ion pairs.
Differences in the glass transition temperature ($T_g$) of 50 °C or more were reported for PILs by exchanging the respective counter-ions of the poly(ionic liquids) [27,28]. Thus, the properties of soluble PILs can be further tuned by anion exchange, even after polymerization [28–30]. Exchanging the anion can also be utilized to adjust the solubility of the polymer [31,32] and the mechanical properties [33]. Depending on the monomer structure and the ion pair, PILs can be designed to be hydrophobic [34,35], antibacterial [36], and non-flammable [37] polymers. While the anion selection offers many opportunities to create task specific materials, some anions result in a significant increase in brittleness and the PILs cannot be processed as a consequence [38–40].

![Figure 1](image.jpg)

**Figure 1.** Schematic representation of different poly(ionic liquid) (PIL) structures and terminology thereof.

In many cases, PILs exhibit high CO$_2$ solubility and solubility selectivity [41,42], especially in combination with free ionic liquids as solid-liquid composites [42–44]. This renders them very interesting candidates for gas separation membranes. The reported loadings of free ILs in the PIL membranes are often above 40 wt.% [45,46], whereas the PIL matrix increases the maximum filler capacity by intermolecular bonding and prevents leakage [28]. Moreover, PILs have shown to be potential candidates as catalysts for solvent free reactions [47,48]. Another interesting property of PILs is their ionic conductivity for applications as polyelectrolytes [49].

Several different approaches to polymerizing ILs have been reported in literature: (i) As pendant group of a polymerizable monomer [50,51], (ii) block copolymers of IL monomers with non-ionic monomers [52,53], (iii) by protonation of amine functional groups and subsequent cross-linking with an epoxy resin [54], (iv) or by using cross-linkable ionic liquid monomers [46]. While polymerizable cations are more common, it is also possible to functionalize the anion or even both ions [55,56]. The majority of the reported PILs are obtained from monomers with only one polymerizable group with the ionic moieties as pendant groups [57–59]. By polymerizing these monomers, a polyelectrolyte is formed.

Similar structures can be produced by grafting an ionic group to the backbone as a side chain [57,60–62] or via polymerization of a dihalide with bisimidazole molecules [63–66]. The latter results in ionenes, which are polymers with ionic moieties in their backbone [67]. PILs and ionenes are generally very similar and share the same field of applications [68]. However, while PILs are often based on mono-functional alkenyl moieties, and therefore, possess an aliphatic, non-ionic backbone, literature reports on ionenes increasingly focus on the introduction of ionic groups in high performance polymers such as polyimides [63,69] or aromatic polyamides [64,70]. Furthermore, the position of the ions influences the ion aggregation [71] and an increased ion conductivity was found for ionenes in comparison to a polyelectrolyte [72].

IL-based ionenes can also be obtained from multifunctional ILs [73]. Difunctional ILs equipped with epoxy-functional groups (epoxy-IL) represent a very versatile class of materials. The oxirane ring exhibits reactivity towards a multitude of species and can be used for polymerization with amines, phenols, thiols, isocyanates or carboxylic acids following a step-growth mechanism. However, epoxies can also be polymerized by a chain-growth reaction mechanism, either as homopolymerisation or as alternating copolymerization with anhydrides [74,75]. Epoxy based polymers with thermoplastic properties can be produced by the homopolymerisation of monofunctional epoxy resins or by curing a difunctional epoxy resin with a difunctional hardener [76]. If the epoxy
resin or the hardener have a functionality of more than two, cross-linked PILs/ionenes are acquired.

Most reported epoxy functional ILs are based on imidazolium cations, due to their high thermal stability [77], combined with bis(trifluoromethane)sulfonimide (Tf$_2$N$^-$) anions, which ensure that even the large aromatic monomer molecules are liquid at lower temperatures [78], and yield hydrophobic polymers [34,36,79]. In contrast, epoxies based on the widely used diglycidyl ether of bisphenol A (DGEBA) tend to be more hydrophilic [34,80]. The starting compounds are usually functionalized using halides or tosylates, either by direct epoxidation with epichlorohydrin/epibromohydrin [81–85] or by addition of an alkenyl group and subsequent oxidation of the double bond [42,46,49,78].

Table 1 summarizes literature examples of epoxy-IL monomers and their respective glass transition temperature after polymerization. McDanel et al. [42,46] synthesized PILs 1A–D for CO$_2$ capture. The networks were produced by curing the respective monomers with tri- or hexa-functional amine hardeners. Through variation of the hardeners and their stoichiometry, the authors were able to significantly influence the CO$_2$ uptake of the materials as well as the respective glass transition temperatures.

