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## I. Introduction

Many biotechnological applications rely on the retarded release of active ingredients that are embedded in amorphous solid dispersions or on the liberation of drugs and pesticides from polymeric carriers.<sup>1–3</sup> These phenomena require a detailed characterization of the underlying diffusion processes that take place in the amorphous matrices. Numerous theoretical approaches have been developed in the context of plasticizer transport, yet their predictive power leaves room for improvements. This is understandable, for instance in view of the longknown and still debated mutual decoupling of various relaxation processes.<sup>4–14</sup> Decoupling phenomena are not at all confined to the field of polymers, but observed in a diverse array of condensed matter systems.<sup>15–19</sup> In fact, for instance in the field of polymer electrolytes and other ion conducting systems

# Relaxation and diffusion of an ionic plasticizer in amorphous poly(vinylpyrrolidone)<sup>+</sup>

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The present work focuses on the dynamics of the ionic constituents of 1-propyl-3-methyl-imidazoliumbis-(trifluormethylsulfonyl)-imide (PT), a paradigmatic ionic liquid, as an additive in poly(vinylpyrrolidone) (PVP). Hence, the resulting product can be regarded as a polymer electrolyte as well as an amorphous dispersion. Leveraging dielectric spectroscopy and oscillatory shear rheology, complemented by differential scanning calorimetry, the spectral shapes and the relaxation maps of the supercooled PVP-PT mixtures are accessed in their full compositional range. The study also presents dielectric and shear responses of neat PVP with a molecular weight of 2500 g mol<sup>-1</sup>. We discuss the plasticizing role of the PT additive and the decoupling between ionic dynamics and segmental relaxation in these mixtures. The extracted relaxation times, steady-state viscosities, and conductivities are employed to estimate the translational diffusivities of the ionic penetrants by means of the Stokes–Einstein, Nernst–Einstein, and Almond–West relations. While some of the estimated diffusivities agree with each other, some do not, pointing to the importance of the chosen hydrodynamic approximations and the type of response considered for the analysis. The present extensive dielectric, rheological, and calorimetric study enables a deeper understanding of relaxation and transport of ionic ingredients in polymers, particularly in the slow-dynamics regime which is difficult to access experimentally by direct-diffusivity probes.

decoupling of charge and structure dynamics is highly desired.  $^{\rm 20\text{-}23}$ 

Therefore, further experimental studies of small-molecule diffusion in polymers are warranted. Consequently, a wide variety of techniques have been devised for this purpose. Many of them employ light as a probe such as in studies using forced Rayleigh scattering,<sup>24,25</sup> fluorescence recovery after photobleaching,<sup>26–29</sup> fluorescence nonradiative energy transfer,<sup>30</sup> and fluorescence correlation microscopy.<sup>31</sup> Among various other approaches, nuclear magnetic resonance was also exploited.<sup>32</sup> Furthermore, dielectric and viscosimetric techniques were explored to assess diffusion in neat supercooled liquids.<sup>33,34</sup>

In the present work, these latter approaches are exploited for amorphous solid dispersions based on polyvinylpyrrolidone (PVP). This biocompatible polymer plays a prominent role. *e.g.*, in the field of pharmaceutical formulations. PVP was studied in pure and hydrated forms for various molecular weights.<sup>35-46</sup> PVP-based amorphous solid dispersions featuring pharmaceuticals such as nifedipine,<sup>47,48</sup> indomethacin,<sup>49-51</sup> and others were also investigated.<sup>52</sup>

Another area of research, which is particularly relevant for energy storage technologies, is the transport of penetrants that act as electric charge carriers in polymer matrices.<sup>23,53–55</sup> The workhorse material here is polyethylene oxide,<sup>56</sup> a semicrystalline polar polymer doped with various salts in a rather reduced

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<sup>&</sup>lt;sup>†</sup> Electronic supplementary information (ESI) available: As supporting information we present loss modulus spectra for  $PVP_{0.34}PT_{0.16}$ , *G'*, *e'*, and *e''*<sub>der</sub> spectra for  $PVP_{1-x}PT_x$  with x = 0.16. 0.33, and 0.66, as well as details regarding the DSC measurements and how they were analyzed. Furthermore, the data in Fig. 7 are presented in a Walden-type plot. See DOI: https://doi.org/10.1039/d4cp01001a

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concentration range.<sup>57</sup> While the resulting polymer electrolytes mitigate several drawbacks of inorganic conductors by providing mechanical flexibility and good contact with the electrodes, they often suffer from reduced electrochemical stability.<sup>58</sup> For example, the degradation of polyethylene oxide above 4 V precludes its use in combination with high-voltage electrodes for high performance batteries.<sup>59</sup> This situation warrants the current search for alternative polar polymers with a reduced degree of crystallization propensity, such as PVP, as ion matrices for energy storage applications.

In the present work, we first study neat PVP with a molecular weight near 2500 g mol<sup>-1</sup> (on average featuring 23 monomer units per chain) and then mixtures of an ionic liquid with PVP. While many applications have emerged for mixtures of polymers and ionic liquids,<sup>60,61</sup> here we employ them to track the ion diffusion *via* dielectric spectroscopy. Thus, we exploit that mass and charge transport are intimately coupled<sup>62,63</sup> in ionic liquids such as 1-propyl-3-methyl-imidazolium-bis(trifluormethyl-sulfonyl)-imide (for brevity here denoted PT, previously also abbreviated as PMIM-TFSI).

