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Phenol, the simplest aromatic monohydroxy alcohol, displays a faint Debye-like process when mixed with a nonassociating liquid†

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Solvated in propylene carbonate, viscous phenol is studied using dielectric spectroscopy and shear rheology. In addition, several oxygen-17 and deuteron nuclear magnetic resonance (NMR) techniques are applied to specifically isotope labeled equimolar mixtures. Quantum chemical calculations are used to check the electrical field gradient at phenol's oxygen site. The chosen combination of NMR methods facilitates the selective examination of potentially hydrogen-bond related contributions as well as those dominated by the structural relaxation. Taken together the present results for phenol in equimolar mixtures with the van der Waals liquid propylene carbonate provide evidence for the existence of a very weak Debye-like process that originates from ringlike supramolecular associates.

1. Introduction

Hydrogen bonding is at the core of molecular associations in all sorts of condensed matter. Depending on the kind of moieties that are in the hydrogen bonding “soup”¹ and subject to the geometrical constraints imposed by the molecular architecture, a wealth of different aggregates can emerge in solids and liquids. The formation of hydrogen bonding patterns has been studied intensively in self-associating liquids such as linear and branched monohydroxy alcohols.² Due to the typically small hydrogen bonding strength,³ transient supramolecular patterns often evolve⁴ that can be modulated by changes in external parameters such as temperature, pressure, external field, or the degree of dilution.^{5,6}

The monohydroxy alcohols are probably among the molecularly simplest glassforming liquids for which, depending on the position of the OH group on the aliphatic backbone, ring- or chain-type association patterns are favored. For a long time it was thought that phenyl groups are effective in reducing the impact of hydrogen bonding, not the least because the presence of the relatively bulky aromatic rings can act as a source of considerable steric hindrance for the association of OH groups.⁷ Recent scrutiny, involving systematic variations of the “side-group” that is attached to the phenyl ring, shows that matters are more complex: In addition to steric effects and O-H···O hydrogen bonding,^{3,8} also the π-π interactions among the phenyl rings,⁹ as well as O-H···π based associations need to be taken

into consideration,^{10–15} albeit not always conforming to the classical notion of “π stacking”.¹⁶ Owing to their ability to supercool easily and to form glasses, recently the isomeric phenyl propanols received particular attention: Using light-scattering-assisted,¹⁷ shear mechanical,¹⁸ as well as high-field electrical experiments,¹⁹ an important finding for these liquids is that, at least in their highly viscous state, a supramolecular Debye-type process exists that is slower than the structural relaxation.

For a long-time, the Debye process in monohydroxy alcohols was considered a phenomenon that is only accessible *via* dielectric spectroscopy.⁵ In these liquids, the amplitude of the Debye relaxation can be about 100 times larger than that of the structural relaxation,²⁰ but also three times smaller than the latter.^{6,21,22} It has been recognized that the Debye process leaves its mark in other quantities such as an OH-related mode as detected using NMR relaxometry⁴ and neutron spin-echo techniques²³ as well as in a polymer-like viscosity enhancement.²⁴

Without reference to these latter observations, but rather by comparison with results from light scattering, more recently it has been suggested that a Debye process could be featured in the dielectric response also of nonassociating glass forming liquids.²⁵ This recent suggestion underscores that for an identification of a Debye process that is akin to that in associating liquids, in particular to that in aliphatic monohydroxy alcohols, it is advisable to rely not just on dielectric measurements. Therefore, in addition to dielectric spectroscopy, in the present work we will apply shear rheology and several nuclear magnetic resonance (NMR) techniques to check whether or not in a mixed liquid in which the only associating component is phenol (PhOH), the simplest aromatic monohydroxy alcohol, a hydrogen-bond related Debye process does exist.

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From the present perspective, a comparison with other members of the homologous $C_6H_5-(CH_2)_n-OH$ series is particularly intriguing.^{11,17} While initially it was thought that due to effects of sterical hindrance the Debye-type relaxation might not generally be observable in liquids like 1-phenyl-2-propanol,⁷ recent evidence shows that also here a Debye relaxation does exist, but it weakens as the OH group gets closer to the phenyl ring.¹⁷ These findings raise the question whether in the $n = 0$ (monohydroxybenzene) limit, *i.e.*, for PhOH, the sterical hindrance becomes overwhelming so that the formation of hydrogen bonded structures is precluded or whether the hydrogen bond interactions are sufficiently effective, so that a Debye process is observable nevertheless.

X-ray diffraction and infrared absorption techniques were exploited to map out the interplay between dispersive and hydrogen bond interactions in the $C_6H_5-(CH_2)_n-OH$ series upon systematically varying the length of the linear alkyl chain ($n = 1-5, 7$) that is attached to the ring moiety.¹¹ Obviously, the $n = 0$ case, PhOH, where the OH group is directly attached to the aromatic ring, was not considered in this context, so far. Presumably, this is because C_6H_5-X with $X = OH$, which melts near 314 K,²⁶ but also the substances with $X = SH$ or SeH , display a pronounced crystallization tendency.²⁷ At ambient pressure, the molecules in crystalline PhOH form three-fold hydrogen bonded helices.²⁸ Along them the OH-related dipole moments are aligned in an antiparallel fashion.²⁹ With reference to the fact that the ambient-pressure structure of PhOH is built from trimeric repeat units, a Raman study of the pure liquid, emphasizes the important role that intermolecular hydrogen bonding plays in stabilizing three-membered phenol clusters.³⁰ When PhOH is compressed to 0.16 GPa, two-fold ribbon-like structures evolve, along which the dipole moments are arranged in a parallel fashion.^{29,31}

Glass formation has been reported for phenol dissolved in relatively concentrated mixtures with hydrogen bonded liquids such as ethanol,³² glycerol/water solutions,³³ or deep eutectic choline chloride binaries.^{34,35} Furthermore, when embedded in a host of solvents near and above room temperature, PhOH was examined using vibrational spectroscopy also under more highly diluted conditions.^{30,36-38} PhOH does not form vitrifying mixtures with other simple aromatic ring molecules such as toluene³² or (as a result from the present work also not with) 2-picoline, while for PhOH diluted in an *ortho*-terphenyl/benzene solution glass formation could apparently be enforced by shock freezing to liquid nitrogen temperature.³⁹

Attempts that we undertook to vitrify pure PhOH by quenching from about 350 to 77 K with rates exceeding 15 K s⁻¹ failed. Therefore, with the goal to examine the relaxational contributions associated with the phenolic hydrogen bonds nevertheless, for PhOH a nonassociating mixing partner devoid of hydroxyl groups is to be sought. In this context, it should be emphasized that studies of supramolecular association in aliphatic monohydroxy alcohols often exploit dilution with a mixing partner that is unable to sustain hydrogen bonds: Typically, in 50:50 mixtures and sometimes even for smaller alcohol concentrations, the Debye process can persist almost unaltered, albeit reduced in intensity.^{6,40-47}

The solvent that is suitable to dilute PhOH should exhibit a good ambient-pressure glassforming propensity⁴⁸ to avoid the need for the application of high cooling or heating rates. Furthermore, we presumed that in order to ensure miscibility, the mixing partner should be polar and possess a molecular weight comparable to that of PhOH. Conforming to these provisions, in the present work we will show that propylene carbonate (PC) is a suitable choice. This van der Waals glass former, with a glass transition temperature T_g of about 160 K,^{49,50} was studied already by, *e.g.*, dielectric spectroscopy,⁵¹⁻⁵⁴ shear rheology,⁵⁵ and NMR.⁵⁶ Furthermore, PC was studied by numerous other techniques as well.⁵⁷⁻⁶⁰ PhOH, a very weak acid, and PC, a very weak base,⁶¹ are not expected to react with each other.⁶² A potential drawback of choosing PC as a mixing partner may be its large dipole moment of about 5 D,⁶³⁻⁶⁵ while for PhOH some solution studies indicate an “infinite dilution” value of “only” 1.5 D. The sizeable dependence of PhOH’s dipole moment on concentration and solvent could be taken as a hint for self-association effects.^{66,67}

The dipolar response of PhOH mixtures with PC are thus expected to comprise significant contributions also from the van der Waals liquid. To enable an additional perspective on the relaxation properties of PhOH in PC that is not sensitive on the molecular dipole moments, apart from dielectric spectroscopy, we performed shear mechanical measurements. Furthermore, to achieve a fully selective detection of the various components in the mixture, we will exploit isotope edited NMR. In other words, for the present NMR studies unlabeled substances were mixed with ring-deuterated PC, ring-deuterated PhOH, or ¹⁷O labeled PhOH. Thus, while the PC-d₃ and the PhOH-d₃ molecules can be expected to be most sensitive to the structural relaxation, the oxygen-17-labeled probe is closest to and should therefore be most sensitive to the dynamics at the hydrogen bonding site.

2. Experimental details

Fully protonated PC and PhOH (nominal purities $\geq 99\%$) were acquired from Alfa Aesar and Sigma Aldrich, respectively. PC-d₃ was taken from the batch used in ref. 56. Phenol-2,4,6-d₃ (PhOH-d₃, ≥ 98 atom% D) was purchased from Sigma Aldrich. The Ph¹⁷OH sample (¹⁷O enrichment 35.8 atom%, purity $\geq 98\%$) was obtained from Eurisotop. To prepare samples with the desired molar fraction, x_{PhOH} , directly in the dielectric cell or in the NMR glass tube, PhOH was first weighed under a fume hood. Then, the appropriate amount of PC was pipetted into the cell or tube where during stirring in or shaking of these containers, PhOH quickly dissolved in PC. To remove potentially adverse effects of paramagnetic O₂, the filled NMR tubes were degassed *via* the freeze-pump-thaw method and hermetically flame-sealed afterwards.

For the rheological measurements of a sample with $x_{PhOH} = 0.5$ we employed an MCR 502 rheometer from Anton-Paar. A premixed PC:PhOH solution was pipetted onto the lower shear plate of a 4 mm parallel-plate geometry. To control and stabilize the temperature to within 0.2 K, an EVU20 unit and a



CTD450 oven were used. Dielectric spectroscopy experiments were carried out on fully protonated PC:PhOH mixtures (or an equimolar mixture of PhOH and 2-picoline, data not shown) using a Novocontrol Alpha Analyzer in conjunction with a Quatro temperature controller from Novocontrol Co. The utilized sapphire/invar cell was stabilized to the desired temperatures to within ± 0.1 K.

