Polarization of acetonitrile under thermal fields via non-equilibrium molecular dynamics simulations

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ABSTRACT
We show that thermal gradients polarize liquid and supercritical acetonitrile. The polarization results in a stationary electrostatic potential that builds up between hot and cold regions. The strength of the field increases with the static dielectric constant or with decreasing temperature. At near standard conditions, the thermal polarization coefficient is \( \sim -0.6 \text{ mV/K} \), making it possible to induce significant electrostatic fields, \( \sim 10^3 \text{ V/m} \), with thermal gradients \( \sim 1 \text{ K/\mu m} \). At supercritical conditions, \( \sim 600 \text{ K} \) and \( 0.249 \text{ g/cm}^3 \) (the critical isochore), the electrostatic field is of the same order, despite the low dielectric constant of the fluid. In this case, the electrostatic field is determined by the enhanced rotational diffusion of the molecules and stronger cross-coupling between heat and polarization fluxes. We show that the coupling between the heat and polarization fluxes influences the thermal conductivity of acetonitrile, which becomes a worse heat conductor. For the thermodynamic states investigated in this work, the thermal polarization effect leads to a \( \sim 2\%–5\% \) reduction in thermal conductivity.

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I. INTRODUCTION

Temperature gradients are ubiquitous in nature and industry. There is ongoing interest in the use of thermal gradients to induce thermophoretic transport in solutions or to recover waste heat. The application of a thermal gradient to an ionic solution induces a concentration gradient, the Soret effect, and electrostatic potential, the Seebeck effect. In addition to these coupling effects between heat, charge, and mass fluxes, it has been demonstrated that water becomes polarized when exposed to a thermal flux. This phenomenon, called thermal polarization, increases linearly with the magnitude of the thermal gradient and in the case of water has been shown to be particularly strong at near critical conditions. Thermal polarization, or more generally thermal orientation, has also been observed in models of simple polar fluids, \(^{1,2} \) as well as non-polar fluids \( ^{3,4} \) and colloids, \(^{5} \) when the particles are anisotropic in mass and/or size.

Although most investigations to date dealing with real molecular fluids have focused on water, thermal polarization should also be observed in other highly polar fluids. Examining the response and physical behavior of different fluids is necessary to explain more generally the behavior of polar fluids under thermal fields and the first step in rationalizing correlations between thermal polarization and molecular structure. Moreover, this understanding might inspire avenues for applications. Thermal orientation effects have been discussed in the electroproporation of biological membranes, \(^{6,7} \) the dynamical ordering of quantum fluids, \(^{8,9} \) and the generation of thermal monopoles. \(^{10} \) The importance of these effects in other polar fluids has, however, not been fully addressed.

In this work, we investigate the thermal polarization effect in acetonitrile (ACN). ACN is a highly polar fluid, which is widely used as an aprotic solvent in synthesis and energy applications. ACN also features a strong response to electromagnetic fields, e.g., the optical Kerr effect, with a Kerr constant about half of that of
o-nitrotoluene, which is among the liquids featuring the strongest birefringence response.\(^1\) Shelton\(^1\) measured the depolarized hyper-Rayleigh scattering spectra of ACN. The results were interpreted as indicative of a ferroelectric transition with the formation of small ferroelectric domains. This motivated computer simulation studies to test the existence of such domains.\(^2\) It was concluded that acetonitrile does not feature local ferroelectric domains, in disagreement with the conclusions of the experimental studies. However, these investigations were performed under equilibrium conditions. The impact of transient or stationary thermal fields on ACN is unknown.

Thermal gradients have been shown to induce polarization in liquid water. This coupling phenomenon can be formalized with non-equilibrium thermodynamics theory and described in terms of phenomenological coefficients and the dielectric permittivity. This phenomenon depends on the general properties of polar fluids, and we therefore expect that it will be present in acetonitrile as well. In this work, we investigate the thermal polarization of acetonitrile and quantify the strength of the effect at standard and supercritical conditions. Furthermore, we quantify the reduction in the thermal conductivity of ACN, as induced by the thermal polarization effect.