Radchenko et al. [34] synthesized the aromatic PIL 2 that can be used for shape memory applications due to its two relaxation temperatures at 60 °C and 109 °C. The authors reported comparably high mechanical properties, with a Young’s modulus of 1.63 GPa, a yield strength of 96 MPa, 28 % strain at break as well as a storage modulus of 1.7 GPa at room temperature. In another publication, the authors produced the three monomer materials 3A–C, which have a high potential for 3D-printing applications in combination with diaryliodonium hexafluoroantimonate as the UV or thermally activated initiator. The cationically cured monomer 3B led to a polymeric network with a storage modulus of 1.18 GPa [86].

Livi et al. [36,49] produced PILs from monomers 4A–C for antibacterial coatings or surfaces and reported a biofilm inhibition of more than 80 % for all three polymers. The PILs exhibited rather high glass transition temperatures between 53 °C and 62 °C, considering that an aliphatic hardener was used. The highest value was reported for 4B, which does not contain an ether bond in the epoxy monomer. The storage moduli are similar for all three polymers, with 700 MPa reported for 4A at room temperature [36,49]. Apart from a reduced stiffness of PILs compared to commercially available epoxy monomers, the ionic character is expected to have an influence on the reactivity of functional groups. While the peak of the exotherm of 4A during curing is observed at lower temperatures than that of DGEBA [49], it is less reactive than monomer 4C with its two aromatic ligands. The authors attribute this lower reactivity to the higher distance of the oxirane ring to the cation [36]. The substituents do not only influence the properties of the monomer but also the synthesis. Electron donating groups on aromatic rings reduce the reaction speed during the oxidation of the alkenyl group to yield epoxide rings. The highest effect was observed for ester groups in the synthesis of various monomers [78].
Table 1. Overview of PILs based on epoxy functional imidazolium ionic liquids reported in literature.

| Monomer Structure | $T_f^\circ\text{C}$ | Ref. |
|-------------------|------------------|-----|
| (1A) | [87] | |
| (1B) | 9/17/38 (A) | [42,46] |
| (1C) | −5/12 (B) | |
| (1D) | −18/−27 (C) | |
| (2) | 3/20 (D) | |
| (3A) | 60 & 109 | [34] |
| (3B) | 85 (3B) | [86] |
| (3C) | | |
| (4A) | 55 (A) | [36,49] |
| (4B) | 62 (B) | |
| (4C) | 53 (C) | |
| (5) | 61 | [79] |

$\text{Tf}_2\text{N}^− = \text{Bis(trifluoromethane)sulfonimide.}$

Grugel and co-workers [79] investigated the properties of epoxy-IL resins in detail to examine their suitability as matrix materials in carbon fiber reinforced composites for the storage of liquid hydrogen and for cryogenic applications. The thermoset resulting from curing of monomer 5 is reported to exhibit low flammability, hydrogen permeability and thermal expansion, as well as increased adhesion, lap shear and tensile strength. Increased impact resistance and fracture toughness were observed, especially under cryogenic conditions in comparison to a commercially available DGEBA based epoxy resin. The reported tensile strength of approximately 60 MPa is significantly lower than for PIL 2 but still within the range of technical resins [79]. It has to be noted that the IL was cross-linked with an aromatic amine hardener, while the reference DGEBA resin was cured with polyoxypropylenetetramine, an aliphatic polyether. PIL 5 was exposed to aging tests in space, facing earth for approx. 2 years, in which it went through 12,500 cycles between $−40{\text{ °C}}$ to $40{\text{ °C}}$ under...
the influence of radiation and oxygen. The investigated specimen on aluminum substrates exhibited no cracks, delamination or loss of adhesion after the tests [87]. Furthermore, the material was tested with fiber reinforcements and exhibited increased strain under liquid nitrogen conditions and better adhesion to the fibers compared to a common DGEBA based thermoset. A different crack growth behavior was observed for DGEBA based thermosets and PIL 5. The DGEBA based material exhibited failure along the interface between fiber and epoxy, while the crack propagated through the epoxy matrix for PIL 5 and even plastic deformation before failure was reported [88]. In contrast, PIL 5 did not show any cracks or delamination from temperature cycles between cryogenic and room temperature, and surpassed commercial systems in terms of tensile and bond strength [89].

Another possibility to synthesize IL based epoxy networks is to functionalize the cation with a group that can react with epoxy monomers, such as amines or carboxyl groups. Amine functional ILs can also be obtained by quaternization of an amine curing agent [54], by exchanging the anion to an amino acid, which allows the synthesis of ILs with both functional anions and cations [22,90]. Another route involves the Radziszewski imidazole synthesis using functionalized amines, which is reported for the production of both amine and carboxylic acid functional ILs [91,92].

