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Shear-Modulus Investigations of Monohydroxy Alcohols: Evidence for a Short-Chain-Polymer Rheological Response

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Liquids composed of small-molecule monohydroxy alcohols are demonstrated to display rheological behavior typical for oligomeric chains. This observation was made possible by rheological experiments in which more than seven decades in frequency and more than five decades on the mechanical modulus scale are covered. The singly hydrogen-bonded monohydroxy alcohols were chosen because they display significant, but surprisingly poorly understood effects of intermolecular association. Based on the present shear study, one can apply theoretical concepts of polymer science to understand the anomalous physical behavior of a wide range of hydrogen-bonded liquids.

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Rheological experiments have long been thought to allow for a clear-cut distinction between simple small-molecule liquids and macromolecular melts [1]. The dynamic viscosity of small-molecule fluids is typically governed by local flow processes reflecting short-ranged interactions. Conversely, in melts of chain molecules it is the covalently bonded backbone structure which governs the slowest relaxation of externally imposed mechanical stresses [2,3]. This leads to an increase of the overall mechanical stiffness, eventually giving rise to flow behavior typical of polymer melts. Supramolecular behavior can further be fortified by noncovalent interactions mediated by, for instance Coulombic forces, metal-ligand coordination, and hydrogen bonding, the latter, e.g., in the form of acid-base pairing [4–6]. Recently, Lou et al. studied multiply bonded donor acceptor motifs, found evidence for supramolecular polymeric liquid behavior based on nuclear magnetic resonance (NMR), dielectric relaxation spectroscopy, as well as rheological measurements [7], and compared their findings with those of the singly hydrogen-bonded monohydroxy alcohols (MAs). Albeit indications for supramolecular characteristics in MAs have been reported from NMR and dielectric studies [8], this behavior is in seeming contradiction with many data collected for these small-molecule liquids during the last five decades [9–12]. Based on shear mechanical measurements and by comparing with data of nonhydrogen-bonded small-molecule liquids the present Letter shows that MAs indeed do exhibit a rheological signal demonstrating short-chain polymerlike aggregation of MAs. Our work paves the way for their description in terms of concepts developed for transient (living) polymers.

The MAs studied in the present work are the octanols 4-methyl-3-heptanol (4M3H) and 2-ethyl-1-hexanol (2E1H) (both obtained from Sigma-Aldrich and used as received), well studied using various techniques; see, e.g., [12–16]. In Fig. 1 we present the complex shear modulus $G^\prime(\nu) = G^\prime(\nu) + iG^\prime\prime(\nu)$ of 4M3H, covering more than 7 decades in frequency (from $10^{-3}$ to $10^4$ Hz) using a piezoelectric shear modulus gauge (PSG) technique [17]. A single-peak structure in $G^\prime(\nu)$ is observed shifting through the frequency window, indicating the temperature dependence of the structural (or $\alpha$-) relaxation time originating from the local flow processes of this viscous liquid.

At first glance, the data look unsuspicuous, except that a kink appears on the low-frequency flank of the dominant peak, a feature reminiscent of a terminal mode reported to occur for a variety of short-chain polymers [18–20]. All spectra from Fig. 1 can be collapsed to a master curve, see Fig. 2(a), indicating that for 4M3H time temperature superposition is obeyed for the storage component and the loss component of $G^\prime(\nu)$ [21]. Here, we used the peak frequency $\nu_{\text{max}}$ identified with $\nu_\alpha$, as a scaling variable, i.e., as an effective shift factor. Figure 2(a) also includes shear data recorded using the rheometer ARES G2 (from TA Instruments) that provides mechanical excitation from 0.1 to 100 Hz [22]. Overall, the scaled frequency range extends over about 10 decades. For 2E1H, shear data from the rheometer and the PSG technique are combined yielding the master plot given as Fig. 2(b). Again, the $G^\prime(\nu)$ data exhibit excellent scaling behavior for all frequencies near or below the peak in $G^\prime(\nu)$. The deviations from time-temperature superposition visible at higher frequencies for 2E1H are caused by a secondary relaxation stemming from localized molecular motions [12]. The shapes of the master curves for the two MAs differ, pointing to differences in their supramolecular organization. As will be discussed below, these originate from differences in the molecular structures, sketched in Fig. 2.
illustrates that, unlike 4M3H, 2E1H features a hydroxyl group in a terminal position.

Since the focus here is on the spectral regime in which supramolecular relaxation may be expected and to facilitate detailed comparisons with the behavior of simple, non-polymeric liquids [23], in Figs. 3(a) and (b) we show MA data in the scaled frequency range near and below \( \nu_\alpha \). The shift factors used to scale the rheological data in Fig. 2 and Fig. 3 comply with the temperature dependent structural relaxation frequencies determined directly from the PSG measurements; see the Arrhenius plot, Fig. 4.