Many active ingredients are however not intrinsically charged, but often display a residual conductivity only, mostly originating from minute ionic contaminants left over from their production. This calls for tests to find out whether charge transport and the diffusion-related mass transport are coupled or not. Interestingly, for several viscous liquids it has been demonstrated how rheological timescales can be used as a suitable predictor of diffusion phenomena over a temperature range down to the calorimetric glass transition at  $T = T_{gr}^{34}$ 

Therefore, in the present article, we generalize the approach from that work and test it for a polymer electrolyte. Thus, we examine the extent to which rheology in conjunction with conductometry can be exploited to track the diffusion of PT in the PVP matrix. More specifically, based on dielectrically detected DC conductivities, we use the Nernst–Einstein relation to estimate the plasticizer's diffusion coefficients and compare the results with those from a rheology based determination using the Stokes–Einstein equation. Apart from these classical approaches relying on steady-state responses we also employed the original<sup>64</sup> and the recently introduced modified Almond–West concepts. Here, the elementary diffusivity length of the plasticizer is assessed based on the material's frequency-dependent conductivity<sup>65</sup> as well as on its fluidity.<sup>34</sup>

While the dielectric and rheological properties of  $PT^{66}$  have been examined already, it appears that neat PVP with a molecular weight of 2500 g mol<sup>-1</sup> was not studied previously in these respects. Therefore, after providing the experimental details, we present and discuss our results regarding the dynamics of neat PVP. Then, based on measurements and analyses of the dielectric and shear mechanical spectra for the PVP-PT mixtures, we compare their spectral widths, glass transition temperatures, viscosities, and electrical conductivities. Finally, based on the relaxation times for all systems, we compute the related diffusivities and discuss the relative merits of the various approaches used to this end.

## II. Experimental details

For the present experiments we use PT (nominal purity >98%, Sigma Aldrich) and PVP (also termed PVP K-12 = Kollidon 12PF from BASF). The latter is specified by the supplier with a weight average molecular weight of 2000–3000 g mol<sup>-1</sup>.

Prior to the measurements, pure PVP was dried overnight at 80  $^{\circ}$ C and melted at 180  $^{\circ}$ C directly on the lower plate of the dielectric or the rheology cell. Then, for the dielectric measurements the (room-temperature) upper electrode was put onto the sample, while for the rheological measurements the upper plate was priorly heated to 180  $^{\circ}$ C.

To prepare the ionic liquid mixtures, the appropriately weighed components PVP powder and liquid PT were mixed, stirred, and dried at 100 °C until a clear (slightly yellowish) liquid was formed. The premixed PVP powders were melted at 160 °C because at higher temperatures, the samples will quickly develop a brownish color. As soon as the mixtures have become liquid they are kept under vacuum at a temperature of 80 °C. For the ionic-liquid mixtures we studied molar PT fractions (with respect to the PVP monomers) of *x* = 0.10, 0.16, 0.33, and 0.66, corresponding to weight fractions of *w* = 28.6, 41.2, 64.0, and 87.8 wt%, respectively.

The dielectric response was probed using an Alpha-A impedance analyzer from Novocontrol and thermostatted using a Quatro unit. The samples were investigated in a parallel-plate cell made of invar and sapphire.<sup>67</sup> For the shear mechanical oscillation or rotation experiments, an MCR 502 rheometer from Anton-Paar was employed. The gap between the two shear plates was 1 mm.

Differential scanning calorimetry (DSC) was conducted with a Q2000 from TA Instruments. The onset glass transitions were determined for a heating rate of 10 K min<sup>-1</sup>. For the mixtures, the glass transition temperatures range from that of pure PVP ( $T_g = 375 \text{ K}$ ;<sup>68</sup> this work:  $T_g = 373 \text{ K}$ ) to that of neat PT ( $T_g = 180 \text{ K}$ ).<sup>66</sup> Further experimental details as well as the DSC thermograms are given as ESI.†

### III. Results and discussion

#### A. Dynamics of neat PVP

The segmental relaxation of PVP was probed using dielectric spectroscopy and Fig. 1 shows the results obtained for the dielectric function,  $\varepsilon^* = \varepsilon' - i\varepsilon''$ . In panel (a) the typical relaxation step is observed in  $\varepsilon'$ . As the temperature is lowered, it moves through the frequency window, signaling the slow-down of the segmental motion. Towards lower frequencies an upturn in  $\varepsilon'$  is seen. This is the hallmark of blocking electrode effects which in this polymer arise as a consequence of a residual ionic conduction. The conductivity itself is seen more directly as a low-frequency feature in  $\varepsilon''$  which, in particular at low temperatures, see Fig. 1(b), masks the dielectric loss peak completely. For sufficiently broad spectra, one can nevertheless render the underlying process visible by using the logarithmic derivative<sup>69</sup>

$$\varepsilon''(\omega) \approx -\frac{\pi d\varepsilon'(\omega)}{2d\ln(\omega)} = \varepsilon''_{der}$$
 (1)



**Fig. 1** (a) Real and (b) imaginary part of the dielectric function of PVP K-12. The lines in panel (a) and (b) are fits using eqn (2), those in (a) are augmented by the electrode polarization term described in the text. Panel (c) presents  $e'_{der}$ , *cf.* eqn (1), and the lines are fits using eqn (2).

adapted from rheology,<sup>70,71</sup> where apparently it was first introduced. For the whole accessible temperature range the emergence of  $\varepsilon''_{der}$  peaks is demonstrated in Fig. 1(c).