II. METHODS

We performed equilibrium and non-equilibrium molecular dynamics simulations of acetonitrile at thermodynamic conditions ranging from standard to supercritical. Acetonitrile molecules were modeled as rigid bodies using the TraPPE force field [see Fig. 1(c)], which predicts thermophysical properties in good agreement with experimental results.\(^3\) The molecule is modeled using a united atom representation and rigid bonds. The equations of motion were integrated using the velocity Verlet algorithm and a standard time step of 2 fs. Rigid body dynamics were evolved using two virtual sites located along the ACN axis,\(^13,15\) whose distance was kept constant using the LINCS algorithm.\(^17\) The coordinates of the CH\(_3\), C, and N interaction sites were represented relative to the position of these virtual sites such that the total force and the total torque are conserved.\(^15,16\) Electrostatic interactions were calculated using the particle mesh Ewald method.\(^1\) A cutoff of 1.06 nm was used for the dispersion interactions. All trajectories were generated with the code Gromacs v. 2020.\(^18,19\)

Boundary-driven non-equilibrium molecular dynamics (NEMD) simulations using a restraint approach were used to model the fluids in the stationary state. An orthorhombic simulation cell of dimensions \((L_x, L_y, L_z) = (3.564, 3.564, 35.639)\) nm was used, with 3D periodic boundary conditions. The two thermostating regions, hot and cold, were located in the centre and edges of the box, respectively (see Fig. 1). The thermostating regions extend in the \((x, y)\) plane such that the temperature gradients are generated along the \(z\)-direction. In the stationary state, this setup results in two equal but opposite temperature gradients, and therefore in equal and opposite heat fluxes, such that the system is completely periodic.

The thermostats were applied, every time step, to acetonitrile molecules located in the thermostating regions. We restrained the position of the carbon atom bonded to nitrogen to its initial \(z\)-coordinate using a harmonic potential with a force constant of \(10^3\) kJ mol\(^{-1}\) nm\(^{-2}\). The restraint was applied only in the \(z\)-direction: the molecules were free to rotate and also translate in the \(x\)- and \(y\)-directions. The temperature difference between the hot and cold thermostats was set to 100 K for all the NEMD simulations reported in this work.

The thermostating process was implemented using the Nosé-Hoover thermostat, which reproduces the canonical (NVT) ensemble. A single thermostat was applied to each of the thermostating regions only, using one chain, while the trajectories of all other molecules were generated following standard Newtonian dynamics. The Nosé–Hoover equations include an equation of motion for the dynamic variable \(p_i\), which models the dynamic exchange of heat of the system at temperature \(T\), with a reservoir that is set to the desired temperature \(T_r\),

\[
\frac{dv_i}{dt} = \frac{f_i}{m_i} - \frac{p_i}{Q} \frac{d}{dt} \left(\frac{p_i}{Q}\right),
\]

\[
\frac{d(p_i/Q)}{dt} = \frac{T - T_r}{Q},
\]

where \(Q = \tau_T / \pi T_r^4\), \(\tau_T\) defines the period of the temperature fluctuations, which was set to 1 ps. \(v_i\) and \(f_i\) are the velocity and the force acting on particle \(i\) with mass \(m_i\). The Nosé–Hoover Hamiltonian incorporates the contributions due to the reservoir,

\[
H(r^N, p, \xi) = \sum_{i=1}^N \frac{p_i^2}{2m_i} + U(r^N) + \frac{2Q}{Q} N_{dof} k_B T \xi,
\]

