Molecular interpretation of the non-Newtonian viscoelastic behavior of liquid water at high frequencies

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Using classical as well as ab-initio molecular dynamics simulations, we calculate the frequency-dependent shear viscosity of pure water and water-glycerol mixtures. In agreement with recent experiments, we find deviations from Newtonian-fluid behavior in the THz regime. Based on an extension of the Maxwell model, we introduce a viscoelastic model to describe the observed viscosity spectrum of pure water. We find four relaxation modes in the spectrum which we attribute to i) hydrogen-bond network topology changes, ii) hydrogen-bond stretch vibrations of water pairs, iii) collective vibrations of water molecule triplets, and iv) librational excitations of individual water molecules. Our model quantitatively describes the viscoelastic response of liquid water on short timescales, where the hydrodynamic description via a Newtonian-fluid model breaks down.

I. INTRODUCTION

Liquid water is an ubiquitous medium on earth, and of fundamental importance for all organisms. 1, 2 A standard model for the large-scale hydrodynamics of liquid water is the Newtonian fluid, 3–5 where one assumes a linear relationship between local instantaneous stresses and rates of strain, with the proportionality constant given by the viscosity. The combination of the momentum conservation equation with this relation is then known as the Navier-Stokes equation, and it is the basis for hydrodynamics.

Despite its success for describing dynamics of water and other liquids, this model has a limited range of applicability. At high frequencies, when time scales are comparable to those of molecular kinetics within the liquid, real water deviates from the Newtonian fluid model. Slie et al. 6 showed by ultrasound absorption measurements that in aqueous glycerol solutions, at high frequencies the shear viscosity decreases and the mixture starts to have an elastic response under shear deformation. More recently, Pelton et al. 7 showed that the same occurs for pure water. Both studies replaced the Newtonian fluid model by a viscoelastic Maxwell fluid, which in the low frequency limit reduces to a Newtonian fluid but can account for the experimentally observed elastic behavior at high frequencies. Molecular dynamics simulations of water-glycerol mixtures 8 and pure water 10, 11 also find a non-Newtonian regime at high frequencies, the onset of which is well-described by a Maxwell model. 9, 11

Understanding this non-Newtonian behavior is becoming increasingly important. First, with the advancement of nanotechnology, small structures, when in an aqueous environment, start to probe the regime where the Newtonian-fluid model of water breaks down. 7,12 Second, the THz regime probed by modern spectroscopic methods constitutes the boundary between collective and single-water dynamics. 12–16 If one seeks detailed insight into dynamics of molecular or collective processes of solutes on such fast time scales, an important aspect is therefore understanding how water itself behaves on the relevant time and length scales.

In the present work, we use both force field molecular dynamics (MD) and ab-initio molecular dynamics (aiMD) simulations to probe the high-frequency behavior of both pure water and water-glycerol mixtures. From our simulations, we extract the frequency-dependent shear viscoelasticity. We verify our method by comparing to the experimental results of Slie et al. 6 for the viscoelasticity of water-glycerol mixtures. Then we investigate the MD spectrum of pure water in more detail. We propose a viscoelastic model to account for deviations of the observed water viscosity spectrum from the Maxwell model. We identify four independent relaxation modes in the THz regime and link them to molecular processes, namely i) hydrogen-bond network topology changes, ii) hydrogen-bond stretch vibrations, iii) collective vibrations of water molecule triplets, iv) librational excitations of individual water molecules. Our viscosity spectrum based on ab-initio MD (aiMD) simulations shows the same high-
frequency features as the spectrum obtained from force field MD simulations, and thus validates the latter. The aiMD spectrum additionally contains features originating from intramolecular degrees of freedom (OH stretching, OH bending) at large frequencies not included in the rigid water model used for the force field MD simulations.

II. FRIQUENCY-DEPENDENT VISCOSITIES

In this section we recall some generalizations of the standard Green-Kubo relation for the shear viscosity \[^{17,22}\] These generalizations can be used to calculate the frequency- and wave number dependent shear viscosity in terms of velocity and stress tensor correlation functions. Detailed derivations can be found in the Supplemental Material (SM) as well as in the literature\[^{18,20}\] We start from the linearized continuum-mechanical momentum conservation equation\[^{8}\]:

\[
\rho \dot{v}_\alpha(x,t) = \sum_{\beta=1}^{3} \partial_\beta \sigma_{\alpha\beta}(x,t), \quad \alpha \in \{x, y, z\},
\]