For a more quantitative analysis we employ the empirical Havriliak–Negami (HN) function<sup>72</sup>

$$\varepsilon^*(\nu) = \varepsilon_{\infty} + \frac{\Delta\varepsilon}{[1 + (i2\pi\nu\tau_{\mathrm{HN},\varepsilon})^{\alpha_{\varepsilon}}]^{\gamma_{\varepsilon}}} - i\frac{\sigma_0}{2\pi\nu\varepsilon_0}$$
(2)

Here,  $\varepsilon_0$  denotes the vacuum permittivity and the dielectric loss part is augmented by a term relating to the electrical conductivity  $\sigma_0$ . In eqn (2)  $\tau_{HN,\varepsilon}$  denotes the characteristic time and  $\alpha_{\varepsilon}$ and  $\gamma_{\varepsilon}$  account for the broadening and the asymmetry, respectively, of the loss peaks. From the PVP data shown in Fig. 1 we find  $\alpha_{\varepsilon}$  = 0.73 ± 0.04 and  $\gamma_{\varepsilon}$  = 0.35 ± 0.05, so that on the highfrequency flank the dielectric loss follows a power-law  $\varepsilon'' \propto$  $\nu^{-\gamma_{\epsilon,\text{eff}}}$  with an effective exponent  $\gamma_{\epsilon,\text{eff}} = \alpha_{\epsilon}\gamma_{\epsilon}$  which here is 0.26. For the fits shown in Fig. 1(a) the blocking-electrode effects were taken into account in terms of a power-law contribution, justified in detail in ref. 73 and 74. In effect, this phenomenon leads to a low-frequency power-law,  $\varepsilon' \propto \nu^{-s}$ , with an exponent s, typically in the range between about 1 and 2; here we find s = 1.5. For PVP and subsequent dielectric data, this power law contribution is shown for the  $\varepsilon'$  fits, but not for  $\varepsilon''_{der}$ , in order to gain a better impression of the fitted spectral peak shape.



Fig. 2 (a) Real and (b) imaginary part of the shear modulus of PVP K-12. The lines are fits using eqn (4) and the numbers indicate temperatures in Kelvin.

The dielectric relaxation time  $\tau_{\varepsilon}$  corresponding to the position of maximum loss can be assessed from<sup>75</sup>

$$\tau_{\varepsilon} = \tau_{\rm HN} \left[ \frac{\sin(\alpha_{\varepsilon} \gamma_{\varepsilon} \pi / (2 + 2\gamma_{\varepsilon}))}{\sin(\alpha_{\varepsilon} \pi / (2 + 2\gamma_{\varepsilon}))} \right]^{1/\alpha_{\varepsilon}}$$
(3)

The relaxation times resulting for neat PVP will be summarized together with those from the mixtures, in Section III.C. This comparison will also involve results from DSC (see the ESI†) and the rheological experiments which we present in Fig. 2. There, we show the real and the imaginary part of the complex shear mechanical modulus  $G^* = G' + iG''$  of neat PVP. The present data cover dynamics in the range of the segmental relaxation. In Fig. 2(a) one recognizes that toward high frequencies G' approaches a plateau modulus,  $G_{\infty}$ . Furthermore, in Fig. 2(b) well resolved shear loss peaks are detected over a wide range of temperatures.

To describe the rheological data we employed a (suitably adapted) HN function

$$G^{*}(\nu) = G_{\infty} \left\{ 1 - \frac{1}{[1 + (i2\pi\nu\tau_{\mathrm{HN,G}})^{\alpha_{\mathrm{G}}}]^{\gamma_{\mathrm{G}}}} \right\}$$
(4)

Analogous to eqn (2), the exponents appearing in eqn (4) describe the shape of the rheological spectra. From the fits shown as solid lines in Fig. 2 we find  $\alpha_G = 0.68 \pm 0.05$  and  $\gamma_G = 0.25 \pm 0.03$ . Furthermore,  $\tau_{HN,G}$  denotes the characteristic shear relaxation time, and a relation analogous to eqn (3) holds for the rheological time constants  $\tau_G$ .

# B. Dielectric and shear mechanical spectra for PVP mixed with PT

As an example for a mixture of PVP with PT, in Fig. 3 we present dielectric data for  $PVP_{0.9}PT_{0.1}$ . The permittivity spectra, shown

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in panel (a), resemble those of the neat polymer. However, due to the plasticizing effect of PT, in the binary melt the dynamics is faster so that their spectra were measured at lower temperatures. Furthermore, since the admixture of PT introduces numerous charge carriers into the system, for  $PVP_{0.9}PT_{0.1}$  the relative contribution of the electrical conductivity is larger than that of neat PVP. Thus, even at high temperatures the dielectric loss peaks of  $PVP_{0.9}PT_{0.1}$  are fully swamped. Therefore, in Fig. 3(b), we present  $\varepsilon''_{der}$  instead of  $\varepsilon''$ . But even then, the  $\varepsilon''_{der}$ . spectra display a significant broadening in the peak region, which may indicate a dynamical disparity between the two kinds of ions or between their translational and reorientational (since the present ions carry permanent dipole moments)<sup>76</sup> degrees of freedom.