Jiang et al. [93] synthesized diamine imidazolium hardeners with dicyanamide and Tf$_2$N$^-$ anions and used it to cure a trifunctional epoxy resin. The authors determined the ideal amount of hardener to be 15 wt.% of IL, which is non-stoichiometric. Therefore, a significant number of cross-links can be expected to be based on homopolymerisation, which is likely the reason for the higher glass transition temperature (212 $^\circ$C/196 $^\circ$C) of the network compared to other reported PILs. Apart from a high $T_g$, the thermoset exhibited good mechanical properties with a tensile strength of up to 40 MPa and a strong influence of the anion. Using Tf$_2$N$^-$ as counterion resulted in a lower cross-link density and tensile modulus but increased tensile strength and elongation at break. Interestingly, despite the decreased cross-link density, a higher $T_g$ were reported for the Tf$_2$N$^-$ anion than for the dicyanamide.

Jin et al. [60] reported a seven-fold increase of impact strength upon the addition of a carboxyl functional polyether with pending imidazolium IL groups to a commercial epoxy resin. It has to be noted that the addition of the polyether also introduces long alkyl chains into the network, which might play a significant role in the impact strength enhancement. Apart from epoxy groups, amines can also react with anhydrides to form polyimides. Li, Zhang et al. [73,94,95] produced random and block-copolyimides with mono- and di-cationic amine ILs. The addition of different ratios of the amine IL led to an increase in density while it significantly decreased the molecular weight and the glass transition temperature from initial 296 $^\circ$C to as low as 177 $^\circ$C for 30 mol% of IL.

Due to this high tunability and sheer endless combinations, numerous reports on the synthesis and characterization of task-specific PILs are available in literature. This, in turn, requires custom syntheses for most PILs. Livi and co-workers simulated the properties of homopolymerized epoxy PILs based on the monomer structure to reduce the amount of different syntheses required [96]. In the present work, the versatility and considerable potential of epoxy-functional ILs is further demonstrated via the selection and implementation of different hardeners, resulting in an effective tailoring of the properties of the prepared epoxy-IL networks. Specific focus is laid on a systematic evaluation of the thermal and thermo-mechanical properties of representative epoxy-IL networks, cured with different types of hardeners. For this purpose, an epoxy-functional IL monomer with comparably short side-chains was synthesized and cured with selected multifunctional amine hardeners. The thermal and thermo-mechanical properties of the resulting epoxy-IL networks are examined and compared to a reference DGEBA based epoxy resin as a benchmark. To the best of the authors’ knowledge, this has not been extensively done so far. The potential for a flexible tailoring of the network properties of epoxy-IL resins is further demonstrated by an investigation of resin systems prepared with varying mixing ratios between epoxy-functional IL and DGEBA as a commonly used benchmark resin.
2. Materials and Methods

2.1. Chemicals and Materials

1H-imidazole, 3-chloroperoxybenzoic acid (77%), 3-(aminomethyl)-3,5,5-trimethylcyclohexanamine (IPD), 4,4’-diaminodiphenylsulfone (DDS), and trimethyl-1,6-hexanediamine (TMHMDA) were obtained from Sigma-Aldrich (St. Louis, MO, USA). 4-bromo-1-butene and lithium bis(trifluoromethane)sulfonimide were purchased from ABCR (Karlsruhe, Germany). 4-methyl-hexahydro-phthalic anhydride (MHHPA) and bisphenol A diglycidyl ether (DGEBA) were supplied by Huntsman (The Woodlands, TX, USA). \(N, N\)-dimethylbenzylamine (BDMA) was purchased from Fisher Scientific (Hampton, NH, USA). All chemicals were used without further purification.

2.2. Synthesis and Thermal Curing of Ionic Liquids

1,3-bis(2-oxiranyl-ethyl)imidazolium Tf\(_2\)N\(^-\) (IL-E) was synthesized according to a procedure published by McDanel et al. \([46]\). 20.07 g Imidazole (294 mmol) was dissolved in 170 mL acetonitrile and 24.8 g NaHCO\(_3\) (294 mmol) and 83.57 g 4-bromo-1-butene (617 mmol) were added to the solution and refluxed for 15 h. The reaction mixture was allowed to cool to room temperature, subsequently filtered and the solvent was removed under reduced pressure. 175 mL of de-ionized water were added to the residue, followed by washing with ethyl acetate and heptane. An amount of 88.83 g (309 mmol) of lithium bis(trifluoromethane)sulfonimide was added to the aqueous phase and the solution was stirred for 4 h at room temperature. The aqueous phase was separated, and the remaining oily layer was extracted with 500 mL ethyl acetate. The organic phase was washed with 1 M aqueous HCl solution and saturated NaHCO\(_3\) solution, followed by washes with de-ionized water until the test with AgNO\(_3\) was negative for halides. MgSO\(_4\) and activated charcoal was afterwards added to the organic phase. After stirring for 3 h, the solution was filtered through basic alumina and the solvent was removed under reduced pressure to yield 122 g (90% yield) of 1,3-di(3-butenyl)imidazolium Tf\(_2\)N. The epoxidation was performed batchwise: 45 g (98 mmol) 1,3-di(3-butenyl)imidazolium Tf\(_2\)N was dissolved in 200 mL acetonitrile, 90 g (394 mmol) 3-chloroperoxybenzoic acid was added and the solution was then stirred for 48 h at room temperature. The reaction mixture was filtered, and the solvent was removed under reduced pressure. Residual 3-chlorobenzoic acid and unreacted 3-chloroperoxybenzoic acid was removed by washing with 2-methoxy-2-methylpropan. The product was dried under vacuum to yield 35 g (73%) of IL-E. The chemical structure was confirmed via \(^1\)H- and \(^13\)C-NMR in DMSO-d\(_6\), and FTIR spectroscopy using literature reports \([46,97,98]\) (Figures S1–S3, Supplementary Materials).