where \(\rho\) is the constant equilibrium volume mass density of the fluid, \(\mathbf{v}\) its velocity field, \(\sigma\) its stress tensor, and a dot denotes the time derivative. Note that while for a compressible fluid the density is not constant, deviations from the equilibrium volume mass density \(\rho\) on the left-hand side of Eq. (1) would constitute nonlinear effects and are therefore not considered in our linear treatment. For a linear, homogeneous, isotropic compressible fluid the stress tensor is given as

\[
\sigma_{\alpha\beta}(x, t) = -\delta_{\alpha\beta} P(x, t) + 2 \int \int \eta(|x'|, t') \epsilon_{\alpha\beta}(x - x', t - t') \, d^3x' \, dt' + \delta_{\alpha\beta} \sum_{\gamma=1}^{3} \int \int \left( \eta'(|x'|, t') - \frac{2}{3} \eta(|x'|, t') \right) \epsilon_{\gamma\gamma}(x - x', t - t') \, d^3x' \, dt',
\]

where \(P\) is the pressure, \(\delta_{\alpha\beta}\) is the Kronecker delta, the components of the rate of strain tensor \(\dot{\epsilon}\) are

\[
\dot{\epsilon}_{\alpha\beta} = \frac{1}{2} \left( \frac{\partial v_\alpha}{\partial x_\beta} + \frac{\partial v_\beta}{\partial x_\alpha} \right), \quad \alpha, \beta \in \{x, y, z\},
\]

and \(\eta, \eta'\) are the shear and volume viscosity kernels, which for an isotropic medium only depend on the modulus of the vector \(x\).

If the viscosity kernels decay on length- and time scales that are small compared to those on which the rate of strain tensor varies, the stress tensor defined by Eq. (2) can be approximated at\[^{23}\] \n
\[
\sigma_{\alpha\beta}(x, t) \approx -\delta_{\alpha\beta} P(x, t) + 2\eta \epsilon_{\alpha\beta}(x, t) + \delta_{\alpha\beta} \sum_{\gamma=1}^{3} \epsilon_{\gamma\gamma}(x, t),
\]

where

\[
\eta = \int \int \eta(|x'|, t') \, d^3x' \, dt',
\]

\[
\eta' = \int \int \eta'(|x'|, t') \, d^3x' \, dt',
\]

are the standard shear and volume viscosities, which do not depend on space and time. A fluid with stresses given by Eq. (4) is called a Newtonian fluid. If the stress tensor Eq. (4) is used in the momentum conservation equation (1), the linearized compressible Navier-Stokes equation is recovered. While in this work we mostly consider the spatial average of the shear viscosity kernel, we are precisely interested in the dynamics on time scales where the approximation Eq. (1) breaks down, and non-Markovian effects become relevant.

From momentum conservation Eqs. (1), (2), and the equipartition theorem it follows that the shear viscosity kernel \(\eta\) is given in terms of the trace free part of the stress tensor,

\[
\Pi_{\alpha\beta} = \sigma_{\alpha\beta} - \delta_{\alpha\beta} \frac{1}{3} \sum_{\gamma} \sigma_{\gamma\gamma}, \quad \alpha, \beta \in \{x, y, z\},
\]

and\[^{19,24}\]

\[
\bar{\eta}(k, \omega) = \frac{\beta V}{10} \int_0^\infty e^{-i\omega t} \sum_{\alpha,\beta} \left\langle \Pi_{\alpha\beta}(t) \Pi_{\alpha\beta}(0) \right\rangle \, dt,
\]

where \(V\) is the volume of the fluid, \(\beta^{-1} = k_B T\) is the thermal energy with \(k_B\) the Boltzmann constant and \(T\) the absolute temperature, the tilde denotes a combined spatial Fourier transform (with wave vector \(k\))
and temporal half-sided Fourier transform transform (with angular frequency $\omega$), and the average on the right-hand side is to be understood as an ensemble average over space and time. The real part of $\tilde{\eta}$ yields the viscous response under shear, whereas the imaginary part models the elastic response under shear.