By contrast to the  $\varepsilon''_{der}$  spectra of neat PVP, for PVP<sub>0.9</sub>PT<sub>0.1</sub> a low-frequency flank cannot be discerned. Hence, for a quantitative description of the data in terms of eqn (2), the exponent  $\alpha_{\varepsilon}$  was fixed. Here, we have chosen a value of 0.5 which provides a good description in the peak region of the  $\varepsilon''_{der}$  spectra. Fig. 3(a) and (b) show that this approach provides a satisfactory fit to the spectra for frequencies near the peak and higher. Similarly broadened  $\varepsilon''_{der}$  peaks, typically devoid of a clearly discernible maximum, are observed for all PVP<sub>1-x</sub>PT<sub>x</sub> samples. The corresponding  $\varepsilon'$ ,  $\varepsilon''$ , and  $\varepsilon''_{der}$  spectra are summarized as ESI.† As will be discussed in detail in Section III.C, for all mixtures the spectral widths (more precisely the high-frequency flanks of their dielectric spectra) are similarly broad.



**Fig. 3** Dielectric data of  $\text{PVP}_{0.9}\text{PT}_{0.1}$  presented in terms of (a)  $\varepsilon'$ , (b)  $\varepsilon''_{\text{der}'}$  and (c)  $\sigma'(\nu)$ . The lines are fits as described in Fig. 1. For this mixture the fit parameters, *cf.* eqn (2), are  $\gamma_{e,\text{eff}} = \alpha_{e}\gamma_{e} = 0.42 \pm 0.05$  and s = 1.5.

Returning to  $PVP_{0.9}PT_{0.1}$ , a close inspection of Fig. 3(a) reveals that upon cooling the relaxation strength slightly decreases and consequently also the amplitude of the  $\varepsilon''_{der}$  spectra, see Fig. 3(b). This anti-Curie-type temperature evolution of the dielectric strength is much more pronounced for the samples with x = 0.33 and 0.66, see the ESI.† This anti-Curie effect is known from the highly viscous regime of glassforming electrolytes,<sup>66,77–79</sup> where the dielectric absorption is largely governed by the conductivity relaxation of the ionic species. This observation also rationalizes why this effect is particularly pronounced in more highly PT doped PVP samples.

The frequency dependent conductivity is directly related to the dielectric permittivity  $\nu ia \ \sigma^*(\nu) = 2\pi\nu\varepsilon_0\varepsilon^*(\nu)$ . Fig. 3(c) displays the real part of the complex conductivity. From the lowfrequency ( $\nu \rightarrow 0$ ) plateau of these  $\sigma'(\nu)$  curves one can extract the DC conductivity  $\sigma_0$  in a straightforward manner. The temperature evolution of  $\sigma_0$ , together with that of  $\tau_{\varepsilon}$ , extracted from the peak of the  $\varepsilon''_{der}$  spectra is discussed in Sections III.C and III.D, respectively.

Shear loss spectra,  $G''(\nu)$ , of  $PVP_{1-x}PT_x$  with PT fractions of x = 0.10, 0.33, and 0.66 are presented in Fig. 4. The corresponding *G'* spectra and the full set of shear data for x = 0.16 are provided as ESI.† Fig. 4 shows that the  $G''(\nu)$  responses for x = 0.1 and x = 0.66 are rather broad. Nevertheless, their shear loss spectra can still satisfactorily be described using eqn (4).

The spectra for x = 0.33 display, however, a much larger width and obviously they develop a two-peak structure toward low temperatures which also in other contexts signaled the existence of a microphase separation.<sup>80</sup> This circumstance renders a reliable determination of time constants cumbersome. Corroborating the conjecture of a microphase separation, we mention that for x = 0.33 a bimodal calorimetric feature is observed (see the ESI†). Near 220 K, the separation of the processes that are comprised in the rheological two-peak structure is on the order of 3 decades. In view of the complexity of the rheological spectra for x = 0.33, we refrained from quantitatively describing their shape and restrict ourselves to determining time constants  $\tau_G$  from the G'' maxima, whenever possible.

Apart from frequency dependent measurements which mostly cover the low-frequency or high-viscosity range of these glassforming binary melts, we also gathered rotational viscosities, extending the probed range down to values of about  $10^2$  Pa s.

# C. Comparison of spectral widths, glass transitions, viscosities, and electrical conductivities

After introducing the data base for the different PVP mixtures and in order to prepare for the analyses regarding the charge and molecular diffusion within them, we first compare the width of their rheological and dielectric spectra, their glass transition temperatures, as well as their DC conductivities and zero-shear viscosities.

To be able to compare the shapes of the different spectra, it is advantageous to rescale them appropriately. Following ref.



**Fig. 4** Shear loss modulus spectra measured for  $PVP_{1-x}PT_x$  with *x* given by (a) 0.1, (b) 0.33, and (c) 0.66. The lines are fits using eqn (4) with  $\alpha_G = 0.54 \pm 0.04$  and  $\gamma_G = 0.38 \pm 0.04$  for x = 0.1 and with  $\alpha_G = 0.55 \pm 0.04$  and  $\gamma_G = 0.56 \pm 0.03$  for x = 0.66. The arrow in panel (b) highlights the approximate position of the broad low-temperature peak that can be identified for 222.5 K.