IL-E was cured with selected amine and anhydride hardeners in stoichiometric ratios. TMHMDA, IPD, and DDS were used in a molar ratio of 0.5 to the epoxy resin. MHHPA was added in a molar ratio of 1.7 to IL-E, with 3 wt.% BDMA in relation to the epoxy resin as accelerator. DGEBA was cured with DDS in a molar ratio of 1:0.5 as a reference thermoset. For the production of resin containing both IL-E and DGEBA, the resins were mixed in the desired molar ratios prior to the addition of DDS in a stoichiometric ratio. Liquid hardeners were thoroughly mixed into the resin at room temperature. DDS was dissolved in the epoxy monomers by stirring the components at 150 °C until a homogenous mixture was achieved (between 5 and 20 min). The formulations were then homogenized in a vortex mixer and degassed under vacuum before curing. An overview of the chemical structure of the implemented resins and hardeners, their respective abbreviation and cure conditions of the networks is listed in Table 2.
Table 2. Overview of the composition of the prepared epoxy-based resins and their respective cure cycles.

| Abbreviation | Epoxy Resin | Hardener | Cure Cycle |
|--------------|-------------|----------|------------|
| IL-TMHMDA    | IL-E        | TMHMDA   | 60 °C (16 h) + 130 °C (1 h) |
| IL-IPD       | IL-E        | IPD      | 130 °C (16 h) + 160 °C (3 h) |
| IL-MHHPA     | IL-E        | MHHPA    | 80 °C (1 h) + 140 °C (2 h) + 180 °C (30 min) |
| IL-DDS       | IL-E        | DDS      | 160 °C (16 h) + 180 °C (2 h) |
| DGEB-DDS     | DGEBA       | DDS      | 160 °C (16 h) + 180 °C (2 h) |
| 90-10        | 90 mol% IL-E | DDS      | 160 °C (16 h) + 180 °C (2 h) |
| 50-50        | 50 mol% IL-E | DDS      | 160 °C (16 h) + 180 °C (2 h) |

2.3. Characterization Methods

Infrared spectra were recorded on a Vertex 70 spectrometer (Bruker, Billerica, MA, USA) in attenuated total reflectance (ATR) mode with the software OPUS 7.5 (Bruker, Billerica, MA, USA). A total number of 16 scans were gathered from 500 cm⁻¹ to 4500 cm⁻¹ with a resolution of 2 cm⁻¹.

Measurements of the viscosity of the investigated resins were done using an MCR 501 rheometer (Anton Paar, Graz, Austria). The plate-plate measurements were conducted with a shear rate of 10 s⁻¹ and a deformation of 7% at 30 °C.

A dynamic mechanical analysis (DMA) was done on a DMA/SDTA861 instrument (Mettler-Toledo, Columbus, OH, USA) in tensile mode with a clamping length of 9 mm in accordance to DIN EN ISO 6721-4 [99]. Rectangular specimens of 20 mm × 2.5 mm × 1 mm were tested in a temperature range of 50 °C up to a maximum temperature of 250 °C or at least 100 °C above the maximum tan delta with a heating rate of 2 K/min under nitrogen. All measurements were performed in the strain-controlled mode with 3 μm deformation, 1 Hz frequency and 150 % off-set.

The thermal analysis equipment DSC 4000 (PerkinElmer, Waltham, MA, USA) was used for differential scanning calorimetry (DSC) measurements according to DIN EN ISO 11357-2 [100]. The weight of the samples in the standard aluminium pans (40 μL) was in the range of 12 mg to 14 mg for cured samples and 4 mg to 6 mg for uncured samples. All samples were first heated from −50 °C to 200 °C or 300 °C, for resins containing DDS, with 20 K/min under nitrogen atmosphere. Afterwards, each sample was cooled
down to −50 °C. In the subsequent second run, all samples were heated up again to the above-mentioned temperatures using the same heating rate and atmosphere. The glass transition temperature was determined as midpoint temperature in accordance to the relevant standard using the software Pyris (PerkinElmer, Waltham, MA, USA).

$^1$H-NMR and $^{13}$C-NMR spectra were recorded on a Bruker (Billerica, MA, USA) Advance III 300 MHz NMR spectrometer in DMSO-$d_6$.