Note that, in view of Eqs. (4), (5), $\tilde{\eta}$ evaluated at $\mathbf{k} = 0$ can be thought of as the viscosity kernel for a viscosity that decays on a length scale much smaller than the length scale on which $\dot{\epsilon}$ varies, but including memory effects in time. If additionally the limit $\omega \to 0$ is taken, memory effects in time are also neglected and the Green-Kubo relation for $\tilde{\eta}$ is obtained from Eq. (8),

$$
\tilde{\eta} = \frac{\beta V}{10} \int_0^\infty \sum_{\alpha \beta} \left\langle \Pi_{\alpha \beta}(t) \Pi_{\alpha \beta}(0) \right\rangle \, dt.
$$

(9)

While Eqs. (8), (9), can be used to calculate the viscosity within force field MD simulations, where the space-averaged pressure tensor is available, this is not the case for the ab-initio simulations we also consider in this work. To calculate shear viscosities also from ab-initio simulations, we use that for small wave number $k = |\mathbf{k}|$, the shear viscosity is given by

$$
\tilde{\eta}(k, \omega) = -\frac{\rho}{k^2} \int_0^\infty e^{-i\omega t} \tilde{C}^-\!(k, t) \, dt,
$$

(10)

where a hat denotes a spatial Fourier transform and $\tilde{C}^-\!(k, t)$ is the normalized autocorrelation function of the $\alpha$-component of the transversal velocity,

$$
\tilde{C}^-\!(k, t) = \frac{\langle \tilde{v}^-\!(\mathbf{k}, t) \tilde{v}^-\!(\mathbf{k}, 0) \rangle}{\langle \tilde{v}^-\!(\mathbf{k}, 0) \tilde{v}^-\!(\mathbf{k}, 0) \rangle}, \quad \alpha \in \{x, y, z\},
$$

(11)

with $\tilde{v}^-\!(\mathbf{k}, t) := \sum_\beta (\delta_{\alpha \beta} - k_\alpha k_\beta/k^2) \tilde{v}_\beta$ the transversal velocity. Note that for an isotropic medium, the right-hand side of Eq. (11) does not depend on the component $\alpha$ used and only depends on $\mathbf{k}$ via the modulus $k$, which is why $\tilde{C}^-\!(k, t)$ has no index $\alpha$ and is written as a function of $k$.

In the context of Eq. (10), small wave number means

$$
|k^2 \tilde{\eta}| \approx 1,
$$

(12)

which for realistic values for water, $|\tilde{\eta}| \approx 0.1$ mPa $\cdot$ s, $\rho = 10^3$ kg/m$^3$, gives $|k|^2 n^2/\rho \ll |\omega|$ . Thus, for $|k| = 4$ nm$^{-1}$, corresponding to the smallest wave number resolvable in a typical box in an MD simulation, formula (10) is valid for $\omega \gg 1.6$ ps$^{-1}$, i.e. for frequencies well above 1 THz, which severely limits the applicability of Eq. (10). While in the limit $k \to 0$, condition (12) is always fulfilled and the approximate Eq. (10) is valid, obtaining $\tilde{v}^-\!(\mathbf{k} = 0, t)$ from simulations is typically difficult because large simulation boxes are needed to resolve small wave numbers, and thereby to extrapolate to $\mathbf{k} = 0$.

### III. GLYCEROL SPECTRA

We simulate glycerol solution with glycerol mass fractions 0, 0.2, 0.4, 0.6, 0.8 in TIP4P/2005 water, for details see SM. From the simulations we calculate the respective viscosity spectra at $\mathbf{k} = 0$ using Eq. (8). The resulting spectra are shown as blue and green solid curves in Fig. 1.

For low frequencies, the real parts (blue) of the spectra are constant and the imaginary parts (green) vanish. As the frequency is increased, the real parts of the spectra decrease to zero, while the imaginary parts show peaks. For low glycerol mass fractions, the real part of the shear viscosity shows a non-monotonic behavior with a second peak at around 10 THz, accompanied by a peak in the imaginary part at slightly higher frequencies. For large glycerol mass fractions, this high-frequency peak disappears, and the decay in the real part with its corresponding peak in the imaginary part shift to lower frequencies.

These spectra illustrate the limits of the Newtonian fluid model, as defined by Eq. (1), because for such a fluid one would expect a spectrum with constant real part and vanishing imaginary part over the whole frequency range. Our spectra exhibit deviations from this behavior for high frequencies, indicating that the assumption of temporal locality breaks down once the rate of strain tensor varies on the picosecond time scale.

To go beyond Newtonian hydrodynamics, we fit Maxwell models, defined by

$$
\tilde{\eta}(\omega) = \frac{\eta_0}{1 - i\omega \tau},
$$

(13)

with $\eta_0 = \tilde{\eta}(0)$ the steady state shear viscosity and a timescale $\tau$, to the frequency range $\omega < 1$ THz of the glycerol-water spectra, see Fig. 1.