81, G'' masterplots were constructed with respect to a reference temperature  $T_{ref}$ . For the masterplots presented in Fig. 5(a),  $T_{ref}$ has been chosen close to  $T_g$ , so that the spectra can be considered isochronal with respect to a structural relaxation time of roughly 100 s. The resulting masterplots, further normalized with respect to their peak amplitude  $G''_{max}$  and peak frequency,  $\nu_{max}$ , are shown in Fig. 5(a). Except for PVP<sub>0.67</sub>PT<sub>0.33</sub>, *cf.* Fig. 4(b), for which the construction of a master curve is far from unambiguous, the spectral width of all mixtures are similar to each other and with respect to PVP. Only the G''spectrum of neat PT<sup>66</sup> is significantly narrower than for the



**Fig. 5** Amplitude- and frequency-scaled (a) rheological and (b) dielectric spectra of  $PVP_{1-x}PT_x$ . The plots essentially correspond to isochronal representations with rheological time constants of  $\sim 10^2$  s and dielectric time constants of  $\sim 10^{-2}$  s.

samples containing PVP. Hence, disregarding PVP<sub>0.67</sub>PT<sub>0.33</sub> and slightly simplifying, one may say that the spectral width of the shear response is essentially governed by the segmental relaxation of the polymeric component.

For the dielectric (derivative) spectra, where in Fig. 5(b) the displayed frequency window is shifted to higher values, it is more appropriate to choose a shorter reference time scale. The spectra displayed in Fig. 5(b) are isochronal with respect to about  $10^{-2}$  s. These dielectric loss data again emphasize that only the high-frequency flank of the spectra is well defined. Interestingly, these flanks are broadest for PVP. If one considers that for PVP the dielectric loss peaks reflect structural rearrangements, while for PVP-PT they reflect the conductivity relaxation, this difference is highly plausible.

Furthermore, Fig. 6 shows that PT acts as plasticizer in PVP, thereby significantly reducing  $T_g$  of the mixtures with increasing *x*. Based on the weight fraction *w* of the components and on the specific heat steps  $\Delta c_p$  of PVP (0.37 J g<sup>-1</sup> K<sup>-1</sup>)<sup>40</sup> and PT (1.3 J g<sup>-1</sup> K<sup>-1</sup>),<sup>82</sup> the observed composition dependence of  $T_g$  is well approximated by the Gordon–Taylor equation<sup>83</sup>

$$T_{\rm g} = \frac{w_{\rm PT}\Delta c_{\rm p,PT} T_{\rm g,PT} + w_{\rm PVP}\Delta c_{\rm p,PVP} T_{\rm g,PVP}}{w_{\rm PT}\Delta c_{\rm p,PT} + w_{\rm PVP}\Delta c_{\rm p,PVP}}$$
(5)

see Fig. 6. Here, we compile the glass transition temperatures  $T_{g,DSC}$  as determined from DSC as well as  $T_{g,\varepsilon}$  and  $T_{g,G}$ , *i.e.*, the temperatures at which the (usually slightly extrapolated) relaxation times  $\tau_{\varepsilon}$  and  $\tau_{G}$ , respectively reach 100 s. The inset of Fig. 6 shows that  $T_{g,DSC}$  can be larger by up to about 30 K than  $T_{g,\varepsilon}$ . The difference between the dielectric and the DSC estimates is maximum for *w* close to 40% (corresponding to *x* = 0.16). This difference again highlights the existence of a dynamical decoupling concerning the rearrangements of the free charge carriers (giving rise to the dielectric relaxation) with respect to their embedding matrix.<sup>20,63</sup>

In the PT-containing electrolytes, the concentration of the (intrinsic) charge carriers should be directly proportional to *x*.



**Fig. 6** Composition dependent glass transition temperatures  $T_g$  of  $PVP_{1-x}PT_x$  as determined from dielectric spectroscopy, rheology, and DSC experiments. The lines are calculated using eqn (5) and the parameters given above that equation. The inset shows that for intermediate weight fractions *w*,  $T_{q,DSC}$  and  $T_{q,e}$  differ significantly.

Therefore, it is interesting to find out whether this expectation is reflected in the conductivity of these mixtures and whether, like in other ionic liquids, factors like the glass transition temperature<sup>84</sup> or the fragility<sup>85</sup> need to be considered in addition. To examine these parameters, we present a  $T_{g,e^-}$ scaled plot, see Fig. 7(a), where we compile the temperature dependent DC conductivities for all PT-containing samples. This Angell plot reveals that the curvature and even the scaled values of the DC conductivities  $\sigma_0$  are the same for all PT mixtures. In this context it is important to recall that  $T_{g,e^-}$ reflects the freezing of the dielectric degrees of freedom, which for decoupled systems corresponds to ionic rearrangements.

The solid line in Fig. 7(a) demonstrates that the DC conductivities  $\sigma_0$  of the binary mixtures can be described using the Vogel–Fulcher law

$$\sigma_0 = \sigma_\infty \exp\left(-\frac{D_\sigma T_0}{T - T_0}\right) = \sigma_\infty \exp\left(-\frac{D_\sigma C_\sigma}{T/T_{g,\varepsilon} - C_\sigma}\right)$$
(6)

here rewritten with the  $T_{g,e}$ -scaled divergence temperature  $C_{\sigma} = T_0/T_{g,e}$ . This way, the data for the  $PVP_{1-x}PT_x$  mixtures can be described by a common set of parameters. These are the prefactor  $\sigma_{\infty} = 2.9 \text{ S cm}^{-1}$ , the strength parameter  $D_{\sigma} = 7.85$ , and  $C_{\sigma} = 0.82$ .