The NMR experiments were conducted using home-built equipment. The ^2H spectrometer was operated at a Larmor frequency of $\omega_D = 2\pi \times 45.6$ MHz. The $^{17}\text{O}/^1\text{H}$ double-resonance spectrometer was operated at $\omega_O = 2\pi \times 54.2$ MHz and $\omega_H = 2\pi \times 400.1$ MHz in conjunction with a probe head from NMR Service Co. Below 200 K, ^1H decoupling employed a power level corresponding to a Rabi frequency of 167 kHz in order to effectively remove unwanted hydroxyl-proton impacts on the oxygen magnetization. The oxygen-17 spectra are referenced with respect to liquid $^1\text{H}_2^{17}\text{O}$ at 291 K. The ^2H pulses were $t_{D,\pi/2} = 2.0$ to 2.5 μs long and for the excitation of the central ($-1/2 \leftrightarrow +1/2$) ^{17}O transition $t_{O,\pi/2}$ was 1.5 μs . During each NMR experiment the temperature stability was typically better than ± 0.2 K.

3. Results and analyses

Below, in Section 3A we will first present dielectric measurements on PC:PhOH mixtures and show how their spectra can be decomposed into a Debye-like and another contribution. Then, based on rheology, we will determine the so-called viscosity enhancement which is able to signal the presence of even weak supramolecular modes. Section 3B focuses on PC:PhOH's NMR spectra. These are required, for instance, for the detailed analysis of the ^{17}O spin relaxation times that are dealt with in Section 3C. On a similar basis, Section 3D provides time-scale information by employing deuteron NMR. For aliphatic monohydroxy alcohols, close to the site where hydrogen bonding occurs, the correlation times (determined using ^{17}O NMR) are found to be longer than those at the alkyl rest (accessible, *e.g.*, using ^2H NMR) that probes the structural relaxation.^{4,5,68} Sections 3C and D aim at gathering all the (mostly time scale) information that is needed to check whether or not PC:PhOH displays a pattern similar to that which characterizes the well-established Debye liquids.

A. Dielectric spectroscopy and shear rheology

Fig. 1 shows the real and the imaginary part of the complex dielectric function $\epsilon^* = \epsilon' - i\epsilon''$ of PC:PhOH. Upon cooling, well defined steps in ϵ' , *cf.* Fig. 1(a), and well defined asymmetric loss peaks in ϵ'' , *cf.* Fig. 1(b), shift through the accessible frequency window. Above about 175 K the spectral shapes remain more or less invariant, implying that frequency temperature superposition is approximately fulfilled. Then at lower temperatures, steps and peaks which refer to the dominant relaxation of PC:PhOH are no longer detected. Now, another process that is characterized by an enormously large spectral width and a very small relaxation strength is visible. This symmetrically broadened relaxation can be assigned to a Johari–Goldstein process.

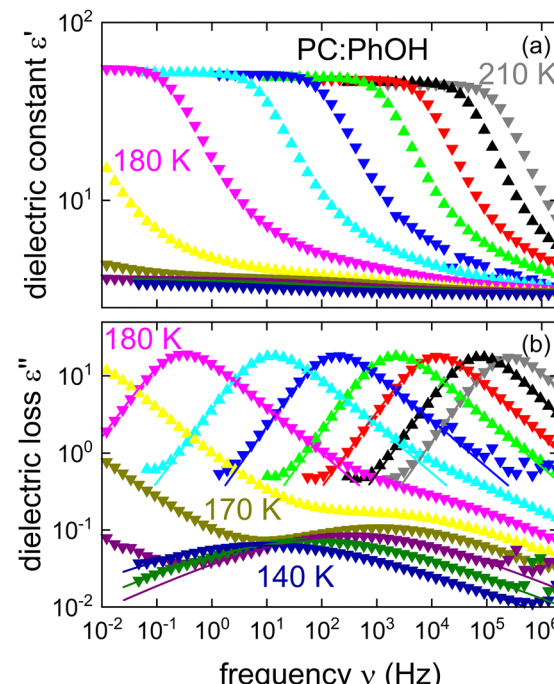


Fig. 1 Dielectric constant (a) and dielectric loss (b) of an equimolar mixture of phenol and propylene carbonate. The lines are fits using a Cole–Davidson function (for the dominant process) or Cole–Cole function (for the β process). Data are shown in 10 K steps from 140 to 170 K and in 5 K steps at higher temperatures.

As indicated by the solid lines in Fig. 1, the peaks and steps referring to the dominant relaxation are nicely describable using a Cole–Davidson function, *i.e.*, by a Havriliak–Negami function

$$\epsilon^*(\nu) = \epsilon_\infty + \frac{\Delta\epsilon}{[1 + (2\pi i\nu\tau_{\text{HN}})^{\alpha_{\text{CC}}}]^{\gamma_{\text{CD}}}} + \frac{\sigma_0}{2\pi i\nu\epsilon_0}, \quad (1)$$

in the limit of $\alpha_{\text{CC}} = 1$. Then, the Cole–Davidson exponent γ_{CD} that appears in eqn (1) governs the asymmetry of the spectral shape. For $\alpha_{\text{CC}} < 1$ and $\gamma_{\text{CD}} = 1$ eqn (1) reduces to the Cole–Cole function which can be used to parameterize the symmetrically broadened low-temperature loss curves. In eqn (1) $\Delta\epsilon$ denotes the relaxation strength, where $\Delta\epsilon = \epsilon_s - \epsilon_\infty$ is the difference between the low- and the high-frequency permittivity. From eqn (1) the relaxation time τ corresponding to the maximum dielectric loss is calculated according to⁶⁹

$$\tau = \tau_{\text{HN}} \left[\frac{\sin(\alpha_{\text{CC}}\gamma_{\text{CD}}\pi/(2 + 2\gamma_{\text{CD}}))}{\sin(\alpha_{\text{CC}}\pi/(2 + 2\gamma_{\text{CD}}))} \right]^{1/\alpha_{\text{CC}}}. \quad (2)$$

For the dominant relaxation of PC:PhOH (apart from $\alpha_{\text{CC}} = 1$) we find $\gamma_{\text{CD}} = 0.6 \pm 0.05$, a value typical for many neat supercooled liquids.⁷⁰ However, for binary mixtures often times one also observes an enhanced broadening, hence smaller γ_{CD} .⁷¹ In this context, we note that for the dielectrically detected α -process of neat PC an exponent $\gamma_{\text{CD}} = 0.66$ was reported.⁵¹

A direct comparison of the dielectric function of PC at 180 K with that for the equimolar PhOH mixture is shown in Fig. 2. Here, one recognizes that the alcohol addition slows the

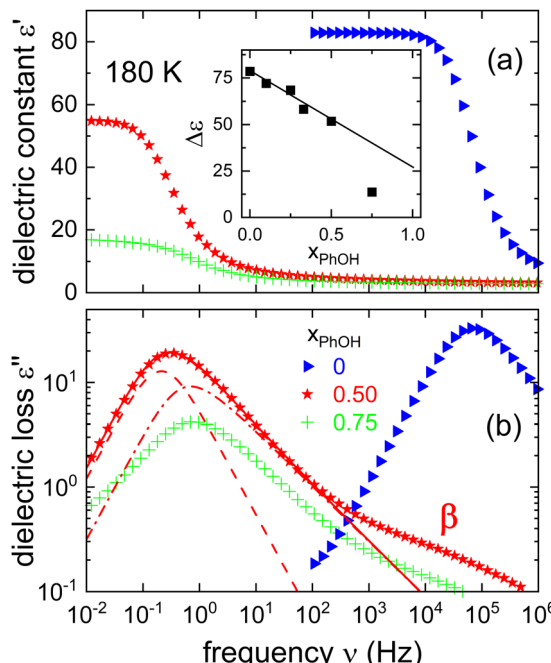


Fig. 2 Isothermal comparison of (a) the real and (b) the imaginary part of the dielectric function ϵ^* as measured for different molar fractions x_{PhOH} at 180 K. For this temperature, the inset shows the relaxation strengths as a function of x_{PhOH} . For $x_{\text{PhOH}} \leq 0.5$ we find that $\Delta\epsilon$ displays a linear trend, while for $x_{\text{PhOH}} = 0.75$ the reduced $\Delta\epsilon$ is indicative for a partial crystallization. The line in the inset is calculated using eqn (23) and the information given below that equation. The dashed and the dash-dotted lines in frame (b) result from a decomposition of the main peak in terms of a Debye-like ($\gamma_{\text{CD,D}} = 1$) and an α -type contribution ($\gamma_{\text{CD,}\alpha} = 0.53$), respectively. The decomposition is not entirely unique, but with the chosen $\gamma_{\text{CD,}\alpha}$ the time constants of the two processes differ by a factor of about 3.3.

dominant dynamics of the mixture by about 5 decades. This finding suggests that if PhOH could be vitrified in pure form, its effective T_g would be much larger than that of neat PC.^{49,50} Fig. 2 also includes a dielectric spectrum recorded for a binary mixture featuring a molar PhOH fraction of 75%. The appearance of a strongly reduced relaxation strength in that sample and of a peak frequency similar to that of the 50% mixture indicate a limited miscibility beyond this latter PhOH fraction. A comparison of the relaxation strengths also including several other compositions (see the inset of Fig. 2) confirms the overall trend and furthermore that the equimolar sample does not show signs of segregation.

So far we assumed that the relaxation strength essentially originates from a monomodal main peak. However, also another interpretation is possible by assuming a recently suggested decomposition into a Debye-like (D) and an α -type contribution that was applied in analyses of various phenyl alcohols.^{17–19} For comparison purposes, the results of such a decomposition are included in Fig. 2 (see also Fig. S1 in the ESI†).

As already pointed out, from Fig. 1(b) it is obvious that PC:PhOH exhibits a well-defined secondary process. To describe the broad peaks referring to this so called β relaxation, eqn (1) was used again but now in the Cole–Cole limit with $\alpha_{\text{CC}} = 0.33 \pm 0.03$ and $\gamma_{\text{CD}} \approx 1$. The appearance of a Johari–Goldstein

relaxation in PC:PhOH is remarkable because in neat PC such a feature could be resolved *via* broadband dielectric spectroscopy only after excessive physical aging.⁷² Yet, for glassformers like toluene or *meta*-fluoroaniline, which involve substituted aromatic rings, the existence of a prominent Johari–Goldstein relaxation is well known.^{73,74}

In the search for dynamical signatures of supramolecular structures, also shear rheology plays an important role. Therefore, we carried out measurements of the complex shear modulus $G^* = G' + iG''$ and in Fig. 3(b) we show $G''(\nu)$ for PC:PhOH at different temperatures. The slowdown of the dynamics that occurs with decreasing temperature leads to a down-shift of the loss peak frequencies that reflect an increase of the associated time constants τ_G .