In Fig. 2 we compare the fitted viscosities and relaxation times $\eta_0$, $\tau$ to experimental results. As can be seen, the viscosity spectrum of the simulated glycerol/water mixtures reproduces very well both the low-frequency shear viscosity $\eta_0$ and the timescale $\tau$ for the Maxwell model. In subplot (a) we additionally include the zero-frequency viscosity, calculated using the Green-Kubo formula Eq. (4).

The good agreement with our Maxwell model serves as a validation that our approach of using Eq. (5) in conjunction with force field MD simulations yields physically meaningful results.
Taking a closer look at the pure TIP4P/2005 water spectrum, shown in Fig. 1 (a), which is very similar to the previously obtained viscosity spectrum using the TIP4P force field\(^{(10)}\), it transpires that a single Maxwell model, which features a monotonically decreasing real part and one peak in the imaginary part, is not able to model the full viscosity spectrum observed in the MD simulation, which contains two peaks in the imaginary part and a non-monotonic real part. A Maxwell model only describes this spectrum for frequencies \( f \lesssim 1 \text{ THz} \), as was noted before\(^{(11)}\).

**IV. THE SHEAR INERTIA MODEL**

To model the observed shear viscosity spectrum of pure water, we consider a general stress-strain relation, which links off-diagonal components of stress and strain rate tensors at \( k = 0 \) via\(^{(12)}\)

\[
\tilde{\sigma}_{\alpha\beta}(\omega) = 2(-i\omega)\tilde{\eta}(\omega)\tilde{\varepsilon}_{\alpha\beta}(\omega) \quad \alpha \neq \beta, \tag{14}
\]

c.f. Eq. (2). This relation is analogous to electrical circuit theory, where one is interested in the total complex impedance \( \tilde{Z}(\omega) \) of a circuit, which links time-dependent voltage \( U \) and time-dependent current \( I \) via\(^{(28)}\)

\[
\tilde{U}(\omega) = \tilde{Z}(\omega)\tilde{I}(\omega). \tag{15}
\]

In analogy to electrical networks, we use Eq. (14) to model the total complex shear viscosity \( \tilde{\eta}(\omega) \) of a viscoelastic network\(^{(28)}\).

The building blocks for electrical circuits are resistor, capacitor and inductor, and each of them has a characteristic complex impedance \( \tilde{Z}(\omega) \), shown in the left column of Table I. In viscoelasticity, usually only viscoelastic analogues of resistor and capacitor,
but not of inductor, are considered. The viscoelastic analogue of the resistor is the dashpot (both resistor and dashpot dissipate energy), while the viscoelastic analogue of the capacitor is the spring (both capacitor and spring can store energy). The rules for calculating the total viscosity of a viscoelastic network, built up by serial and parallel combination of viscoelastic building blocks, are illustrated in Table II.  

\[ \eta = \eta_0/K. \]

As can be seen from Fig. 3 (a), (b), both Kelvin-Voigt and Maxwell model do not exhibit a real part of the complex viscosity that is increasing as a function of frequency. To model the pure water spectrum, and in particular the non-monotonic real part of the viscosity in Fig. 1 (a), we introduce the viscoelastic analogue of the inductor, which we denote by a circle containing a curved arrow, see right column of Table II. This new circuit element models stress generated by shear acceleration. Indeed, substituting \( \theta(\omega) = -i\omega L \) into Eq. (13) and using Eq. (3), it follows that

\[
\sigma_{\alpha\beta} = 2L\gamma_{\alpha\beta} = L\left(\frac{\partial\gamma_{\alpha}}{\partial x_\beta} + \frac{\partial\gamma_{\beta}}{\partial x_\alpha}\right),
\]

meaning that the stress is proportional to the shear acceleration. Substituting Eq. (13) into the momentum conservation Eq. (1), the resulting force can be interpreted as contributing an effective induced mass due to velocity gradients in the fluid. A possible microscopic origin of such an effect is the coupling between shear and rotational degrees of freedom of individual fluid particles (vorticity-spin coupling).  

Using the shear inertia circuit element, we consider the viscoelastic network shown in Fig. 3 (c), which we call the shear inertia model. As we show now, this is a generalization of the Maxwell model which includes shear inertial effects. Using the viscosities of the individual network elements, c.f. Table II and the combination rules, c.f. Table II, the total viscosity of the shear inertia model in Fig. 3 (c) is found to be

\[
\tilde{\eta}(\omega) = \eta_0 \frac{1 + (i\omega)^2 \tau_m}{1 + (i\omega)^2 \tau_0^2}. \tag{16}
\]

where \( \tau_m = L/\eta_0, \tau_0^2 = L/K \). From Eq. (16) it can be seen that this is an extension of the Maxwell model, Eq. (13), which is recovered in the limit \( \omega \tau_m \to 0, \omega \tau_0 \to 0 \), at finite \( \omega \tau_0^2/\tau_m \equiv \omega \tau \).  