**Fig. 7** (a) DC conductivities  $\sigma_0$  and (b) shear viscosities  $\eta_0$  for all currently studied samples are shown as symbols on reduced inverse temperature scales. The lines are fits (a) using eqn (6) and (b) using eqn (7). In panel (b), the circles refer to viscosities from rotation experiments, all others are from oscillation experiments. The open symbols result from master-curve constructions. The PVP<sub>0.67</sub>PT<sub>0.33</sub> data are scaled once with  $T_{g,G}$  = 216 K (crossed symbols) and once with  $T_g$  = 227 K as taken from DSC. Data for neat PT are from ref. 66.

Fig. 7(b) provides a similar compilation for the shear viscosities of  $PVP_{1-x}PT_x$  as determined using rotational and oscillatory measurements. Here,  $T_{g,G}$  was used for rescaling, and in this representation all viscosity datasets appear to be in good agreement with each other. Like for the DC conductivities  $\sigma_0$ , *cf.* eqn (6), after appropriate rescaling, also the zero-shear viscosities  $\eta_0$  essentially obey a suitably adapted Vogel–Fulcher law

$$\eta_0 = \eta_\infty \exp\left(\frac{D_\eta C_\eta}{T/T_{\rm g,G} - C_\eta}\right) \tag{7}$$

here, the coefficients are  $\eta_{\infty} = 1 \times 10^{-4}$  Pa s,  $D_{\eta} = 4.4$ , and  $C_{\eta} = 1.13$ . Fig. 7(b) also shows that for PVP<sub>0.67</sub>PT<sub>0.33</sub> the scaling using  $T_{\text{g,DSC}}$  (= 227 K) works well, while  $T_{\text{g,G}}$  (which is close to  $T_{\text{g,c}}$ ) does not work. This implies that for the x = 0.33 sample with its rheological double-peak structure, not the higher-frequency peak but merely the hard-to-analyze low-frequency contribution appears to be the more relevant process in the present context.

#### D. Relaxation and diffusion

With the steady-state transport-related quantities  $\eta_0$  and  $\sigma_0$  summarized in Fig. 7 at hand, several well-known hydrodynamics-based approximations have been employed to estimate the corresponding diffusivities, at least in the regime where the melt is sufficiently fluid. With the viscosity given, one may start from the Stokes–Einstein (SE) relation,

$$D_{\rm SE} = \frac{k_{\rm B}T}{6\pi\eta_0 R_{\rm H}} \tag{8}$$

Here, the hydrodynamic radius  $R_{\rm H}$  is related to the van der Waals radius  $R_{\rm vdW}$  of the diffusing moiety, but in many instances the effective  $R_{\rm H}$  is somewhat smaller than  $R_{\rm vdW}$ .<sup>33</sup> Alternatively, for electrolytes such as the current PT mixtures, the Nernst-Einstein relation

$$D_{\rm NE} = \frac{k_{\rm B}T}{nq^2}\sigma_0 \tag{9}$$

is frequently applied. This way, the diffusion coefficient related to the charge q and number density n of the mobile (ionic) species can be determined from the dielectrically probed  $\sigma_0$ .

A different approach of extracting translational diffusion coefficients is based on insights developed by Almond and West (AW) in the framework of describing the electrical conductivity in inorganic solid electrolytes.<sup>64</sup> Basically, AW suggested to identify the crossover frequency, at which  $\sigma'(\nu)$  changes from DC to AC conductivity with an ionic 'hopping rate'  $\Gamma$ . If the effective hopping distance l is known, then one can exploit Einstein's relation to obtain

$$D_{AW,\varepsilon} = \frac{\Gamma l^2}{6} = \frac{\Gamma}{6} \left(\frac{3}{4\pi n_i}\right)^{2/3}$$
(10)

This formalism assumes that *l* can be estimated in a modelindependent way from the ionic number density  $n_i$  according to  $l = [3/(4\pi n_i)]^{1/3}$ ,<sup>63</sup> see Table 1, where *l* is taken to represent the effective radius of the Coulombic cage which extends around each ion. Accordingly,  $\Gamma^{-1}$  can be identified with  $\tau_e$ , except for neat PVP. Here,  $\tau_e$  reflects solely dipolar reorientations

**Table 1** Molecular fractions of the ionic liquid *x*, number densities  $n_i$  of ion pairs, and hydrodynamic radii  $R_H$  for the  $PVP_{1-x}PT_x$  mixtures

$n_i \left(10^{27} \text{ m}^{-3}\right)$	$R_{\rm H}$ (Å)	<i>l</i> (Å)
0.986	0.3	6.2
1.593	0.1	5.3
2.543	0.1	4.5
3.187	0.3	4.2
4.312	0.3	3.8
	$\begin{array}{r} n_i \left(10^{27} \text{ m}^{-3}\right) \\ 0.986 \\ 1.593 \\ 2.543 \\ 3.187 \\ 4.312 \end{array}$	$\begin{array}{c c} n_i \left(10^{27} \text{ m}^{-3}\right) & R_{\text{H}} \left(\text{\AA}\right) \\ \hline 0.986 & 0.3 \\ 1.593 & 0.1 \\ 2.543 & 0.1 \\ 3.187 & 0.3 \\ 4.312 & 0.3 \end{array}$

and is not related to ions (translationally) escaping from a Coulombic cage.