While the evolution of $G''(\nu)$ looks similar to that revealed by the dielectric loss, one has to realize that in Fig. 1 we plot ϵ^* , *i.e.*, the dielectric susceptibility, while in Fig. 3(b) a modulus is presented. For a more consistent comparison, it is advisable to use the mechanical equivalent of ϵ^* , *i.e.*, the mechanical compliance $J^* = J' - iJ'' = 1/G^*$. From Fig. 3(a) one recognizes well defined steps in $J'(\nu)$, akin to the situation for $\epsilon'(\nu)$. Peaks in $J''(\nu)$, that one may naively expect in analogy to the situation for $\epsilon''(\nu)$, are not observed. This is because, similar to the dc

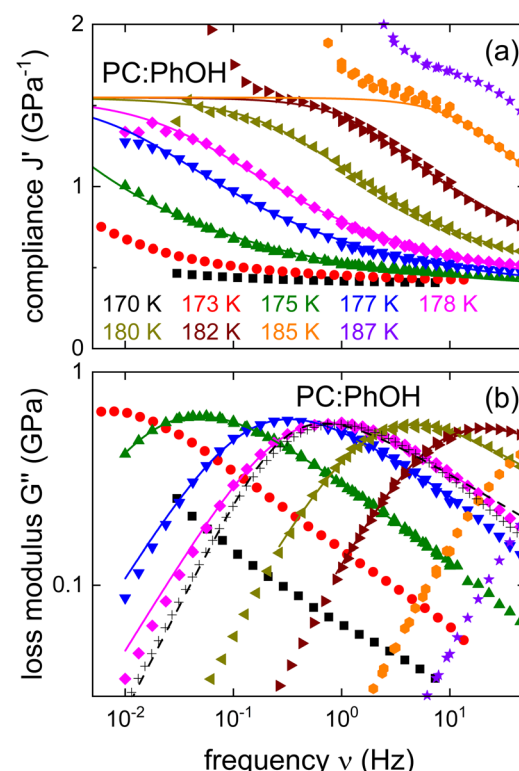


Fig. 3 Shear mechanical response of PC:PhOH, here represented as (a) real part of the compliance and (b) imaginary part of the loss modulus. The solid lines represent fits and calculations using eqn (3) and the parameters given in the text. The black crosses mark appropriately shifted data for DC704 (for $T = 218$ K) a simple liquid devoid of supramolecular modes.^{75,76} For 178 K the dashed line represents a Cole–Davidson curve normalized to the maximum of G'' . The low-frequency upturns seen in J' above about 1.5 GPa^{-1} reflect the low-torque resolution limit of our rheometer.

conductivity in $\epsilon''(\nu)$, for the compliance the viscous flow leads to a $1/\nu$ divergence in $J''(\nu)$. This contribution is so prevalent that it completely masks any peaks in the loss part of the compliance.

Nevertheless, analogous to the dielectric data, one may try to describe the mechanical ones using, *e.g.*, a Havriliak–Negami function here written as

$$J^* = J_\infty + \frac{\Delta J}{[1 + (2\pi\nu\tau_{J,\text{HN}})^{\alpha_J}]\gamma_J} + \frac{1}{2\pi\nu\eta}. \quad (3)$$

In eqn (3) J_∞ and J_0 denote the limiting high-frequency compliance and the recoverable compliance, respectively, so that the compliance relaxation strength is $\Delta J = J_0 - J_\infty$. The distribution parameters employed to calculate the solid lines in Fig. 3 are $\alpha_J = 0.75 \pm 0.05$ and $\gamma_J = 0.40 \pm 0.05$. Analogous to eqn (2) the timescale τ_J is derived from $\tau_{J,\text{HN}}$. From the fits with the fixed parameters $\Delta J = 1.2 \times 10^{-9} \text{ Pa}^{-1}$ and $J_\infty = 3.5 \times 10^{-10} \text{ Pa}^{-1}$, see the solid lines in Fig. 3(a), we determined τ_J . The same fit parameters are employed to calculate the shear loss modulus G'' . While the maximum region is described very well, see Fig. 3(b), deviations of the fits from the experimental data are observed on the low-frequency flank. In fact, the measured shear loss of PC:PhOH displays a well-defined terminal mode leading to $G'' \propto \nu^1$, as expected for low-molecular weight liquids. This is also seen for the data of the pump oil tetramethyl-tetraphenyl-trisiloxane (DC704), *cf.* Fig. 3(b), a liquid completely unsuspicious of sustaining such modes.^{75,76} The above power law is obtained in the Cole–Davidson limit of eqn (3). Therefore, we also included a Cole–Davidson curve in Fig. 3(b) which matches the high-frequency flank very well. Clearly, on the low-frequency side this curve underestimates the observed shear loss. Significant deviations of this kind are the hallmark of so called viscosity enhancements⁷⁷ which hint at the presence of supramolecular modes. For aliphatic monohydroxy alcohols and other systems displaying such modes, it was shown that the viscosity enhancement can be quantified using the scaled zero-shear viscosity⁷⁷

$$z_0 = 2\pi\nu_{\max}\eta'(\nu \rightarrow 0)/G_\infty. \quad (4)$$

Here, $G_\infty = 1/J_\infty$ designates the limiting high-frequency shear modulus and $\eta'(\nu) = G''(\nu)/(2\pi\nu)$ defines the frequency dependent shear viscosity. One finds $z_0 \approx 1.0 \pm 0.05$ for nonassociating liquids like DC704 and $z_0 \approx 10$ for monohydroxy alcohols such as 2-ethyl-1-hexanol that exhibit a strong Debye process.⁷⁷ For PC:PhOH we obtain $z_0 \approx 1.2 \pm 0.1$, *i.e.*, evidence for a weak, yet readily detectable supramolecular mode.

B. ^{17}O and ^2H NMR spectra

Using NMR, information regarding molecular dynamics can for instance be gathered by tracking motional narrowing effects. Furthermore, from the low-temperature spectra, the quadrupolar coupling constant $C_Q = eQV_{zz}/h$ and the asymmetry parameter $\eta = (V_{xx} - V_{yy})/V_{zz}$ become accessible. These quantities involve the components $|V_{zz}| \geq |V_{yy}| \geq |V_{xx}|$ of the electrical field gradient tensor at the nuclear site and are necessary for the analysis of the spin-relaxation times that will be discussed in Section 3C, below.

Fig. 4 shows ^{17}O central-transition spectra that were recorded for PC:Ph ^{17}OH in a wide temperature range. Near ambient temperature a motionally narrowed spectrum is observed, indicating that here the motional correlation times are short on the scale of $\tau \approx 1/\omega_0 \approx 2.9 \text{ ns}$ that is set by the inverse Larmor frequency. Then, upon cooling the absorption spectra broaden, shift their center of gravity, narrow again, and eventually broaden into a well-structured pattern. As the analysis of these data will reveal, this seemingly unintuitive sequence of temperature dependent effects is in fact characteristic for slowing dynamics as probed by nuclei that are subjected to strong quadrupolar interactions.

Above about 250 K, the spectra are peaked near 4.6 kHz, corresponding to an isotropic chemical shift $\delta_{\text{iso}} = 86 \text{ ppm}$ in the high-temperature limit. Due to the appearance of very short T_2 times ($\lesssim 10 \mu\text{s}$), see Section 3C, below, spectra could not be recorded between about 250 and 215 K. From the data below

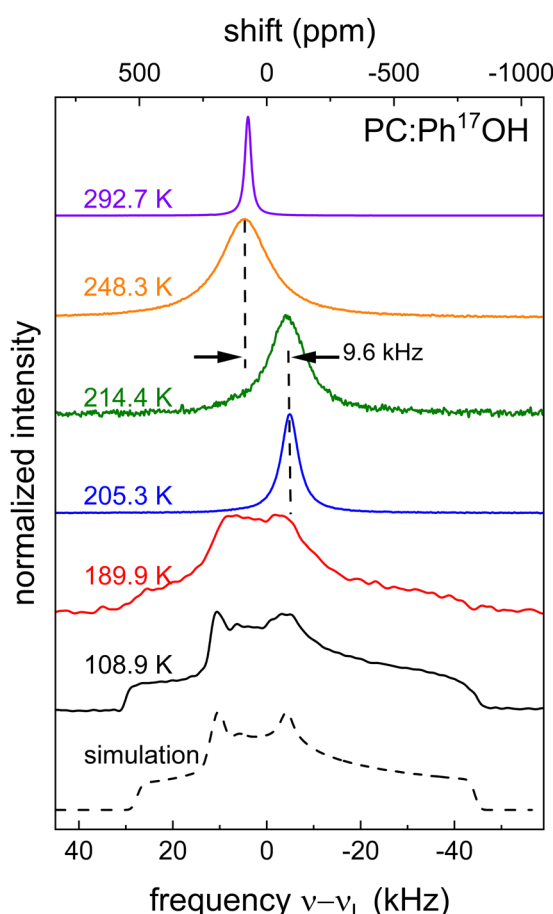


Fig. 4 Temperature dependent ^{17}O central-transition NMR spectra of an equimolar mixture of Ph ^{17}OH and PC. Below 215 K, a 90° – $25 \mu\text{s}$ – 180° pulse sequence was applied. Above this temperature the spectra are excited with a single pulse. Additionally, at the lowest two temperatures proton decoupling was used and zero-filling extended the time signal from 0.5 to 16 ms. A Gaussian apodization with $\sigma/2\pi = 100 \text{ Hz}$ was applied, except for 189.9 K where $\sigma/2\pi = 500 \text{ Hz}$. The dashed line represents a simulation using $C_Q = 8.7 \text{ MHz}$ and $\eta = 0.72$ with an apodization of $\sigma/2\pi = 2 \text{ kHz}$.

this latter temperature it is evident that the spectra have undergone a so-called dynamic frequency shift.⁷⁸ This shift amounts to about 9.6 kHz, see Fig. 4, corresponding to $\Delta\delta = 177$ ppm.