Thermogravimetric analysis (TGA) was performed on a TGA/DSC thermogravimetric analyser (Mettler-Toledo, Columbus, OH, USA) from 25 °C to 900 °C with a heating rate of 10 K/min under nitrogen in aluminium oxide crucibles with a volume of 70 µl and an approximate sample weight of 30 mg. The measured curves were evaluated with the STAR® Evaluation Software (Mettler-Toledo, Columbus, OH, USA).

Density was measured in distilled water using a 50 mL pycnometer with a ground-in thermometer at room temperature. The density of the samples $\rho_s$ was calculated from the sample weight $m_s$, the weight of the pycnometer filled with water $m_w$, the weight of the pycnometer filled with water and the sample $m_{sw}$, and the density of water at the respective temperature $\rho_w$ using the following equation:

$$\rho_s = \frac{m_s}{m_s + m_w - m_{sw}} \ast \rho_w$$

3. Results and Discussion

The synthesis of the ionic liquid 1,3-bis(2-oxiranyl-ethyl)imidazolium trifluoromethane-sulfonimide (IL-E) via alkenylation followed by ion exchange and epoxidation [46] proved to be a very robust and efficient procedure for the production of epoxy-ILs with high purity, as also indicated by the obtained NMR spectra and the nearly colorless appearance of the IL [101]. The monomer was chosen for this study due to its chemical structure with comparably short side chains. This allows for more versatility, since the network properties can be effectively tailored and repeating units between the ionic moieties can be easily varied by the hardener. IL-E exhibits a viscosity of 150 mPa s at 30 °C and was stable during storage for several months. In comparison, a significantly higher viscosity of 860 mPa s was measured for the commonly used epoxy resin DGEBA.

3.1. Thermal Properties of Thermally Cured IL-E

The epoxy-IL resins were thermally cured with aliphatic, cycloaliphatic and aromatic amine hardeners to achieve cross-linked networks with different chemical and physical properties. Their curing behavior was monitored via DSC and complete curing was characterized during the second heating run. Due to the higher reactivity of the systems including aliphatic and cycloaliphatic amines (IL-TMHMDA and IL-IPD), curing already started at room temperature after mixing the materials and the compositions could therefore not be stored for extended periods of time. The systems with the aromatic DDS hardener (IL-DDS) presented a higher latency. IL-DDS was stored in a fridge for several weeks and no pronounced phase separation or curing reaction was observed, which is in good agreement with the longer pot life and gel time of aromatic hardeners compared to aliphatic hardeners reported in literature [102,103]. The anhydride system (IL-MHHPA) also showed a starting curing reaction but can nonetheless be stored over extended periods (at least one week) of time if no accelerator is added. Typical DSC scans for the various epoxy-IL resins are depicted in Figure 2. Table 3 summarizes the characteristic temperatures of the uncured resin-hardener mixtures. The glass transition temperature, midpoint temperature in the second heating run of DSC tests, is denoted as $T_{g,u}$. The results confirm the initial observation of the latency of IL-DDS having significantly higher temperatures of the exothermic peak ($T_{\text{peak}}$) of 283 °C and curing onset ($T_{\text{onset}}$) of 171 °C than the other epoxy-IL systems. A slightly smaller exothermic peak was observed at 230 °C. In consideration of the relatively low $T_{g,u}$ of 65 °C for IL-DDS, the secondary exotherm was assigned to epoxy-IL homopolymerisation reactions. When comparing IL-DDS with DGEBA-DDS, it can be seen that $T_{\text{onset}}$ of 185 °C and $T_{\text{peak}}$ of 238 °C for DGEBA-DDS correspond very well
with $T_{\text{onset}}$ and the first exotherm of IL-DDS. This further indicates that the higher $T_{\text{peak}}$ is derived from etherification [104]. It has to be noted that at such high temperatures some thermal degradation is likely to occur and a slight exotherm is visible when heating the pure IL-E to 300 °C in the DSC, however, no peak was visible. Epoxy-anhydride systems generally require a catalyst such as tertiary amines to initiate the alternating ring-opening polymerization and the onset temperature can be influenced by the variation of the catalyst concentration [105]. With 3 wt.% BDMA as catalyst, a curing onset of 120 °C and $T_{\text{peak}}$ of 161 °C was measured for IL-MHHPA. Attempts to reduce the curing temperature for IL-DDS by addition of 1 wt.% BDMA resulted in a significant reduction of $T_{\text{onset}}$ and $T_{\text{peak}}$ to 88 °C and 135 °C, respectively. However, using the accelerator also resulted in a drastically reduced $T_{g,u}$ value of 38 °C, indicating that BDMA mainly initiated the homopolymerisation of the IL, and was therefore disregarded. This observation is in good agreement with reports on catalyzed DGEBA-DDS systems [106].