An important difference between the Maxwell and the shear inertia model is that the latter features a non-monotonic real part in the complex viscosity; from equating its derivative with zero, it follows that the real part of \( \tilde{\eta}(\omega) \) defined by Eq. (16) is non-monotonic if and only if

\[
\left(\frac{\tau_0}{\tau_m}\right)^2 = \frac{\eta_0^2}{KL} < 1. \tag{17}
\]

In Fig. 3 (c), we show the complex viscosity of the shear inertia model for parameters \( \eta_0 = 1 \text{ mPa} \cdot \text{s}, K = 2 \cdot 10^{12} \text{ mPa} \cdot \text{s}, L = 10^{-12} \text{ mPa} \cdot \text{s}^2 \). In the low-frequency limit the viscosity spectrum approaches that of a Maxwell model, with constant real part.
TABLE I. Electrical and viscoelastic circuit elements

| Electrical circuit | Viscoelastic network |
|--------------------|----------------------|
| Resistor $R$       | $\eta_0$             |
| Capacitor $-i\omega C$ | $K(-i\omega)^{-1}$ |
| Inductor $-i\omega L$ | $-i\omega L$         |

TABLE II. Combination rules for viscoelastic networks

| Combination | Stresses & Strains | Formula for $\tilde{\eta}_{\text{tot}}(\omega)$ |
|-------------|--------------------|-----------------------------------------------|
| Parallel    | $\sigma_{\text{tot}} = \sigma_1 + \sigma_2$ | $\tilde{\eta}_{\text{tot}}(\omega) = \tilde{\eta}_1(\omega) + \tilde{\eta}_2(\omega)$ |
| Serial      | $\sigma_{\text{tot}} = \sigma_1 + \sigma_2$ | $\tilde{\eta}_{\text{tot}}(\omega) = (\tilde{\eta}_1^{-1}(\omega) + \tilde{\eta}_2^{-1}(\omega))^{-1}$ |

TABLE III. Parameters for the viscoelastic model Eq. (18), resulting from a fit to results for TIP4P/2005 water, c.f. Fig. 4 (a-c), (e). Timescales are converted to frequencies via $f = (2\pi \tau)^{-1}$ for ease of comparison with Fig. 3.

| Parameter | Value | Interpretation |
|-----------|-------|----------------|
| $\eta_{0.1}$ | $0.568$ mPa·s | HB network |
| $(2\pi \cdot \tau)^{-1}$ | $0.19$ THz | topology changes |
| $\eta_{0.11}$ | $0.019$ mPa·s | O-O-O vibrations |
| $(2\pi \cdot \tau_{m,11})^{-1}$ | $2.68$ THz | (HB bending) |
| $\eta_{0.111}$ | $0.053$ mPa·s | O-O vibrations |
| $(2\pi \cdot \tau_{m,111})^{-1}$ | $4.03$ THz | (HB stretching) |
| $\eta_{0.1V}$ | $0.052$ mPa·s | Librations |
| $(2\pi \cdot \tau_{1V})^{-1}$ | $28.92$ THz | |

VI. MICROSCOPIC INTERPRETATION OF THE PURE WATER VISCOSITY SPECTRUM

We now give a microscopic interpretation for each of the constituents of the fit of Eq. (18) to the MD viscosities, c.f. Fig. 4 (a), (b), (e).

We first calculate the radial distribution function $g(r)$ for oxygen atoms and use it to obtain the potential of mean force (pmf) $U(r)$ for the effective oxygen-oxygen interaction via Boltzmann inversion,

$$U(r) = -k_B T \ln [g(r)],$$

see Fig. 4 (f). The pmf shows a primary minimum at an oxygen-oxygen distance of approximately $r \approx 2.8$ Å, corresponding to two hydrogen-bound nearest neighbor water molecules.