In the framework of second-order perturbation theory, the dynamic frequency shift can be calculated to be⁷⁸

$$\Delta\delta = -\frac{3}{40} \frac{I(I+1) - \frac{3}{4}}{I^2(2I-1)^2} \frac{C_Q^2 \left(1 + \frac{1}{3}\eta_O^2\right)}{\nu_O^2}. \quad (5)$$

$=3/500$ for $I=5/2$

Thus, from the experimentally determined $\Delta\delta$ a quadrupolar product $C_Q \left(1 + \frac{1}{3}\eta^2\right)^{1/2}$ of 9.3 MHz results for spin $I = 5/2$ nuclei. To disentangle the values of C_Q and η_O , the low-temperature spectrum shown in Fig. 4 can be simulated using a suitable combination of these parameters. The dashed line in Fig. 4 reflects a spectrum simulated by means of WSolids⁷⁹ using parameters that within experimental uncertainty are compatible with the quadrupolar product inferred from $\Delta\delta$.

In addition to recording ¹⁷O spectra, also ²H spectra were sampled. To achieve spectral selectivity we investigated suitably labeled PC:PhOH mixtures, *i.e.*, PC:PhOH-d₃ and PC:PhOH-d₅. It turns out that we do not detect lineshape changes of the type observed, *e.g.*, in binary mixtures comprising constituents of significantly differing molecular weight.⁸⁰ Therefore, as Fig. S2 in the ESI† we provide only low-temperature spectra for PC-d₃ and PhOH-d₃ in the corresponding mixtures. From these spectra we determine the ²H quadrupolar coupling constants which are often expressed in terms of the anisotropy parameter $\delta_Q/2\pi = \frac{3}{4}C_Q$.⁸¹ For neat PC-d₃ an anisotropy parameter $\delta_{PC}/2\pi = 125$ kHz and a vanishing asymmetry parameter were reported.⁵⁶ In the mixture we find the same values. For PhOH (in fact for ring-deuterated phenol-d₅) only rough estimates are given in the literature.^{82,83} For the PhOH-d₃ component in the presently studied mixture we find $\delta_{PhOH}/2\pi = 135 \pm 3$ kHz and $\eta_{PhOH} = 0.05 \pm 0.02$. Expectedly the anisotropy parameter given here is somewhat larger than for aliphatic deuterons, but well within the range expected for aromatic deuterons.^{84,86}

C. Oxygen spin relaxation and line widths

Spin-lattice relaxation times T_1 were monitored using saturation or inversion recovery techniques. For the measurement of the spin-spin relaxation times T_2 and if necessary for refocusing when recording T_1 , 90°–180° (Hahn-echo) and 90°–90° (solid-echo) pulse sequences were applied for the ¹⁷O and the ²H experiments, respectively.

To determine T_1 and T_2 for the equimolar mixture of PC and Ph¹⁷OH, we measured the longitudinal magnetization recovery $M_z(t)$ and the transverse magnetization decay $M_{xy}(t)$, respectively, in a wide temperature range. The corresponding magnetization curves were fitted using the stretched exponential functions

$$M_z(t) \propto \exp[-(t/T_1)^{\mu_1}] \quad (6)$$

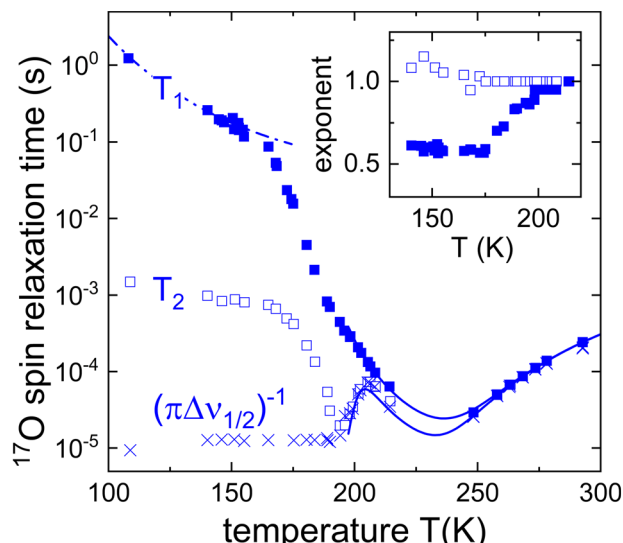


Fig. 5 Spin-lattice relaxation times T_1 (filled symbols) and spin-spin relaxation times T_2 (open symbols) of an equimolar mixture of Ph¹⁷OH and PC. The crosses (\times) refer to $T_2^* = 1/(\pi\Delta\nu_{1/2})$. Below 200 K, the T_2 times and the spectra were measured under continuous-wave proton decoupling. At higher temperatures decoupling was not applied. The inset depicts the Kohlrausch exponents corresponding to the spin relaxation times. The lines are fits using eqn (8), (13), and (14). The dash-dotted line reflects the calculated T_1 contribution that arises from the β -relaxation, see the Discussion section.

and

$$M_{xy}(t) \propto \exp[-(2t/T_2)^{\mu_2}], \quad (7)$$

respectively. Fig. 5 shows the fit results. Above about 250 K, we find that T_1 and T_2 agree and that the magnetization curves are essentially exponential ($\mu_1 \approx \mu_2 \approx 1$). Then, as mentioned, below about 250 K and down to about 215 K, spin relaxation becomes so fast that measurements could not be taken. From the inset of Fig. 5 we infer that the transversal dephasing remains essentially exponential down to the lowest temperatures.

For temperatures below 215 K, Fig. 5 shows that while T_2 exhibit a nonmonotonic behavior upon cooling, T_1 displays a monotonous increase which is somewhat more steep between 185 and 165 K and then tends to become flatter. The region of steeper increase is reminiscent of similar effects observed using ²H NMR, where the analogous phenomenon was assigned to a crossover from rate averaging over a distribution of T_1 times at high temperatures to time averaging at low temperatures.⁸⁵ In other words, at high T ergodicity restoring effects take place on the T_1 scale and lead to an exponential spin-lattice relaxation; conversely, at lower T where this effect ceases to be operative on the scale of T_1 , spin lattice relaxation is expected to proceed in a *nonexponential* fashion.⁸⁶ Indeed, the exponent $\mu_1(T)$ for PC:Ph¹⁷OH conforms exactly to this behavior, see the inset of Fig. 5.

Regarding the applicability of the interpretation just given for $\mu_1(T)$, one should keep in mind that for halfinteger quadrupolar spins, in principle, spin-lattice relaxation proceeds in a *nonexponential* fashion.^{87–89} Yet, even in the slow-motion



regime the expected deviations from exponentiality are small and can roughly be parameterized using a Kohlrausch exponent $\mu_1 = 0.95$, see Appendix A. Disregarding these minor deviations, the spin-lattice relaxation rate can be expressed as^{90,91}

$$\frac{1}{T_1} = K_0 [2J(\omega_0) + 8J(2\omega_0)] \quad (8)$$

with⁹²

$$K_0 = \frac{12\pi^2}{400} \frac{2I+3}{I^2(2I-1)} \Delta C_Q^2 \left(1 + \frac{1}{3}\eta_0^2\right). \quad (9)$$

For $I = 5/2$, the right hand side of eqn (9) reduces to a numerical factor $6\pi^2/625$ multiplied by the fluctuating part of the quadrupolar product $\Delta C_Q^2 \left(1 + \frac{1}{3}\eta_0^2\right)$. For $I = 1$ the numerical factor is $K_Q = 3\pi^2/20$.

In eqn (8) the spectral density function $J(\omega)$ depends on the molecular correlation time. For typical glassforming liquids, a distribution of correlation times needs to be taken into account. For the structural relaxation, we will use a Cole–Davidson spectral density⁹³

$$J_{CD}(\omega) = \frac{\sin[\gamma \arctan(\omega\tau_c)]}{\omega(1 + \omega^2\tau_c^2)^{\gamma/2}}. \quad (10)$$

From this expression, for $\gamma = 1$ the classical Bloembergen–Purcell–Pound form $J_{BPP}(\omega) = \tau_c/[1 + (\omega\tau_c)^2]$ is recovered.

In the regime of fast and intermediate motion, $\omega\tau_c \lesssim 1$, the spin–spin relaxation is “nearly exponential”,⁹⁴ as well so that one can define a rate

$$\frac{1}{T_2} = K_0 [3J(0) + 5J(\omega_0) + 2J(2\omega_0)]. \quad (11)$$

Under the condition of extreme narrowing, *i.e.*, for $\omega\tau_c \ll 1$, the terms in the square brackets of eqn (8) and (11) reduce to $10J(0)$ so that here

$$\frac{1}{T_1} = \frac{1}{T_2} = \frac{12}{125} \pi^2 C_Q^2 \left(1 + \frac{1}{3}\eta_0^2\right) J(0) \quad (12)$$

results. For $\omega \rightarrow 0$, eqn (10) yields $J(0) = \gamma\tau_c$. Hence, with the quadrupolar product known, for high temperatures eqn (12) provides a simple means of extracting molecular correlation times.

Outside the intermediate motion regime, where in the presence of a strong quadrupolar coupling essentially only the central transition is excited, the first-order quadrupolar contribution to the effective transverse relaxation rate is⁹⁵

$$\frac{1}{T_{2c}^{(Q1)}} = 5K_0 [J(\omega_L) + J(2\beta\omega_L)]. \quad (13)$$

For $I = 5/2$ the coefficient $\beta = \sqrt{2/\left[2/\left[I(I+1) - \frac{7}{4}\right]\right]}$ is ≈ 0.53 .

As τ_c gets longer towards the μ s scale, also T_2 tends to become longer so that for the transverse relaxation eventually the second-order quadrupolar contribution^{96–98}

$$\frac{1}{T_{2c}^{(Q2)}} = \frac{298}{875} \left(\frac{3}{16} \frac{I(I+1) - \frac{3}{4}}{I^2(2I-1)^2} \right)^2 \frac{4\pi^2 \Delta C_Q^4 \left(1 + \frac{1}{3}\eta_0^2\right)^2}{\nu_0^2} J(0) \quad (14)$$

will prevail. Eqn (14) has been derived under the assumption that the central-transition line displays essentially a Lorentzian shape.⁹¹ This condition becomes violated as the molecular correlation times exceed $\tau_{exc} = 1/(2\pi M_2^{1/2})$, where M_2 denotes the central second moment of the second-order broadened central-transition line. Using the approach set out in ref. 91, for the present parameters τ_{exc} is estimated to be about 16 μ s.