where \(r^N = (r_1, r_2, \ldots, r_N)\), \(\xi = dx/dt = p_i/Q\), and \(N_{dof}\) is the total number of degrees of freedom of the region being thermostatted. The last two terms in Eq. (3) represent the energy exchanged with the reservoir. The heat flux, \(J_q = (0, 0, z J_q)\), can be obtained from the time-averaged value of \(\xi\) using the continuity equation

\[
J_q = -\frac{\langle \xi \rangle N_{dof} k_B T}{A},
\]

where \(A = (L_x \times L_y)^2 = 3.564^2\) nm\(^2\) \((L_x \equiv L_y)\) is the cross-sectional area of the simulation box. The simulations were performed with systems consisting of 1590–5050 molecules to target different densities (see Fig. 2 and Table I for a summary of the systems investigated in this work). Each thermostat was coupled to 33 molecules, giving \(N_{dof} = 165\) (5 degrees of freedom per molecule). Similar numbers of molecules were used in our previous NEMD simulations, providing well-defined heat fluxes and thermal gradients.\(^7\)

After the stationary state was reached, each NEMD simulation involved trajectories of either 4 ns or 8 ns. Longer trajectories were used for the low density systems. To ensure good statistics, for each state point, we simulated several replicas, 5 for high densities and up to 25 (200 ns of sampling) for low densities \((\rho = 0.249\) g cm\(^{-3}\)). The fluctuations for these low density systems are large, particularly as the critical temperature is approached. Extensive sampling was needed to obtain aver-
FIG. 1. Non-equilibrium molecular dynamics simulation setup. (a) Snapshot of the simulation box: thermostat regions are represented by enlarged red (hot) and cold (blue) molecules; acetonitrile molecules are comprised of C atoms (gray), N atoms (sky blue), and CH$_3$ groups (cyan). (b) Representative temperature (middle) and density (bottom) profiles for the systems investigated in this work. The blue (cold) and red (hot) shaded areas indicate the location of the thermostats in the simulation box. Average simulation cell densities are shown next to the corresponding density profile. The density profiles include all the molecules in the system, while the temperature profiles exclude the restrained molecules used to thermostat the system in the hot and cold regions. The pressure tensor component $P_{zz}$ parallel to the heat flux vector is shown in the legend for each profile. (c) Graphical representation of the TraPPE force field of acetonitrile. The charge $q$ and Lennard-Jones parameters $\epsilon$ and $\sigma$ are shown under their corresponding species, while rigid bond distances are shown above the molecule. The same color coding as in (a) is used.

ages within acceptable statistical uncertainty. The profiles were computed using 500 bins. The mass and charge densities were obtained by assigning each atom to its corresponding bin. The electrostatic field was obtained from the integration of the charge density.

We also performed equilibrium simulations in the isothermal-isobaric (NPT) ensemble to test the equation of state obtained from NEMD. Simulations in the canonical (NVT) ensemble using the CSVR$^{21}$ (v-rescale) thermostat were performed in order to compute (1) coexistence lines from simulations of explicit liquid–vapour interfaces, using the direct coexistence method, and (2) dielectric properties and Debye relaxation times. A typical simulation consisted of 1000 molecules, 1 ns of equilibration time, and 9 ns of production, with typically 6–10 statistically independent replicas. We employed a smaller time step, 1 fs, to calculate the dipole moment correlation functions.
TABLE I. Thermodynamics states simulated in this work using non-equilibrium molecular dynamics. \( \rho_{box} \), \( T_{box} \), and \( P_{zz} \) represent the average density, temperature, and pressure parallel to the direction of the heat flux, respectively. \( T_{r} \) and \( T_{c} \) are the temperatures of the hot and cold thermostats, \( \lambda \) is the thermal conductivity, \( S_{TP} \) is the thermal polarization coefficient, and \( E \) and \( E_{P} \) are the total electrostatic field and the dipolar contribution to the total field, respectively. All local properties (columns 8–12) were computed at the thermodynamics conditions specified by density \( \rho \) and temperature \( T \). Statistical uncertainties (standard error of the mean) in the least significant figure are given in parentheses.