We start with the microscopic interpretation of
FIG. 4. Viscosity spectrum of pure water. (a): Complex viscosity \( \tilde{\eta} \) from MD simulations of TIP4P/2005 water, calculated using Eq. (5), together with a fit of the viscoelastic network shown in subplot (d) with complex viscosity Eq. (15). The fitting parameters are given in Table III. (b), (e): Real and imaginary parts of the individual constituents of the fitted viscosity from subplot (a); (c): Complex viscosity \( \tilde{\eta} \) from MD simulations of TIP4P/2005 water (replot of subplot (a)), together with the viscosity calculated from ab-initio molecular dynamics (aiMD) simulations using Eq. (10) at finite \( k = 4.016 \, \text{nm}^{-1} \). (d): Viscoelastic network used for modeling the shear response of pure water. The viscoelastic network is a parallel combination of two shear inertia models and two Maxwell models, the resulting complex viscosity is given by Eq. (18); (f): Black line: Potential of mean force \( U(r) \) for the oxygen-oxygen distance in TIP4P/2005 water. Blue and yellow lines: Harmonic potentials, fitted around the first and second minima of the pmf. The colored arrows indicate the microscopic interpretation of the constituents of the viscoelastic model shown in subplot (d), namely escape from the nearest neighbor pmf minimum (I), oscillations around the minima (II, III) and librations of individual water molecules (IV).

Maxwell model I, which shows a peak in the elastic response at \( f \approx 0.2 \, \text{THz} \), c.f. Fig. 4 (e). This elastic response has been observed experimentally for water and in the context of a Yukawa liquid it was recently uncovered that this peak is linked to nearest-neighbor escape barrier hopping within the fluid, with the peak frequency corresponding to the inverse lifetime of the nearest-neighbor topology of a given molecule. Mean first-passage times for the dissociation of a water-water pair from the nearest neighbor pmf minimum for SPC/E water are about \( \tau_{\text{escape}} \approx 4 - 5 \, \text{ps} \), which translates to frequencies \( 1/\tau_{\text{escape}} \approx 0.20 - 0.25 \, \text{THz} \), in very good agreement with the position of the leftmost peak of the imaginary part in Fig. 4 (f). Multiply- ing Eq. (16) by the denominator of the right-hand side, and performing an inverse Fourier transform, it can be seen that \( \eta(t) \) is the solution of the damped harmonic oscillator equation

\[
\ddot{\eta}(t) + \frac{1}{\tau_m} \dot{\eta}(t) + \frac{1}{\tau_0^2} \eta(t) = 0 \quad (20)
\]

with initial conditions \( \eta(0) = \eta_0 \tau_m / \tau_0^2 \), \( \dot{\eta}(t) = 0 \). With the fitted parameters for shear inertia model III, given in Table III, it follows that the damped harmonic oscillator solution to Eq. (20) is indeed the shear inertia model III. In Table IV we give frequencies for various modes in liquid water obtained from experiments and simulations. Both Raman and infrared (IR) spectroscopy find hydrogen bond stretch vibrations at frequencies \( f \approx 4.5 - 5.5 \, \text{THz} \), while simulation works find them slightly higher, at 6 THz (aiMD) and 6.9 THz (TIP4P/2005), respectively. This suggests that the microscopic origin of the corresponding resonance in our spectrum are vibrations of hydrogen-bound water pairs around the minimum at \( r \approx 2.8 \, \text{Å} \) in Fig. 4 (f). Multiplying Eq. (16) by the denominator of the right-hand side, and performing an inverse Fourier transform, it can be seen that \( \eta(t) \) is the solution of the damped harmonic oscillator equation

\[
\ddot{\eta}(t) + \frac{1}{\tau_m} \dot{\eta}(t) + \frac{1}{\tau_0^2} \eta(t) = 0 \quad (20)
\]
an underdamped oscillation, which is why an extension of the standard (overdamped) Maxwell model is required to describe this feature. The resonance frequency of the underdamped harmonic oscillator defined by Eq. (20) evaluates to

$$f_r = \frac{1}{2\pi} \sqrt{\frac{1}{\gamma_0} - \frac{1}{4\tau_m^2}} \approx 6.7 \, \text{THz}, \quad (21)$$

which is close to the values for hydrogen-bond stretching vibrations found in the literature, see Table 4. Furthermore, the timescales $\tau_0$, $\tau_m$, obtained from the fitted shear inertia model III are in agreement with the intuitive picture of a hydrogen-bound water pair oscillating around the first minimum of the pmf shown in Fig. 4 (f), as we will explain next. To proceed, according to Eq. (20), the frequency for undamped oscillations obtained from the shear inertia model III fit is given by

$$f_{osc,III} = \frac{1}{2\pi\tau_{0,III}} \approx 7.83 \, \text{THz}, \quad (22)$$

and not very different from the result including damping in Eq. (21). The frequency Eq. (22) is comparable to the frequency for an undamped harmonic oscillation of a bound water pair around the first minimum of the pmf,