Returning to the line width, in the range in which $\tau_c < \tau_{exc}$ the overall width is expected to be given by

$$\Delta\nu_{1/2}^{\text{total}} = \frac{1}{\pi T_{2c}^{(Q1)}} + \frac{1}{\pi T_{2c}^{(Q2)}}. \quad (15)$$

Owing to the differing τ_c -dependences of the two contributions appearing in eqn (15), a minimum line width (hence maximum overall dephasing time) arises. Assuming that the spectral densities are of the Bloembergen–Purcell–Pound form ($\gamma = 1$), by inserting eqn (13) and (14) into eqn (15), a minimum line width of ref. 91

$$(\Delta\nu_{1/2}^{\text{total}})_{\min} \approx \frac{13.5\pi}{\nu_0^2} \left(\frac{C_Q \sqrt{1 + \eta_0^2/3}}{2I(2I-1)} \right)^3 \quad (16)$$

has been predicted that appears when the correlation time is

$$\tau_{c,T_{2\max}} \approx \frac{2.8}{C_Q \sqrt{1 + \eta_0^2/3}}. \quad (17)$$

According to ref. 99 a local minimum in T_2 (corresponding to a local maximum line width) will appear at a temperature at which the motional correlation time is $\tau_{c,T_{2\min}} \approx 1/(2.56M_2^{1/2}) \approx 40 \mu$ s. The corresponding time constant is included in the Arrhenius plot that is presented in Section 4, below.

Using the theoretical framework outlined in the present section, for $T \geq 200$ K we fitted eqn (8) to the spin-lattice relaxation times with the quadrupolar coupling set to $C_Q = 8.6$ MHz, and the asymmetry parameter to $\eta_0 = 0.7$ and assumed that the time constants follow the Vogel–Fulcher law that is displayed in the Arrhenius plot in Section 4, below. From the fit curve in Fig. 5, we can read out that the minimal oxygen T_1 relaxation time that would be expected to appear at 237 K. Taking into account the underlying time constants derived from the T_1 values together with the estimated Cole–Davidson parameter $\gamma_{CD} = 0.36 \pm 0.04$ as well as C_Q and η_0 as given above, we calculated the T_2 values according to eqn (13) and (14) and present the results in Fig. 5 as well.

For the current parameters (but with $\gamma = 1$), eqn (16) and (17) would yield 1.4 kHz and 0.3 μ s, respectively. If $\gamma < 1$, then

eqn (15) has to be analyzed numerically (see also Fig. S3 in the ESI†) and for the current γ exponent we find $(\Delta\nu_{1/2}^{\text{total}})_{\min} = 5.4$ kHz and $\tau_{c,T_{2\max}} = 0.6$ μ s.

The line widths $\Delta\nu_{1/2}(T)$ from the experimentally recorded spectra such as those shown in Fig. 4 are included in Fig. 5 in the form of effective spin–spin relaxation times, $T_2^* = 1/(\pi\Delta\nu_{1/2})$. From Fig. 5 one recognizes that T_2^* (which lacks the refocusing effects inherent in the present acquisition of T_2) agrees with T_2 down to about 190 K. Upon further cooling, $\Delta\nu_{1/2}(T)$ approaches its rigid-lattice value and thus becomes temperature independent.

D. Deuteron spin relaxation and stimulated echoes

Like for the ^{17}O labeled mixture, also for PC-d₃:PhOH and for PC:PhOH-d₃, longitudinal and transverse magnetization curves were recorded and analyzed using eqn (6) and (7), respectively. The corresponding spin relaxation times are presented in Fig. 6 (together with the data for PC:Ph ^{17}OH) for the two differently ^2H labeled equimolar mixtures. The Kohlrausch exponents are given as Fig. S4 in the ESI.† The ^2H spin–lattice relaxation times display temperature dependent patterns that are similar to those seen for ^{17}O . However, due to the relatively small quadrupolar ^2H couplings, for the deuteron probes the T_1 minima can be fully resolved. Since PC-d₃ and PhOH-d₃ show somewhat different quadrupolar parameters, see Section 3B, the two axes in Fig. 6 were shifted to achieve a good overlap in the high-temperature regime. This way any remaining differences in the (scaled) spin relaxation times should reflect differences in the temperature dependence or distribution width of the correlation times.

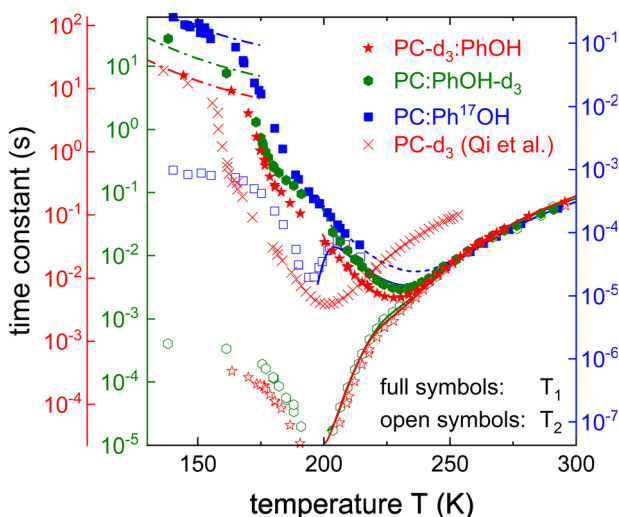


Fig. 6 Spin–lattice and spin–spin relaxation times of PC:PhOH-d₃, PC:PhOH-d₃, and PC:Ph ^{17}OH . For neat PC-d₃ T_1 was measured at a slightly lower Larmor frequency (40.2 MHz).⁵⁶ The y-axes are shifted so that the high-temperature flanks of all spin relaxation times superimpose. Thus, the axes for the deuterated substances differ by a factor 2.26, those for phenol in PC:PhOH-d₃ and in PC:Ph ^{17}OH by a factor of 235. The dashed lines (barely visible due to excellent agreement with the data) are fits to T_1 using eqn (8) for $T \geq 200$ K. The solid lines describing the ^2H T_2 times were calculated according to eqn (11). The dash-dotted lines reflect the calculated T_1 contribution arising from the β -relaxation, see Section 4, below.

The difference in the distribution width is nicely brought out in a plot of $1/T_1$ vs. $1/T_2$,¹⁰⁰ see Fig. 7. In this representation, the limiting slopes refer directly to those of the corresponding frequency dependent NMR susceptibility. For instance assuming a Cole–Davidson form, a power law with an exponent +1 will be observed on the left flank and on the right flank the exponent will be $-\gamma$. Fig. 7 shows that for PC-d₃:PhOH one has $\gamma = 0.42$ and for PC:PhOH-d₃ one has $\gamma = 0.51$. This means that in the mixture, PC senses a somewhat broader distribution of correlation times than PhOH.

Using these Cole–Davidson exponents γ (and $\eta_D = 0$), eqn (8) was fitted to the experimentally determined longitudinal relaxation times for $T \geq 200$ K. Fig. 6 shows that good agreement is achieved when setting the quadrupolar coupling to $\Delta C_Q = 144$ kHz for PC-d₃:PhOH and to $\Delta C_Q = 180$ kHz for PC:PhOH-d₃. From the Vogel–Fulcher laws underlying these fits, see the ESI,† and using eqn (11), we calculated the solid T_2 lines shown in Fig. 6 which describe the transverse relaxation times very well down to the T_1 minimum.

To assess the molecular correlation times also the procedure outlined in ref. 56 is often applied: Here, not only the fluctuating part of the quadrupolar coupling, but also the smallest measured longitudinal relaxation time ($T_{1,\min}$) plays an important role. For PC:PhOH-d₃ one has $T_{1,\min} = 3.0$ ms at 230 ± 3 K and $T_{1,\min} = 4.9$ ms for PC-d₃:PhOH at 228 ± 3 K. The correlation times at the T_1 minimum depend slightly on the exponent γ according to the empirical expression, $\omega_D \langle \tau_c(\gamma) \rangle \approx 0.425 + 0.467\gamma - 0.548\gamma^2 + 0.385\gamma^3 - 0.113\gamma^4$ that is applicable if T_1 is of the form given by eqn (8).¹⁰¹ For the given γ exponents this expression yields 1.95 ns for the PhOH-d₃ component and 1.8 ns the PC-d₃ component. Thus despite the slightly different maximum positions apparent from Fig. 7 the resulting time scale agree within experimental uncertainty.

Using spin relaxometry the molecular correlation function is probed most efficiently on the nanosecond scale (T_1) or on the microsecond scale (T_2). To monitor dynamical process directly on millisecond (and longer) time scales, deuteron stimulated-echo

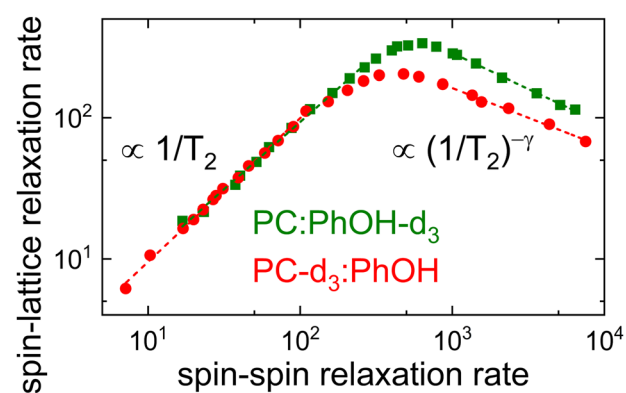


Fig. 7 In this double logarithmic plot of $1/T_1$ vs. $1/T_2$ the limiting behavior of a Cole–Davidson susceptibility, i.e., power laws with slopes of +1 and $-\gamma$ can be inferred from the dashed lines. For the equimolar mixture of PC and PhOH one recognizes that the PC component senses a somewhat broader distribution of time constants than the PhOH component does.

spectroscopy can be exploited to measure the (rotational) correlation functions of the quadrupolarly perturbed frequency^{81,102}

$$\omega_Q(t) = \pm \frac{1}{2} \delta_Q (3 \cos^2 \theta - 1 - \eta_Q \sin^2 \theta \cos 2\phi). \quad (18)$$

In eqn (18) the (in general time dependent) angles θ and ϕ describe the orientation of the electrical field gradient tensor that characterizes the molecular CD bonds with respect to the externally applied magnetic field in the usual fashion. The anisotropy parameter δ_Q and the small asymmetry parameter η_Q can be identified with those given in Section 3B for PC-d₃:PhOH and PC:PhOH-d₃.