To tackle, in a generalized manner, also situations in which dielectric spectroscopy is not or not clearly providing the relevant information, recently a "rheological" AW formalism has been introduced to characterize the molecular flow in liquids. Here, the crossover frequency is not determined from the conductivity, but from the shear fluidity response.<sup>34</sup> In this approach,  $n_i$  is given by the molecular number density,  $\Gamma^{-1}$  can be identified with  $\tau_G$ ,<sup>34</sup> and the resulting diffusivity is called  $D_{AW,G}$ . Since in the present work the matrix is not a simple liquid but a polymer, in the following the number density relevant for the microscopic flow events will be identified with that of the monomeric units. Note that  $n_m = 6.502 \times 10^{27} \text{ m}^{-3}$  for neat PVP.

To discuss the applicability of these approaches for the  $PVP_{1-x}PT_x$  mixtures, Fig. 8(a) collects the dielectric time scales  $\tau_{e}$ , the rheological time scales  $\tau_G$ , and the ones estimated from DSC,  $\tau_{DSC}$ . As described in the ESI,† the calculation of  $\tau_{DSC}$  follows the procedure reviewed by Hodge.<sup>86</sup> Fig. 8(a) shows that the calorimetric and rheological time scales agree relatively well, but they differ from the dielectric ones. For x = 0.1 we find that  $\tau_{e}$  becomes shorter than  $\tau_{G}$ , signaling an additive-matrix decoupling as pointed out in relation to the inset of Fig. 6.

In Fig. 8(a) another hallmark of this decoupling phenomenon<sup>63</sup> becomes obvious: the temperature variation of the dielectric time scales changes from a relatively strong one above, to a weaker one below the temperature  $T_{g,G}$  at which the rheological dynamics freeze in. As observed in Fig. 8(a), the degree of decoupling increases for x = 0.16 (in agreement with the inset of Fig. 6), but decreases for x = 0.33 and 0.66. In the regime where PT is highly concentrated, the ions themselves constitute the matrix or at least the dominant fraction of it. The decoupling is also reflected by the Walden-type relation,  $\eta_0 \propto \sigma_0^{-\zeta}$ , which, except for x = 1, reveals fractional exponents  $\zeta$  (see Fig. S7, ESI†). Such a fractional relationship or an enhanced translational dynamics are observed also in many other glass-forming materials (see ref. 4–19).

The diffusivities  $D_{\text{SE}}$ ,  $D_{\text{NE}}$ , and  $D_{AW,\varepsilon}$  assessed according to eqn (8), (9), and (10), respectively, as well as  $D_{AW,G}$  estimated according to the rheological counterpart of eqn (10) are shown in Fig. 8(b). These reflect the rheological and conductivity (ionic translational) responses of  $\text{PVP}_{1-x}\text{PT}_x$ . For the compositions containing the ionic liquid,  $R_{\text{H}}$  was obtained by imposing an agreement between  $D_{\text{SE}}$  and  $D_{\text{NE}}$  at the highest investigated temperatures. The corresponding  $R_{\text{H}}$  values are included in



**Fig. 8** (a) Arrhenius plot of the time scales  $\tau_{er}$ ,  $\tau_{Gr}$ , and  $\tau_{DSC}$  determined for  $PVP_{1-x}PT_x$  in the present work, except for neat PT (x = 1) where  $\tau_e$  was taken from ref. 66. Full symbols reflect time scales directly from the loss peaks, open symbols are from shear modulus master curves. The  $\times$  symbol reflects the relaxation time corresponding to the low-frequency shear loss peak marked in Fig. 4(b) for x = 0.33. (b) The diffusion coefficient  $D_{SE}$  (solid squares) was calculated using eqn (8) from  $R_{\rm H}$  in Table 1 and from  $\eta_0$ . For the half-filled squares,  $\eta_0$  was extracted from the shift factors used for the shear modulus master curves.  $D_{\rm NE}$  (asterisks) was computed using eqn (9) from  $\sigma_0$  as shown in Fig. 7(a) and  $n_i$  in Table 1. The calculation of  $D_{\rm AW}$  is based on eqn (10) and the data presented in panel (a): For  $D_{\rm AW,G}$  (stars) we used  $\Gamma = 1/\tau_G$  and for  $D_{\rm AW,e}$  we used  $\Gamma = 1/\tau_e$ . The open stars refer to time scales from the shear modulus master curves. For the determination of  $D_{\rm AW,e}$  for neat PVP see the text.

Table 1, together with the calculations for the number density  $n_i$  of the ion pairs.

For the current  $D_{\text{NE}}$  estimates the correlated charge-carrier dynamics, which is known to reduce the conductivity in concentrated electrolytes,<sup>63,87</sup> have been neglected. Consequently, the extracted  $R_{\text{H}}$  values may be underestimated, at least for the systems with large concentrations of ionic liquid. Similar rheological cooperativity effects have been recently advocated to explain the large differences between  $D_{\text{SE}}$  and  $D_{\text{AW,G}}$  determined for several van der Waals liquids near their glass transition.<sup>34</sup> Cooperativity effects may also rationalize the relatively large contrast between  $D_{\text{SE}}$  and  $D_{\text{AW,G}}$  seen in Fig. 8(b) for x = 0.66 and 1. These samples correspond to "liquid"-matrix materials. Since in ionic liquids mass and charge flows are coupled with each other,<sup>66</sup> one may expect similarly large differences between the conductivity-related diffusivities  $D_{\text{NE}}$  and  $D_{\text{AW,e}}$ , as also revealed by Fig. 8(b). However, this

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observation contrasts with the knowledge gained from investigations of weakly and moderately supercooled regimes of ionic liquids that charge correlation effects are not that strong in these materials.<sup>62,63</sup> Further investigations, preferably involving direct diffusivity measurements of deeply supercooled ionic liquids would be beneficial in clarifying these seemingly contrasting observations.