<table>
<thead>
<tr>
<th>( \rho_{box} ) [g cm(^{-3})]</th>
<th>( T_{r} ) [K]</th>
<th>( T_{c} ) [K]</th>
<th>( T_{box} ) [K]</th>
<th>( P_{zz} ) [bar]</th>
<th>( \rho ) [g cm(^{-3})]</th>
<th>( T ) [K]</th>
<th>( \nabla T ) [K nm(^{-1})]</th>
<th>( \lambda ) [W/(K m)]</th>
<th>( S_{TP} ) [mV K(^{-1})]</th>
<th>( E ) [mV nm(^{-1})]</th>
<th>( E_{P} ) [mV nm(^{-1})]</th>
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<td>750</td>
<td>699</td>
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<td>700</td>
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<td>−0.3(1)</td>
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</table>

III. RESULTS

The thermodynamic states investigated in this work are shown in Fig. 2. We targeted the standard state, high temperature/density systems, and states along the critical isochore. It has previously been shown that the TraPPE force field accurately reproduces the phase diagram of acetonitrile, including the critical point.\(^{14}\) Our coexistence densities, obtained using the direct coexistence method, are in good agreement with these previous simulations.\(^{13}\) With regards to the NEMD data, we show in Fig. 1 representative temperature and density profiles for selected thermodynamic states. Only spatial bins located within 4 nm to the centre of the simulation cell were used in order to exclude any possible artifacts arising from the thermostats. Each set of profiles corresponds to the equation of state (EOS) at the simulated pressure, defined by the component \( \rho_{zz} \) parallel to the heat flux vector. The EOS obtained from NEMD (see Fig. 2) reproduces the behavior of the fluid at equilibrium. This is equivalent to stating that the NEMD simulations generate a collection of states that fulfill local equilibrium, a key assumption in linear non-equilibrium thermodynamics. We demonstrate that this is the case in the inset of Fig. 2. In all cases, the NEMD simulations employed temperature gradients between 1 K nm\(^{-1}\) and 5 K nm\(^{-1}\). This magnitude le the thermal gradient is widely used in computer simulations as it provides a small noise to signal ratio, making it feasible to compute the thermal polarization.\(^{14,23,24}\) Although experimental setups at the micro and nanoscale are approaching gradients of this magnitude,\(^{23,24}\) the simulated gradients are large by macroscopic standards. However, the good agreement between NEMD and equilibrium-NPT results indicates that the response of our systems is in the linear regime and that the local equilibrium hypothesis is fulfilled by our simulations. This observation agrees with previous conclusions using different molecular fluids.

We have demonstrated above that locally the NEMD simulations reproduce the equation of state of the fluids. We took

FIG. 2. Thermodynamic states investigated in this work with reference to the coexistence curve of acetonitrile. Literature values from previous simulations\(^{1}\) are represented as crosses with dashed lines (→). The colored triangle-down (▽) symbols along the coexistence line represent coexistence densities obtained in this work using the direct coexistence method. We also show the temperature/density data obtained from the NEMD simulations [dots (•), which may appear as thick lines]. Our data are color coded according to the pressure of the system. The inset compares the equation of state obtained from NEMD at \( P = (43 ± 8) \) bar [dots (•)] and the data obtained from equilibrium (NPT) simulations at the same pressure [circles (○)].
advantage of this to calculate the thermal conductivity of acetonitrile at different thermodynamic conditions using Fourier’s law ($\lambda = -1/\rho dT/dz$) and the local temperature gradient, at the temperature and density of interest. The temperature gradient was determined by fitting a straight line to the temperature profile within a range of ±2 nm around the selected state point. We note that the computation of the thermal conductivity via NEMD includes all possible coupling effects (e.g., those arising from mass-heat flux coupling"). We discuss later in this work the coupling between heat and polarization fluxes and how it influences the thermal conductivity of acetonitrile.