Suitably phase cycled stimulated-echo pulse sequences allow one to record the two-time orientational correlation function^{81,102}

$$F_2(t_p, t_m) = \langle \cos[\omega_Q(0)t_p] \cos[\omega_Q(t_m)t_p] \rangle. \quad (19)$$

Here, the $\langle \dots \rangle$ brackets indicates an average over all deuteron spins. An analysis of $F_2(t_p, t_m)$ as a function of the mixing time t_m yields access to the molecular correlation times. Geometrical aspects of the molecular motion can be mapped out – *i.e.*, the angular sensitivity of the experiment can be modulated – by varying the evolution time t_p .^{102,103}

In Fig. 8 we show experimentally determined $F_2(t_p, t_m)$ curves for the two differently deuteron labeled PC:PhOH mixtures as a function of temperature for a fixed evolution time. For both samples one recognizes that upon cooling the decay of the stimulated-echo amplitude shifts to longer times and becomes more stretched. For a quantitative analysis of these data we

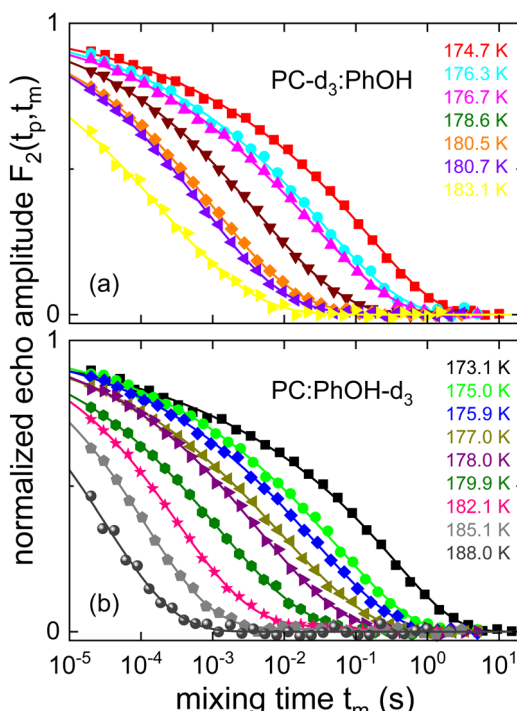


Fig. 8 Stimulated cos-cos echo functions measured for (a) PC-d₃:PhOH and (b) PC:PhOH-d₃. A three-pulse sequence with an evolution time of $t_p = 20 \mu\text{s}$ was used. The lines are fits using eqn (20).

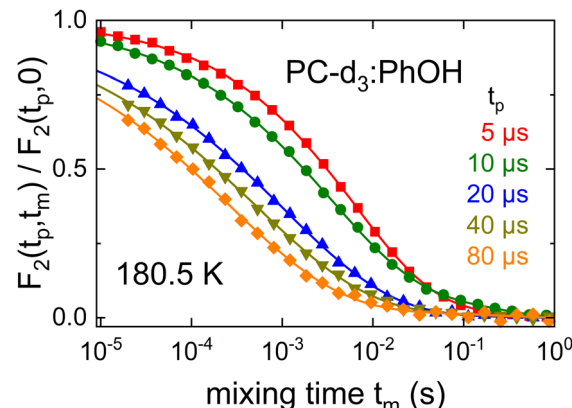


Fig. 9 Evolution time dependent stimulated cos-cos echo functions recorded for PC-d₃:PhOH. The lines represent fits using eqn (20). For $t_p \leq 10 \mu\text{s}$ a four-pulse sequence was utilized.

took the independently measured spin-lattice relaxation times into account and employed the stretched exponential function

$$F_2(t_p, t_m) = A_1 + A_2 \exp[-(t_m/\tau_c)^{\beta_K}]. \quad (20)$$

Here, $A_1 + A_2$ and A_1 denote the limiting values of the F_2 functions for short and long mixing times t_m , respectively. The Kohlrausch exponent β_K quantifies the deviations from an exponential decay, and, like before, τ_c designates the reorientational correlation time. The fits to the data shown in Fig. 8 reveal that β_K increases from 0.25 to 0.34 when T increases from 175 to 181 K.

While temperature dependent data are presented in Fig. 8 for a fixed evolution time, Fig. 9 shows stimulated-echo decays for PC-d₃:PhOH taken at $T = 180.5$ K for a range of evolution times. From least-squares fits based on eqn (20), we find that the correlation time $\tau_c(t_p)$ strongly depends on t_p , see Fig. 10.

This figure also includes data for the complementarily deuteron labeled sample, PC:PhOH-d₃. One recognizes that

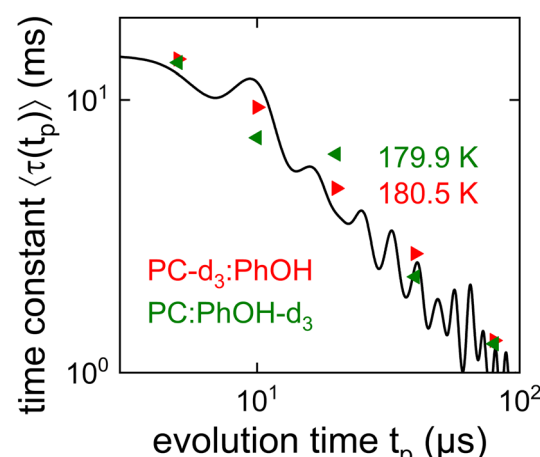


Fig. 10 Evolution time dependent cos-cos time constants as measured for PC-d₃:PhOH and for PC:PhOH-d₃. The solid line is adapted from ref. 104 and reflects simulations based on a motional model featuring a bimodal jump-angle distribution.

both mixtures exhibit very similar $\tau_c(t_p)$ dependences. The solid line in Fig. 10 reflects simulations based on a simple model which involves a bimodal distribution of small-angle (2°) and large-angle (30°) jumps, where on average the former are 50 times more frequent. Simulations with this reorientational scenario, after adjusting the scale for $\tau_c(t_p) \rightarrow 0$, were previously found suitable also for other glass formers including PC-d₃.^{56,104}

4. Discussion

Based on the results from the various experimental methods, Section 4A collects the evidence for the occurrence of a Debye process in PC:PhOH. Insights regarding the molecular conformation and arrangement as obtained from NMR and from dielectric spectroscopy are discussed in Sections 4B1 and 4B2, respectively. Then, Section 4C deals with the implications of these considerations regarding the supramolecular association in PC:PhOH. Finally, Section 4D is concerned with the finding of a clearly discernible secondary relaxation in these mixtures.

In order to facilitate the further discussion, the temperature dependent relaxation times of PC:PhOH that we determined in the previous sections are collected in the Arrhenius plot shown as Fig. 11. This includes the time constants obtained from dielectric and mechanical spectroscopy, from the spin-relaxation times, as well as from stimulated-echo spectroscopy. Except for the data relating to the β process, overall and typical for glass forming liquids, a curved temperature dependence is obvious from this representation. Such a dependence can be described using the Vogel–Fulcher law

$$\tau = \tau_0 \exp[B/(T - T_0)], \quad (21)$$

as well as other expressions.¹⁰⁵ Indeed, eqn (21) with $T_0 = 140 \pm 3$ K, $B = 1300 \pm 100$ K, and $\tau_0 = 10^{-13.2 \pm 0.4}$ s provides a good fit to the dielectric data of PC:PhOH. The Vogel–Fulcher law yields $T_{g,e} = T(\tau_e = 100$ s) ≈ 174 K as the dielectrically detected glass transition temperature. Vogel–Fulcher fits for the spin-relaxation based correlation times, and a fit including also the NMR based time constants that are longer than 1 μ s are included in Fig. 11 as well. The related fit parameters can be found in Table S1 of the ESI.[†]

A. Evidence for the Debye process in PC:PhOH

The pattern displayed in Fig. 11 is reminiscent of that observed for aliphatic monohydroxy alcohols and like for those hydrogen bonded liquids it provides evidence for the existence of a Debye process also in PC:PhOH.^{4,5}

(i) The relaxation times originating from the dominant dielectric loss peaks (*cf.* the dash-dotted line in Fig. 11) are longer than those from the NMR based time scales.

(ii) The correlation times detected using ¹⁷O NMR at the hydroxyl group are intermediate.

(iii) Statement (ii) is corroborated not only by the deuteron spin relaxation data but by the stimulated-echo results as well (also in the $t_p \rightarrow 0$ limit, *cf.* the solid line in Fig. 11).

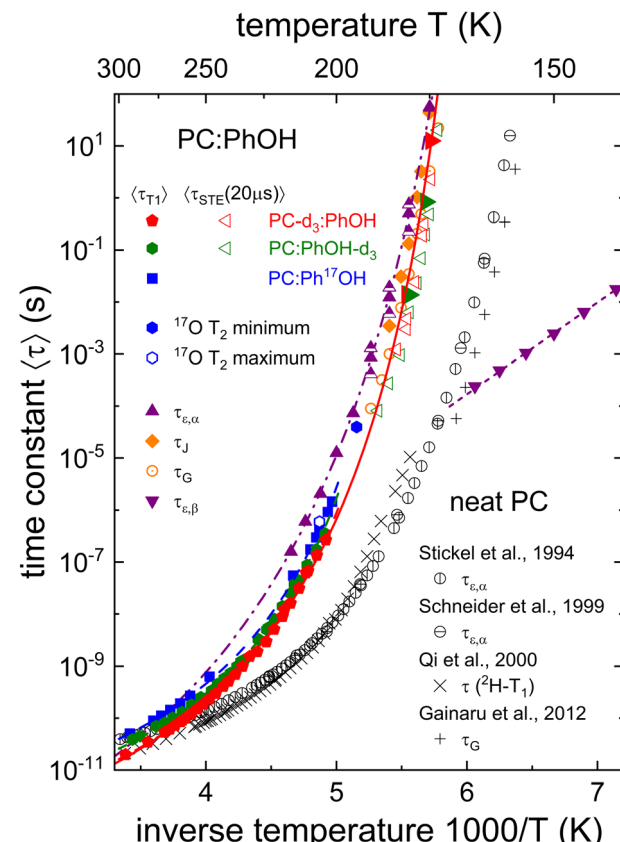


Fig. 11 Dielectric, mechanical, and NMR-based relaxation times of PC:PhOH with $x_{\text{PhOH}} = 0.5$. Half-filled triangles mark the results from the decomposition of the dielectric spectra as illustrated in Fig. 2. The τ_G time scales represented without a central dot are determined using frequency-temperature superposition, the others from observed shear loss peaks. Most stimulated-echo (STE) decays were recorded for evolution times of 20 μ s, those represented by filled triangles pointing to the right for $t_p = 5$ μ s. Lines are fits using Vogel–Fulcher or Arrhenius laws. The dashed lines reflect the Vogel–Fulcher fits used to describe the T_1 curves. Their parameterization as well as those of the dash-dotted and solid lines is given in the text or as ESI.[†] For comparison with the PC:PhOH data, this plot features dielectric, rheological, as well as NMR-based time scales as determined for neat PC taken from ref. 51–53 and 55.