Figure 2. DSC thermograms of the uncured resin-hardener mixtures, including the first (top) and second (bottom) heating. Curves shifted vertically for better visibility.

Table 3. Characteristic temperatures obtained from DSC measurements performed on the uncured systems: $T_{\text{onset}}$ and $T_{\text{peak}}$ of the curing reaction (first heating), and glass transition temperature ($T_{g,u}$) (second heating).

| Resin/Hardener | $T_{\text{onset}}, \degree C$ | $T_{\text{peak}}, \degree C$ | $T_{g,u}, \degree C$ |
|----------------|-------------------------------|-----------------------------|------------------|
| IL-TMHMDA      | 58                            | 109                         | 32               |
| IL-IPD         | 63                            | 108 & 161                   | 61               |
| IL-DDS         | 171                           | 230 & 283                   | 65               |
| IL-MHHPA       | 120                           | 161                         | 99               |
| DGEBA-DDS      | 177                           | 241                         | 200              |

IL-IPD also exhibited a double exotherm at 108 °C and 161 °C. In this case, the double exotherm can be attributed to the different reactivity of the aliphatic and cycloaliphatic amine groups of the non-symmetrical IPD hardener [107]. This is further evidenced when comparing IL-IPD with IL-MHHPA and IL-TMHMDA, cured with cycloaliphatic and aliphatic hardeners, respectively. The higher $T_{\text{peak}}$ is at the same temperature as for IL-MHHPA. On the other hand, the $T_{\text{onset}}$ of 63 °C and the lower $T_{\text{peak}}$ at 108 °C are very similar to IL-TMHMDA, with a $T_{\text{onset}}$ of 58 °C and a $T_{\text{peak}}$ at 109 °C. The use of the aliphatic hardener in IL-TMHMDA also resulted in the lowest $T_{g,u}$ of 32 °C. Apart from DGEBA-DDS with 200 °C, the anhydride system IL-MHHPA exhibited the highest $T_{g,u}$ of 99 °C, being even higher than the epoxy-IL system including the DDS hardener with its
high aromatic content. IL-IPD with its mixed structure exhibits a $T_{g,u}$ of 63 °C, ranking it between IL-TMHMDA and IL-MHHPA.

DSC Measurements of Cured Samples

For a further investigation of the thermal properties, additional samples were prepared by resin casting and subsequent curing in a convection oven according to the determined temperature profiles for full curing. The curing temperatures were chosen in consideration of the respective $T_{\text{onset}}$ and $T_{\text{peak}}$ (see Table 3). Although the uncured resin samples did not exhibit an exothermic peak after the first heating run in DSC tests, a prolonged curing (post-curing) in the oven allowed to reach a higher glass transition temperature ($T_{g,c}$) for the amine-cured systems. After reaching the gel-point, the curing reactions changes from kinetics-controlled to diffusion-controlled [108]. The additional curing time allows the diffusion-controlled curing process to proceed to a higher conversion, leading to a higher cross-link density and thus, higher $T_g$. Figure 3 includes a comparison of the second heating curve from DSC tests with the compositions after post-curing to visualize the shift from $T_{g,u}$ to $T_{g,c}$. After post-curing, the glass transition temperature ($T_{g,c}$) of IL-TMHMDA was increased by 8 °C to 39 °C and by 4 °C to 65 °C for IL-IPD, respectively. The discrepancy between $T_{g,u}$ and $T_{g,c}$ is especially pronounced for IL-DDS and a $T_{g,c}$ of 97 °C compared to the $T_{g,u}$ of 65 °C was reached. Interestingly, the IL-DDS system still exhibits a $T_g$ of more than 100 °C lower compared to the respective DGEBA-DDS system with a $T_{g,c}$ of 220 °C. Compared to literature values of DGEBA cured with IPD with up to 155 °C [109], a similar relation can be observed for the $T_{g,c}$ of IL-IPD at 64 °C. The anhydride cured system IL-MHHPA reached nearly 70% of reported values for DGEBA cured with MHHPA [110]. It is expected that this originates from the different molar and consequently weight ratios of the anhydride system. While the molar ratio of IL-E to DDS or IPD is 2:1, it is 1:1.7 for IL-MHHPA. Therefore, the final network properties are more significantly determined by the hardener structure for the anhydride system. Concluding from these results, the characteristic temperatures ($T_{g,c}$, $T_{\text{peak}}$ and $T_{g,u}$) decrease in the order aromatic > cycloaliphatic > aliphatic for the amine curing agents, while the anhydride hardener provides a $T_g$ comparable to the aromatic amine hardener (see Figure 3), which follows the same trend as conventional epoxy systems [110,111].