$$f_{ho} = \frac{1}{2\pi} \sqrt{\frac{k}{m}} \approx 8.82 \, \text{THz}, \quad (23)$$

where the force constant $k = 110.85 \, k_B T/\AA^2$ is obtained by fitting a quadratic potential

$$U(r) = U(r_0) + \frac{k}{2} (r - r_0)^2 \quad (24)$$

to the minimum at $r_0 \approx 2.8 \, \AA$, c.f. the blue dashed curve in Fig. 4 (f), and for $m$ we use the reduced water mass $m = m_{\text{water}}/2 = 9 \, \text{amu}$, appropriate for relative oscillations of two rigid water molecules. From the damping term in Eq. (20), the effective friction coefficient $\gamma$ for a vibrating water molecule pair within the hydrogen bond network is estimated as

$$\gamma \approx m/\tau_m \approx 2\pi \cdot 9 \, \text{amu} \cdot 4.03 \, \text{THz} \approx 0.38 \, \text{pNns nm}^{-1}. \quad (25)$$

This value is almost one order of magnitude smaller than the friction coefficient of a diffusing water molecule, $\gamma \approx 1.62 \, \text{pNns nm}^{-1}$, which physically makes sense because a diffusing water molecule is expected to experience more resistance to motion as compared to a particle oscillating within a local potential minimum. Relative oscillations of hydrogen-bound water molecules contribute to the shear viscosity because of the polarity of an individual water molecule, which couples translation and rotation of individual molecules within the hydrogen-bond network; indeed, as we show in the SM, upon turning off the electrostatic interactions between the water molecules, the feature in the viscosity spectrum disappears. Note furthermore that our interpretation is consistent with the fact that the peak at around 9 THz disappears as the glycerol mass fraction is increased, c.f. Fig. 4, because diffusing glycerol molecules hinder the hydrogen bond network of water. We thus conclude that the microscopic origin of shear inertia model III are hydrogen bond vibrations of neighboring water molecules.

Similarly, we interpret shear inertia model II as oscillations around the second minimum of the pmf shown in Fig. 4 (f), at around $r_0 \approx 4.5 \, \AA$, corresponding to oxygen-oxygen-oxygen (O-O-O) vibrations within the hydrogen bond network of water. Note that, in contrast to shear inertia model III, now the fitted damped harmonic oscillator Eq. (20) is overdamped, so that O-O-O hydrogen bond vibrations are actually overdamped. As we show in the SM, using a standard Maxwell model for feature II is also possible. Indeed, approximating the overdamped mode Eq. (16) by a Maxwell model with relaxation time scale $\tau = \gamma_0/\tau_m$, we find a resonance frequency $f_r \approx (2\pi\gamma)^{-1} \approx 1.7 \, \text{THz}$, which is close to the literature values for hydrogen-bond bending vibrations, c.f. Table 4. Both experiments and simulations locate these vibrations at frequencies slightly higher than our MD water model, see Table 4. While the fitted shear inertia model II describes an overdamped oscillator, employing a shear inertia model instead of a Maxwell model allows us to estimate the effective friction coefficient for O-O-O vibrations, as

$$\gamma \sim m/\tau_m \approx 2\pi \cdot 9 \, \text{amu} \cdot 2.68 \, \text{THz} \approx 0.25 \, \text{pNns nm}^{-1}, \quad (26)$$

where we again use the reduced mass $m = m_{\text{water}}/2 = 9 \, \text{amu}$ as an estimate for the inertia, based on the picture that during O-O-O bending vibrations, the middle molecule does not move significantly. This value for the molecular friction $\gamma$ is of the same order of magnitude as the one obtained for shear inertia model III, c.f. Eq. (25), and is also considerably smaller than the friction coefficient of a diffusing water molecule, $\gamma \approx 1.62 \, \text{pNns nm}^{-1}$. Note finally that also for shear inertia model II, the frequencies obtained from the fit are in agreement with expectations from the pmf in Fig. 4 (f). Indeed, a quadratic fit to the second minimum of the pmf, shown as the yellow dashed line in the figure, leads to an undamped oscillation frequency

$$f_{ho} = \frac{1}{2\pi} \sqrt{\frac{k}{m}} \approx 0.81 \, \text{THz}, \quad (27)$$
TABLE IV. Frequencies for various resonances of liquid water. All frequencies are given in THz.