(iv) The shear rheological results display a small, yet significant viscosity enhancement.

(v) Upon approaching the glass transition temperature, the timescales from the Debye-like and the structural modes approach each other.

In order to appreciate statement (i) better a few comments are warranted: In general, the observed timescale separation seen in Fig. 11 is much larger than the maximum factor of three that is expected when dealing with different Legendre polynomial ranks, *i.e.*, when comparing dielectric and NMR time scales.¹⁰³ Following previous work,^{17–19} the identification of a Debye process can be based on the spectral decomposition illustrated in Fig. 2, see also Fig. S1 in the ESI.[†] Since, like for other examples,¹⁰⁶ such a decomposition is not immediately obvious from the raw spectra, the application of additional experimental methods is required to allow for an unambiguous detection of a Debye process.

Despite decade-long beliefs to the contrary, that the Debye process in monohydroxy alcohols manifests solely in dielectric spectroscopy,⁵ NMR experiments provided the first evidence that this notion needs substantial revision: the Debye relaxation leaves its mark in an OH-related timescale that is intermediate between those of the Debye and of the structural process.⁴ Further confirmed by findings in pure and diluted monohydroxy alcohols,^{46,107} the observations summarized in items (ii) and (iii) underscore that PC:PhOH closely resembles the pattern revealed by well-established Debye liquids. In PC:PhOH the separation of the OH-related time scale from the other modes is, however, less pronounced than it is in liquids that display a *strong* Debye process.

In line with these findings, as indicated in item (iv), the rheological investigation of PC:PhOH (see Fig. 3) reveals a viscosity enhancement, see eqn (4), relative to that of nonassociating liquids and thus the existence of a supramolecular mode. The spectral comparison with a nonassociating liquid shows that this additional contribution is readily visible, but relatively small in amplitude. Consequently, the quantitative assessment of the viscosity enhancement, carried out in Section 3A, yields a scaled zero-shear viscosity z_0 that is only slightly larger than 1. By comparison with other supramolecular liquids,⁷⁷ this means that in PC:PhOH the underlying rheological mode is relatively weak, a statement that corroborates the finding from the decomposition of the dielectric spectra, *cf.* Fig. 2. Further vindicating the existence of a faint Debye-like process, the time scales determined from the shear compliance (often taken as a measure for the structural relaxation time) almost coincide with the shorter time scale inferred from the decomposition of the dielectric loss spectra.

As item (v) highlights, another observation from Fig. 11 agrees with that for many aliphatic monohydroxy alcohols:²⁰ The Debye and the structural relaxation timescales approach each other upon cooling down to the glass transition temperature.²⁰ Taken together, the combination of the results from the various methods all indicate a weak Debye-like process to be present in PC:PhOH.

B. Molecular conformation and arrangement

1. Insights from oxygen magnetic resonance. Regarding the spectral NMR parameters that we experimentally determined in Section 2B, see in particular Fig. 4, it is useful to point out that these can be compared with results obtained from quantum chemical calculations³³ with reference to work for crystalline phenol.¹⁰⁸ After averaging over all lattice sites, the latter work reports $C_Q = -8.686$ MHz, $\eta_O = 0.84$, and $\delta_{\text{iso}} = 81.6$ ppm. For phenol in a glycerol/water mixture $|C_Q| = 8.3$ MHz, $\eta_O = 0.95$, and $\delta_{\text{iso}} = 100$ ppm was found.³³ Note that experiments typically cannot determine the sign of C_Q . Furthermore, it is interesting to note that these parameters are similar to those found for the phenolic oxygen in salicylic acid ($C_Q = -8.3$ MHz, $\eta_O = 0.60$, and $\delta_{\text{iso}} = 89$ ppm).¹⁰⁹ Thus, the NMR parameters that are determined for PhOH in the present work are in harmony with expectations based on the cited literature.

In order to further scrutinize the quadrupolar parameters at the oxygen site, analogous to previous work,^{110,111} for crystalline

phenol,^{28,29} we carried out quantum chemical calculations using the CASTEP 19.11 code.¹¹² These involved the PBE functional¹¹³ as well as a cut-off energy of 800 eV and employed a Monkhorst–Pack grid with a spacing of 0.03 \AA^{-1} .¹¹⁴ Further details regarding the density functional calculations are given in the ESI†.

For the ambient-pressure phase in which the planar PhOH molecules form three-fold hydrogen bonded helical chains²⁸ we find $C_Q = -8.67$ MHz and $\eta_O = 0.82$, thereby confirming the coupling parameters from earlier work³³ (see also the ESI†). In the high-pressure phase which is stable at 0.16 GPa,²⁹ the PhOH molecules are nonplanar with a dihedral CCOH angle of almost 20° . For this phase in which the PhOH molecules are arranged in hydrogen bonded ribbons, our calculations reveal significantly larger quadrupolar parameters ($C_Q = -9.0$ MHz and $\eta_O = 0.96$). Thus, the comparison with the experimentally determined parameters, see Fig. 4, suggests that in the present binary liquid the PhOH molecules are characterized by a planar structure.

2. Insights from dielectric spectroscopy. In order to examine whether, based now on dielectric spectroscopy, the PhOH molecules form ring-like (or rather chain-like) associates, it is instructive to analyze the dielectric results in terms of the Kirkwood factor¹¹⁵

$$g_K = \frac{9\epsilon_0 k_B T (\epsilon_s - \epsilon_\infty)(2\epsilon_s + \epsilon_\infty)}{\tilde{n}\mu^2} \quad (22)$$

For ring-like or antiparallel dipolar arrangements one typically finds $g_K < 1$, while for chain-like or parallel dipolar arrangements $g_K > 1$ results. For pure PC the dispersion strength $\Delta\epsilon = \epsilon_s - \epsilon_\infty$ is almost temperature independent, implying that g_K decreases with T ,⁵³ and close to the glass transition, g_K has been reported to be near or below 0.6.⁵⁴

For PC the magnitude of the electrical dipole moment ($\mu_{\text{PC}} \approx 5$ D) is generally agreed upon.⁶⁴ However, the dipole moments reported for PhOH in the gas phase,^{37,38,116,117} in the low-viscosity liquid state,¹¹⁸ in aqueous^{119,120} and other solutions,¹²¹ as well as for the crystal at ambient²⁸ and high pressures²⁹ range from 1.2 D in the gas phase, to about 1.5...2.5 D in solution, and 3.5 D in the ambient pressure crystal. Interestingly, the dipole moment for geometry-optimized PhOH trimers was found to be even larger (3.75 D).³⁰

In view of the large uncertainty regarding the magnitude of μ_{PhOH} in liquid mixtures that contain PhOH,^{119–121} in the following analysis we can only expect to extract the product of g_K and μ^2 in a reliable way. This quantity can be estimated from⁶⁹

$$g_K \mu_0^2(x_{\text{PhOH}}) = \frac{3\epsilon_0 k_B T}{\tilde{n}(x_{\text{PhOH}})F} \Delta\epsilon(x_{\text{PhOH}}). \quad (23)$$

Here, $\tilde{n}(x_{\text{PhOH}}) \approx (6.8x_{\text{PhOH}} + 7.1x_{\text{PC}}) \times 10^{27} \text{ m}^{-3}$ is the number density of the binary liquid and x_{PhOH} and $x_{\text{PC}} = 1 - x_{\text{PhOH}}$ designate the mole fractions of the two components. Then, the effective dipole moment in the mixture is given by^{122–124}

$$\mu_0 = \sqrt{x_{\text{PhOH}}\mu_{\text{PhOH}}^2 + x_{\text{PC}}\mu_{\text{PC}}^2}. \quad (24)$$



Note that rather than the mole fractions, also volume fractions¹²⁵ have occasionally been used to assess $\mu_0(x_{\text{PhOH}})$. Furthermore, the local-field correction factor⁶⁹

$$F = \frac{\epsilon_s(\epsilon_\infty + 2)^2}{3(\epsilon_\infty + 2\epsilon_s)} \quad (25)$$

appearing in eqn (23) accounts for the difference in the macroscopic and the microscopic electrical fields. With $\Delta\epsilon$ as given at $T = 180$ K in Fig. 2 for the samples with $x_{\text{PhOH}} \leq 0.5$ and taking $\epsilon_\infty = 3$ (leading to $F = 4.1$) as well as $\mu_{\text{PC}} = 4.94$ D,⁶³ the solid line in that figure is calculated.

Now, based on this result, the determination of g_K rests on assumptions regarding μ_{PhOH} and *vice versa*. For instance, taking $\mu_{\text{PhOH}} = 1.5$ or 2.5 D, the minimum or maximum values, respectively, reported for phenol in various solutions,^{119–121} one arrives at $0.66 \leq g_K \leq 0.81$ or at $0.66 \leq g_K \leq 0.70$. Tentatively assuming a chainlike association, characterized by, *e.g.*, $g_K = 2$, which would however be at variance with the findings for neat PC⁵⁴ even at high temperatures,¹²⁶ one would have $\mu_{\text{PhOH}} \approx 1.7$ D and (also at variance with the experimental value) $\mu_{\text{PC}} \approx 2.8$ D. These considerations show that in the present mixtures a chain-like arrangement of the PhOH molecules can be ruled out.