![Figure 3](image_url)

Figure 3. Shift of glass transition temperatures by prolonged post-curing (thick lines) compared to the second heating of uncured resins (thin lines) as determined in DSC. Curves shifted vertically for better visibility.
TGA measurements (Figure 4) were done on fully cured epoxy-IL resins and DGE-DDS systems in order to determine the thermal stability of the cured systems. The results are in good agreement with the high thermal stability of imidazolium based ionic liquids reported in literature [112]. The highest degradation temperature at 5% mass loss ($T_{\text{dec,5}}$) was determined for DGE-DDS at 385 °C. This high stability is explained by the highly aromatic content of the resin-hardener system. For the epoxy-IL samples, a two-step degradation is visible. The initial mass loss can be assigned to the degradation of the aliphatic moieties, initiated by the cleavage of the N-C bonds to the quaternized imidazolium nitrogen atoms. This is underlined by the significantly lower initial weight loss for the pristine, uncured IL-E and by the degradation of DGE-DDS. The aromatic hardener DDS slightly increases the thermal stability of the cured IL-DDS network to a $T_{\text{dec,5}}$ of 334 °C, above the stability of the pristine IL-E ($T_{\text{dec,5}} = 321$ °C). The cycloaliphatic hardener IPD, however, does not improve the thermal stability and is comparable to the linear aliphatic hardener TMHMDA, with $T_{\text{dec,5}}$ of 300 °C and 301 °C for IL-IPD and IL-TMHMDA, respectively. A $T_{\text{dec,5}}$ of 338 °C was measured for IL-MHHPA, indicating that the aliphatic amine of IPD might be the determining factor for the thermal stability.

![Figure 4. TGA curves obtained for IL-E and fully cured epoxy/hardener compositions.](image)

### 3.2. Thermomechanical Properties of Thermally Cured IL-E

Dynamic mechanical analysis (DMA) was performed on cured specimens prepared by resin casting in order to characterize the temperature dependence of the mechanical properties and to estimate the stiffness of the materials. Storage modulus ($E'$) and glass transition temperature values as determined from these tests are shown in Table 4.

| Material        | $T_{g'}$ °C | $E'$, GPa$^{*}$ | $E_{r'}$, Mpa |
|-----------------|------------|----------------|--------------|
|                 | Onset      | Max tan δ      | $T = 0$ °C   | 5%             |
| IL-TMHMDA       | 18         | 28             | 1.3          | 7.8            |
| IL-IPD          | 44         | 73             | 2.2          | 7.0            |
| IL-MHHPA        | 91         | 107            | 1.9          | 10.0           |
| IL-DDS          | 99         | 113            | 2.1          | 9.8            |
| DGE-DDS         | 196        | 220            | 2.2          | 33             |

* $E_{r'}$: Storage modulus in rubbery state ($T = T'(\text{max tan δ}) + 30$ °C).
Curing of IL-TMHMDA led to comparably soft networks with a \( T_g \) of 28 °C and a storage modulus (\( E' \)) of 1.3 GPa. This behavior accounts for the lack of aromatic moieties between imidazolium groups, allowing for a higher chain mobility and decreasing the stiffness of the polymer backbone [113]. McDanel et al. [46] reported a similar \( T_g \) of 38 °C for the IL-E monomer cured with a triamine. However, the authors calculated a conversion of only 67%. An excess amount of hardener increased the conversion to 80% but at the same time reduced the \( T_g \) to 9 °C due to the introduction of further aliphatic groups.

The implementation of the cycloaliphatic hardener in the IL-IPD resin system led to an increase of the \( T_g \) to 73 °C. Due to its high aromatic character, the IL-DDS system shows a further increase of the \( T_g \) to 113 °C. As discussed in the previous section, the comparably high \( T_g \) of 107 °C for the cured resin including the cycloaliphatic MHHPA hardener likely derives from the higher ratio of hardener to epoxy-IL resin. However, when compared to the DGEB-DDS system with a \( T_g \) of 220 °C, the glass transition of the epoxy-IL networks is still located at significantly lower temperatures. Apart from the different chemical structure of the heterocyclic imidazole, the increased volume due to the bulky anions implemented in the epoxy-IL networks and their plasticizing effect is expected to be the major influence on the decreased \( T_g \). While the \( \text{TF}_{2} \text{N}^- \) anion provides many desirable properties e.g., in terms of viscosity of the monomer and thermal stability, if further increase of the \( T_g \) is required, an anion exchange to reduce the free volume might be reasonable.

The non-symmetrical hardener IPD leads to a cured epoxy-IL network with a slightly broader glass transition and nearly 30 °C difference between the onset temperature and the peak of \( \tan \delta \). A secondary relaxation temperature, visible as a shoulder in the \( \tan \delta \) curve (Figure 5), is observed for IL-IPD. This is accounted for by the chemical structure of IPD and the influence of the cycloaliphatic structure.