In Fig. 3 we compare the \( G'(\nu) \) results with those for propylene carbonate and several other nonassociating liquids [23]. Overall, the main peak of their rheological spectra displays a similar shape, but clear differences are recognized on the low-frequency side of the structural relaxation peak. While the simple liquids display a unique power law in \( G'(\nu) \), the shear modulus of 2E1H, and to some extent also of 4M3H, resembles normal-mode spectra of short-chain polymers [3,18]. Several polymerlike regimes can be distinguished in Fig. 3 from the frequency-dependent shear moduli on the low-frequency side written as \( G'_0(\nu) \propto \nu^\alpha \) and \( G''_0(\nu) \propto \nu^\beta \). Generally, the
The interpretation of our results in terms of end-to-end chains is akin to the situation in type A polymers for which along-the-contour electrical dipole moments add up to produce a pronounced dielectric normal mode [28]. Hence, it is instructive to compare the rheological results with dielectric data for the presently studied supramolecular polymers. In Fig. 3(c), dielectric loss spectra are presented, revealing that in addition to the structural relaxation a strong normal mode type of relaxation appears for 2E1H. In the context of MA this feature, due to its single-exponential appearance has been called dielectric Debye relaxation [29]. For 4M3H the low-frequency dielectric response is much weaker and may be affected by a ring-chain equilibrium [14], also a prominent theme in the context of supramolecular polymers [30]. Then, Fig. 3(c) demonstrates that in nonassociating nonpolymeric liquids, with propylene carbonate being an example, dielectric modes slower than the structural relaxation do not exist. This finding is fully compatible with the mechanical response of propylene carbonate and of other liquids devoid of intermolecular hydrogen bonds; cf. Fig. 3(b).

Using results from infrared, NMR, and dielectric techniques it was argued that the supramolecular structure in octanols develops an increased tendency to disintegrate thermally in the high-frequency regime [15,16]. Thus, in the sub-GHz range, detection of supramolecular features in MAs required the application of high-resolution ultrasonic techniques [31]. Brillouin scattering that probes mechanical response at even higher frequencies did not display a significant signature of supramolecular phenomena [11].

The contribution of the “normal mode rigidity” or “Debye rigidity” to the frequency-dependent shear viscosity of the MAs is obtained from \( \eta''(\nu) = G''(\nu)/(2\pi\nu) \). This relation follows from the definitions of the dynamic shear viscosity and modulus. For the \( \alpha \) relaxation we may write \( \eta_\alpha \approx G_\infty/(2\nu_\alpha) \). As the inset of Fig. 4 reveals, for 2E1H the viscosity increase \( \Delta\eta_\alpha/\eta \) or the rigidity enhancement \( \Delta G_D/G_\infty \) due to the supramolecular (Debye or normal mode) process amounts to a factor of \( \sim 10 \). However, the polymerlike effects show up at moduli about 3 orders of magnitude lower than \( G_\infty \), much below the detection limits of typical shear mechanical [12] and ultrasonic experiments [9]. The suprasegmental mechanical contribution observed for 2E1H is comparable to that of short-chain polymers [3], but has been detected at temperatures much lower than usually applied in studies of covalently bonded polymers. Furthermore, the slower the dielectric normal mode is with respect to the \( \alpha \) response, the larger also is the separation of the two modes in the mechanical spectra, and the lower will be the modulus at which the unambiguous \( G''(\nu) \propto \nu^\alpha \) or \( G''(\nu) \propto \nu^\beta \) low-frequency signature of the terminal chain mode shows up. Thus, on the basis of the current work, we predict that small-molecule liquids comprising a single intermolecular peptide-related bond, such as the secondary amides [32,33] for which so far only an intermediate power
law was resolved in mechanical studies [23], also do display end-to-end chain modes.

To summarize, we identified MA as short-chain polymeric liquids. In contrast to typical supramolecular polymers, in MAs the transient-chain association is sustained via only a single intermolecular hydrogen bond. As a consequence, the observation of a chain mode in MAs becomes possible at temperatures as low as 145 K and, thus, very much lower than usually considered in the study of polymeric liquids. Furthermore, our findings show that MAs and predictably many other chain-forming hydrogen-bonded or peptide-bonded liquids can be analyzed theoretically in terms of polymer concepts and possibly by suitably adapted approaches developed for related soft matter systems [34].

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[21] In the temperature range covered by the present study, near-infrared investigations indicate that the H bond concentration changes by only about 5%–6% for 2E1H [14] and 4M3H [15], thereby not affecting time-temperature superposeability in a significant way.
[25] For the Zimm model, see Fig. 9-8(b) on p. 190 and Eqs. (32) and (33) on p. 192 of Ref. [1].
[26] Here we used \( \rho = 0.833 \times g / cm^3 \), \( m = 2.16 \times 10^{-22} g \) or 130.2 g/mol, and \( \nu_R \approx 0.13 \) Hz at 160 K. According to the Zimm model the terminal rate is larger than \( \nu_R \), thus \( M_{eff} \) would turn out somewhat larger if calculated using that approach. In any case, the given estimates have to be considered with some caution, because neither the Zimm nor the Rouse model address aspects characteristic for transient hydrogen-bonded chains such as exchange processes between bonded and nonbonded molecules.