C. Hydrogen bonding and dilution effects in aromatic and aliphatic Debye liquids

As discussed above, the observed ¹⁷O quadrupolar coupling constant of phenol as well as the Kirkwood factors which characterize PC:PhOH favor an interpretation in terms of the existence of ring-like molecular associates or dipole arrangements with antiparallel orientations. Like for monohydroxy alcohols displaying ring-like supramolecular associations,⁶ it is not surprising that for PC:PhOH only a relatively weak hydrogen-bond mediated viscosity enhancement is found.

For phenyl propanols^{12–14} it was emphasized that the tendency to form molecular associates transiently⁴ held together by O–H···O bonds is impeded by the propensity of these molecules to form O–H···π¹⁵ and π–π configurations. Additionally, it has to be taken into account that PC cannot be involved in the formation of supramolecular structures so that this van der Waals glass former merely acts as a diluent in the PC:PhOH mixture. Hence, we anticipate that similar to the pure ambient-pressure liquid,³⁰ also when mixed with PC, the PhOH molecules predominantly form three-membered O–H···O bonded clusters and ring-like structures. From X-ray diffraction experiments, similar bonding motifs were in fact detected for aliphatic monohydroxy alcohols displaying small Debye processes, while for alcohols featuring large Debye processes chain-like clustering prevails.^{6,47,127} To check which supramolecular bonding structures are present in the PC:PhOH mixtures, in the future these should be scrutinized using suitable scattering experiments.

D. Secondary relaxation

So far, the discussion focused on the structural relaxation or on processes which are slower than that. As can be seen in Fig. 11 PC:PhOH exhibits a β relaxation, *i.e.*, a faster process as well. At first sight, this is remarkable since without long-time aging,⁷²

neat PC does not show a discernible β process. For aromatic glass formers it is not clear what to expect: Substances such as *meta*-fluoroaniline and toluene do exhibit a strong β relaxation while others, such as 2-picoline, do not.^{128,129}

The Johari–Goldstein relaxation times that we obtain for PC:PhOH follow an Arrhenius law

$$\tau = \tau_{0,\beta} \exp(E_\beta/k_B T). \quad (26)$$

From the fit shown as a dashed line in Fig. 11, an activation energy of $E_\beta = 33.1 \pm 1.0$ kJ mol^{−1} and a pre-exponential factor $\tau_{0,\beta} = 10^{-13.3 \pm 0.2}$ s can be inferred. The resulting $E_\beta/(k_B T_g)$ ratio of 22.8 is typical for many glass formers.⁷³

We will now demonstrate that the signature of the β process can be found also from NMR, *i.e.*, in the low-temperature T_1 data. To see this, for $T < T_g$ where the α process ceases to lead to significant spin–lattice relaxation and where thus the β process prevails, based on eqn (8) and (10) we calculated T_1 . As input parameters, we used τ_β from dielectric spectroscopy in conjunction with a Cole–Cole spectral density with an exponent $\alpha_{\text{CC}} = 0.2$ so that the only free parameter is the fluctuating part of the coupling constants. To obtain the dash-dotted lines in Fig. 6 (²H data) that reflect these calculation we used $\sqrt{K_Q} = 15$ kHz. Analogously, for the ¹⁷O data we obtain $\sqrt{K_Q} = 140$ kHz. Reduced quadrupolar couplings (here: 5 to 7% of the full ones) are in fact expected in the framework of models for secondary relaxations: Here, the molecular segments perform motions in an angularly restricted range, typically pictured to occur within a cone.^{86,130} From the cited works cone semiangles κ in the 5 to 20° range were found for various glass formers. Indeed, from the relaxation strengths $\Delta\epsilon_\beta$ and $\Delta\epsilon$ referring to the β and the dominant relaxation, respectively, of PC:PhOH using the empirical relationship^{131,132} $\sin^2\kappa = \Delta\epsilon_\beta/\Delta\epsilon$ it is possible to assess κ on the basis of the dielectric data as well. Assuming that $\Delta\epsilon_\alpha$ is constant not only above T_g , see Fig. 1, but also below T_g , for $x_{\text{PhOH}} = 0.5$ for between 140 to 170 K we find $\kappa \approx 6$ to 8° (or 8 to 11° if the relaxation strength relating only to the α-type contribution is taken, *cf.* Fig. 2).

The observation of a β process in binary systems in which one of the glassformers does not show this phenomenon in its neat form is not unprecedented: Mixtures of toluene and 2-picoline do indeed feature a secondary relaxation which was reported to strongly resemble that found for neat glassformers¹²⁸ so that in ref. 128 it was concluded that “all molecular species contribute to the secondary relaxation of the mixtures”. The present results suggest that in PC:PhOH we encounter the same situation, *i.e.*, that also here both molecular constituents contribute to the secondary relaxation.

5. Conclusions

In this work, we studied the relaxation dynamics of phenol. Since in its pure form, PhOH cannot be brought into the amorphous form, even by quenching with large cooling rates, we studied solutions with PC, an excellent van der Waals glass former, mostly focusing on equimolar mixtures. To this end we



employed a large array of techniques including dielectric spectroscopy and shear rheology, NMR line shape analysis including quantum chemical calculations, spin relaxometry using ^2H and ^{17}O as nuclear probes, as well as deuteron stimulated-echo spectroscopy. The present combination of experimental techniques allowed us to probe the dynamics of glass forming PC:PhOH over a large temperature range. Furthermore, the applied deuteron editing enabled a selective detection of the spin relaxation and stimulated-echo behavior of the two molecular components that form PC:PhOH: we find that within the mixture, the two components display essentially the same structural and secondary dynamics, demonstrating that PC is a well suited mixing partner for the present purpose. However, since PC, unlike PhOH, is unable to sustain hydrogen bonds, the van der Waals glass former cannot be directly involved in the formation of supramolecular structures that give rise to the Debye-like process. For PC:PhOH the latter is identified on the basis of the following findings: by examining the behavior at the hydroxyl site using ^{17}O NMR, we find that this moiety displays slightly longer correlation times than inferred from deuteron NMR for the PhOH ring, an observation that hints at a weak hydrogen-bond induced molecular association. Corroborating this finding, the dielectric relaxation times turn out to be somewhat longer than those determined using ^2H NMR, even after taking into account that the different techniques probe different Legendre orders of the underlying reorientational correlation function. ^{17}O NMR, a method that here probes the dynamics at phenol's hydroxyl site reveals an intermediate dynamic, akin to the situation in aliphatic mono-hydroxy alcohols. Analogous to those latter liquids, from our shear rheology study on PC:PhOH evidence for an, albeit weak viscosity enhancement is obtained. Thus, taking together the results from the diverse array of the present experiments and analyses we obtain evidence for the presence of a faint Debye-like process stemming from phenol. In this sense our studies reveal that this process not only occurs in phenyl alkanols with sizeable side chains, but also in a mixture of a nonassociating liquid with the simplest aromatic alcohol.

Conflicts of interest

There are no conflicts of interest to declare.

Appendix

Appendix A: Multi-exponential spin-lattice relaxation of $I = 5/2$ nuclei: selective and nonselective conditions

Generally, ^{17}O spin-lattice relaxation proceeds in a nonexponential fashion and the time dependent part of the multi-exponential magnetization recovery can be written as

$$M_z(t) \propto \sum_{k=1}^3 C_k \exp(-R_k t). \quad (27)$$

To obtain the relative amplitudes C_k (with $C_1 + C_2 + C_3 = 1$) and the recovery rates R_k , the Redfield equations have to be solved.¹³³ For spins with quantum numbers $I > 3/2$ this is not possible analytically. For $I = 5/2$ nuclei the numerical solution for the situation where all transitions are nonselectively excited was variously given for the complete range of time scales $\omega_L \tau_c$.⁸⁷⁻⁸⁹ However, for selective excitation and detection of only the central transition, we are not aware of a published solution.

Therefore, based on the eigenvalues and eigenvectors of the relaxation matrix derived from ref. 134, we solved eqn (2.4a) in conjunction with eqn (A20), both from that reference, and summarize our numerical results in Fig. 12. A close inspection shows that the relative amplitudes C_k calculated using this approach and marked as "nonselective" in Fig. 12(a) as well as the calculated rates, see Fig. 12(b), reproduce the previously published ones.⁸⁷⁻⁸⁹

While the rates do of course not depend on specific excitation and detection conditions, the relative amplitudes C_k do. This is seen in Fig. 12(a) where the amplitudes referring to selective and nonselective conditions are shown. In particular, for the selective case in the slow-motion regime where $\omega_L \tau_c \gg 1$, a fit of $M_z(t)$ using eqn (6) – after the relevant coefficients were plugged into eqn (27) – yields a Kohlrausch exponent $\mu_1 = 0.95$. In other words, under the conditions which are most relevant for the present work, the deviations from nonexponential spin-lattice relaxation are minor.

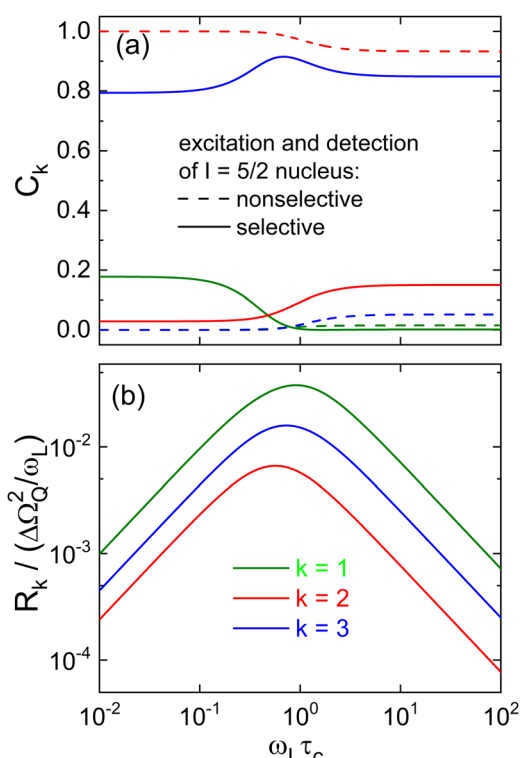


Fig. 12 (a) Amplitudes and (b) rates describing the longitudinal recovery of an $I = 5/2$ nucleus such as ^{17}O after selective as well as nonselective central-transition excitation and detection. The entire range from the fast-motion to the slow-motion regime is covered. Ω_Q denotes the quadrupolar product.



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