Despite the complexity implied by the, in general, differently temperature-dependent time scales that are summarized in Fig. 8(b), several common trends can be recognized:

(i) For all mixtures, the lowest diffusivity estimates are those based on the SE relation, eqn (8). This is barely surprising, considering that the macroscopic viscosity  $\eta_0$  corresponds to the terminal flow of the polymer chains. Although these are not very long (note the relatively low molecular weight of PVP), the probed  $\eta_0$  should be larger than the microscopic viscosity governing the local friction of the monomers (x = 0) or of ionic plasticizers (0 < x)< 1). Additionally, in the high-viscosity regime the SE approach is to be considered with caution since it neglects the decoupling phenomenon. The effects of a dynamic decoupling that were mentioned in the introduction section were shown early on to play a significant role for the diffusion of nonionic additives near the glass transition temperature of polymeric materials.<sup>5,24</sup>

(ii) For the low-concentration electrolytes (x = 0.1 and 0.16), the NE predictions based on the macroscopic conductivity agree excellently with the AW calculations which are based on the microscopic ionic relaxation time  $\tau_{\varepsilon}$ . This observation confirms that the elementary step for single-ion diffusion corresponds to the radius *l* of the Coulombic cage. For the low-charge concentrations l is much larger than the effective steric radius of the monomeric units.

(iii) The agreement between the collective (NE based) and the single-particle (AW based) diffusivities breaks down at high charge concentrations (x = 0.66 and 1). The relationship  $D_{\rm NE} <$  $D_{AW,\varepsilon}$  prevailing in this regime indicates that cooperative effects emerge which suppress the conductivity, an effect usually quantified in terms of the Haven ratio.

(iv) At variance with the situation for x = 0.1 and 0.16, for the electrolytes with x = 0.66 and 1 the diffusivities  $D_{AW,\varepsilon}$  and  $D_{AW,G}$ are similar. This finding indicates again that local-charge and matrix dynamics are decoupled at low x and more coupled in the ion-rich materials.

(v) The time scales from DSC are typically close to those extrapolated from the rheological measurements with the most significant deviation observed for x = 0.33. However, for this sample the relaxation time from the low-frequency shear peak, see the crosses ( $\times$ ) in Fig. 8(a), is not too far from  $\tau_{\rm DSC}$ .

### IV. Concluding remarks

To summarize, the present work explores the frequencydependent dielectric and shear rheological responses of PVP with a molecular weight of  $2500 \text{ g mol}^{-1}$ , and of mixtures of this polymer with the ionic liquid PT. Differential scanning calorimetry has additionally been employed. The dielectric and mechanical responses of PVP are dominated by the structural relaxation process of this material. As is typical for neat glassforming systems, the characteristic time scales extracted from the analysis of the shear modulus function of PVP are somewhat smaller than those from its dielectric compliance. However, with the addition of an ionic liquid the dielectric response of PVP-PT becomes faster than its mechanical counterpart. Corroborated by the significant differences found between the glass transition temperatures accessed via calorimetry and those estimated from the dielectric dynamics, and by the sigmoidal-like variation of the logarithm of dielectric time scales as a function of inverse temperature, this observation indicates that a decoupling of charge dynamics from segmental rearrangements emerges in PVP-PT at low and intermediate ionic concentrations.

Although signs of crystallization have not been detected for any of the investigated systems, an anomalously broad shear response and a bimodality of the calorimetric response indicate the emergence of a microscopic phase separation for a molar PT concentration near 33%. Disregarding this concentration, the analysis of the glass transition temperatures for the other mixtures reveals that the ionic-liquid additive is a good plasticizer for PVP. This is clearly seen from the concentration dependent  $T_{\rm g}$  which can be described well by the Gordon-Taylor equation. Free parameters are not necessary for this approach, since the corresponding specific heat steps are directly available from calorimetric investigations.

The parameterization of the dielectric and rheological spectra allowed us to extract dynamic and transport coefficients such as the characteristic times, steady-state viscosities, and steady-state conductivities. Using these parameters, we estimated the translational diffusivities of the ionic additive PVP, based on the Stokes-Einstein and Nernst-Einstein relations as well as on the conductivity and mechanical versions of the Almond-West formalism. Although the present case - with the dynamics of the additive being faster than the matrix rearrangements - constitutes a clear deviation from the hydrodynamic limit, these approaches provide several interesting observations: first, for all materials, the smallest diffusivities are those estimated using the SE relation. Second, the NE predictions that are based on the macroscopic conductivity and the AW calculations based on the microscopic ionic relaxation time agree well at low ionic concentration, while this agreement breaks down for large ion densities.

Our study reveals that mixing ionic liquids with polar polymers enables the investigation of charge dynamics in the full concentration range by overcoming the dissociation and solubility issues of salt-doped polymer electrolytes. The results obtained in this work thus are relevant for understanding and controlling the microscopic transport in amorphous dispersions and polymer electrolytes near their glass transition.

### Conflicts of interest

There are no conflicts of interest to declare.

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