| Resonance                  | Raman spectroscopy\cite{28} | Infrared spectroscopy (IR)\cite{28,29} | IR from aiMD\cite{30} | IR from MD (TIP4P/2005)\cite{28} | Viscosity from MD (TIP4P/2005, this work) |
|---------------------------|------------------------------|----------------------------------------|-----------------------|-------------------------------|------------------------------------------|
| O-O vibrations (HB stretching) | 4.7                         | ≈ 5.1-5.5                              | 6.0                   | 6.9                           | 6.7                                      |
| O-O-O vibrations (HB bending) | 2.0                         | 1.5                                    | 2.4                   | 1.5                           | 1.7                                      |
| Librations                | ≈ 12-24                     | ≈ 12-21                                | ≈ 18-24               | ≈ 17                          | 28.9                                     |

where as above we use the reduced mass $m = m_{\text{water}}/2$ as an estimate for the inertia. The fitted shear inertia model II yields the comparable frequency

$$f_{\text{osc,II}} = \frac{1}{2\pi\tau_{0,II}} \approx 1.27 \text{ THz.} \quad (28)$$

The highest frequency features, described by Maxwell model IV, we interpret as librational excitations, i.e. rotational vibrations of individual water molecules within the force field of their surrounding molecules. To show this, we in the SM consider orientational spectra calculated for individual water molecules, which show a peak at frequencies comparable to the elastic response frequency of Maxwell model IV\cite{28}. Note that spectroscopy locates such vibrations at frequencies considerably higher than O-O vibrations, c.f. Table IV. That inertial effects can be neglected here is consistent with the fact that for the rotational motion of SPC/E water, inertia starts to dominate only at much higher frequencies, namely at around 90 THz\cite{28}.

In addition to our results obtained from TIP4P/2005, a classical rigid water model, we calculate the viscosity spectrum for pure water also from ab-initio MD (aiMD) simulations, see SM for details\cite{28}. Since no stress tensor is available for the aiMD data, we use Eq. \cite{10} at finite $k = 4.016$ nm$^{-1} = 2\pi/L$, where $L = 1.56$ nm is the edge length of the cubic simulation box. A comparison of aiMD and TIP4P/2005 spectra (the latter as before at $k = 0$) is shown in Fig. \ref{fig:spectra} (c). Qualitative deviations between the spectra appear at frequencies above 50 THz, where intramolecular degrees of freedom\cite{28,30,31} (OH stretching, OH bending), which are not accounted for in a rigid water model, become relevant. Up to these frequencies, however, the spectra show the same features, in particular the peak at around 9 THz with its high-frequency shoulder is present in both the aiMD and the force field MD spectra; this confirms that the spectrum obtained from force field MD is indeed accurate up to about 50 THz. This finding is in agreement with the recent comparison of aiMD and force field MD results for the linear absorption of pure water from 1 MHz to 100 THz\cite{32}.

VII. CONCLUSIONS

In this work we calculate the viscoelastic properties of both pure water and water-glycerol mixtures from force field MD simulations. For water-glycerol mixtures, we find very good agreement of the viscosity and the relaxation time with experimental data\cite{33,34}, for pure water our spectrum agrees with previous theoretical result\cite{10}, and a spectrum calculated from aiMD simulations at finite wave numbers. Using an extension of the Maxwell model for viscoelastic fluids, which includes shear inertial effects, we propose a functional form for the shear viscosity of pure water that is able to describe the force field MD spectrum over the entire frequency range considered. By comparing to Raman and IR spectra, as well as to other observables calculated from our MD data, we subsequently identify the molecular processes underlying this spectrum as water network topology changes, collective vibrations of three water molecules, hydrogen-bond stretch vibrations of water pairs, and librational excitations of individual water molecules.

The viscoelastic circuit we propose describes the short-time non-Markovian behavior of water in the picosecond regime, where the standard Newtonian fluid stress-strain relation becomes a poor approximation to the actual dynamics of the fluid. Only the viscoelastic response at the longest time scales discussed here has been measured until now\cite{35,36,37}, so that our results present a challenge for future experimental investigation.

To obtain an even more detailed picture of the breakdown of the Newtonian fluid picture at small scales, a possible next step would be to also systematically study short-distance non-local effects by considering viscosity kernels at finite wave vector $k$\cite{10,38}.

Note finally that recently it was argued that phonons can travel along the hydrogen bond network, and that elastic peaks in the range of tens of THz should be interpreted as such phonons\cite{39}, which
is consistent with our interpretation of these peaks as bond vibrations. Since at time scales smaller than those of water network topology changes, one could think of water as a static network, possibly with defects, starting from a tetrahedral lattice and calculating the corresponding viscoelastic response.\cite{R. M. Christensen, Theory of Viscoelasticity (Cambridge University Press, 2003).}

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