**Figure 5.** Storage modulus (solid lines) and \( \tan \delta \) (dashed lines) of the PIL networks and DGEB-DDS.
IL-IPD and IL-DDS exhibit storage moduli of 2.2 GPa and 2.1 GPa at 0 °C, respectively, similar to DGEB-DDS with 2.2 GPa, due to the rigid cyclic groups introduced by the hardeners. Interestingly, the anhydride cured network shows a lower E’ of 1.9 GPa at 0 °C, despite its high T_g. At 31 °C, E’ of IL-MHHPA intersects with that of IL-DDS. However, E’ of IL-MHHPA increases less significantly with decreasing temperature. At −48 °C E’ amounts to 2.7 GPa and 2.6 GPa for IL-DDS and IL-IPD, respectively, while the modulus of IL-MHHPA only increased to 2.2 GPa. This indicates that the properties of the anhydride cured systems are less temperature dependent at low temperatures than the amine system, which might be advantageous for applications with large temperature differences.

Above the glass transition region, the epoxy-IL networks exhibit similar storage moduli in the rubbery state (E_r) of 7.0 MPa, 7.8 MPa, 9.8 MPa, and 10.0 MPa, following the order IL-IPD < IL-TMHMDA < IL-DDS < IL-MHHPA, while a E_r of 33 MPa was measured for DGEB-DDS. This indicates that the IL-E determines the E_r of the epoxy-IL networks and the effect of the hardener is less pronounced.

Further networks were prepared by adding DGEBA as co-monomer and compositions with molar ratios of 1:0, 1:1, 9:1, and 0:1 of IL-E/DGEBA were produced and cured with DDS. The influence of the DGEBA content on the thermo-mechanical properties of the cured PIL networks was studied. The aim of these experiments was to determine the influence of the DGEBA content on the thermo-mechanical properties of the resulting networks, since conventional DGEBA resins have been extensively investigated in the past and DGEB-DDS networks provided good thermal stability and thermo-mechanical behavior in our experiments. Such blends of functionalized ionic liquids with other epoxy-based resins, which are not only limited to DGEBA, enable a high flexibility in tailoring the resulting network properties. In addition to the thermo-mechanical properties of the resulting epoxy-IL networks, also the ionic character and ion conductivity can therefore be flexibly tailored. Figure 6 depicts the DMA curves (a) as well as the glass transition temperatures and densities (b) of the networks in dependence of the molar fractions of epoxy monomers. Both the glass transition temperature and the density are observed to change linearly with the mol fractions. With increasing amounts of the IL monomer, the glass transition temperature of the thermoset decreases from 220 °C to 169 °C, 119 °C and 113 °C, while the density increases from 1.24 g/cm^3 to 1.41 g/cm^3, 1.58 g/cm^3 and 1.63 g/cm^3 for 50 mol%, 90 mol% and 100 mol% IL-E, respectively. This is in accordance with findings of other reports where imidazolium Tf_2N derivatives were used as co-monomers [94,95]. The large Tf_2N⁻ anion is reported to act as a plasticizer and leads to polymers with a low T_g [114]. The curing of DGEBA and epoxy-IL successfully yielded a homogenous network, as there are no additional relaxation temperatures visible and the glass transition range does not broaden. Therefore, while decreasing the ionic character of the material, the addition of DGEBA as co-monomer seems to be a suitable method to produce PILs for high thermomechanical demands. Compared to the copolyimides produced by Li et al. [73], the T_g is still significantly lower. However, the system is promising for lower T_g applications as the epoxy monomer is easier to process and does not require solvents, while higher IL loadings are achievable.
The present work, the versatility of epoxy functionalized ionic liquids to achieve cross-linked poly(ionic liquids) with tunable properties was demonstrated. An epoxy-IL network was synthesized and the thermal and thermo-mechanical properties as a function of the applied hardener were evaluated. Due to the vast commercial use of conventional epoxy systems and the reactivity of the oxirane rings towards a multitude of species, cross-linked IL-thermosets can be produced easily, and the properties can be tailored by using commercially available hardeners. This was illustrated by adjusting the glass transition temperature of epoxy-IL networks from 30 °C to 100 °C using four different hardeners. The $T_g$ was shifted to even higher temperatures, by adding DGEB to the formulations. All reported networks only required thorough mixing and thermal curing, without the need of any solvent, for the preparation of free-standing specimens. This demonstrates the
potential of epoxy-IL monomers for multiple technical applications and the possibilities of effectively tailoring the resulting resin properties by a suitable selection of hardener and co-monomer. Further evaluation of suitable alternatives to the implemented Tf$_2$N$^-$ anion opens up possibilities for cost effective epoxy-IL based monomers and possibly higher resulting $T_g$ values, including the beneficial properties of ionic liquid monomers for certain applications.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/polym13223914/s1, Figure S1: $^1$H-NMR spectrum of IL-E, Figure S2: 13C-NMR spectrum of IL-E, Figure S3: FTIR-spectrum of IL-E with corresponding characteristic frequency regions. References [46,97,98].

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