The thermal conductivities computed in this work are shown in Table I and Fig. 3. The TraPPE force field predicts $\lambda = (0.239 \pm 0.007) \text{WK}^{-1} \text{m}^{-1}$ at 303 K and 0.770 g cm$^{-3}$, corresponding to a pressure of $P = (43 \pm 8)$ bars. This result is on the order of the thermal conductivity reported experimentally at 293.15 K and ambient pressure of $P = 0.195 \text{WK}^{-1} \text{m}^{-1}$. The TraPPE force field predicts values of $\lambda$ at different thermodynamic conditions using Fourier’s law ($\lambda = -1/\rho dT/dz$), given by

$$E = \left(1 - \frac{1}{\varepsilon_s} \right) \frac{L_{pp} \nabla T}{T},$$

where $\varepsilon_s$ is the static dielectric constant of the fluid and $L_{pp}$ are the phenomenological coefficients, with $L_{pq}$ measuring the strength of the heat-polarization coupling and $L_{pp}$ being connected to the relaxation time of the polarization (see below). Equation (5) predicts an increase in the TP response with the dielectric constant of the fluid, the ratio $L_{pp}/L_{pp}$, and the thermal gradient and a decrease with temperature. The strength of the TP effect can be quantified using the thermal polarization coefficient, $S_{TP} = E/\nabla T$.

The phenomenological coefficient $L_{pp}$ appearing in Eq. (5) is related to the Debye relaxation time, $\tau$, via

$$P(t) = kE(1 - \exp(-t/\tau)), \quad \text{where } P \text{ is the polarization, which in the absence of free charges is given by } P = -e_0E,$$

$$\text{and } \kappa = (\varepsilon_s - 1)e_0.$$

The phenomenological coefficient $L_{pp}$ is then defined as

$$L_{pp} = \frac{\kappa T}{\tau},$$

hence providing an approximation to obtain $L_{pp}$ within the assumptions used in the Debye model. Once $L_{pp}$ is known, it is possible to extract $L_{pq}$ from Eq. (5), provided $E$, $\varepsilon_s$, $\nabla T$, and $T$ are known. All these quantities are either defined in our simulation setup ($\nabla T$ and $T$) or are accessible through NEMD or equilibrium simulations ($E, \varepsilon_s$).

The electrostatic field induced by the thermal gradient was calculated by integrating the total charge density along the direction of the heat flux,

$$E(z) = \frac{1}{\varepsilon_0} \int_0^z \rho_q(z')dz',$$

where $\varepsilon_0$ is the vacuum permittivity and $\rho_q(z) = (\sum_{i=1}^{N} \delta(z - z_i)q_i)/A$ is the charge density, with $A$ being the cross-sectional area of the simulation box. The sum runs over all $N$ charges in the system. We also calculated the dipolar field, $E_F$ (see Ref. 2),

$$E_F(z) = -\frac{1}{\varepsilon_0} \int_0^z \frac{dP_q(z')}{dz'} dz',$$
The strength of the TP effect depends on the thermodynamic conditions, density and temperature, highlighting the dependence of the phenomenological coefficients with these properties. At high densities, $\rho = 0.770 \text{ g cm}^{-3}$, the polarization is strongest at near standard conditions and decreases with temperature by $\sim 60\%$ from 300 K to 700 K. Similarly, $|S_{TP}|$ decreases with temperature along the critical isochore, decreasing by $\sim 70\%$ from 600 K to 660 K. The enhancement in TP as the system approaches the critical temperature resembles the behavior reported previously for water. At 607 K ($T/T_c \sim 1$ to 0.1), $S_{TP} = -(0.3 \pm 0.1) \text{ mV K}^{-1}$ (see Table I) has a similar magnitude to that near standard conditions. This value is one order of magnitude lower than that obtained for water at the same $T/T_c \sim 1$, $-2.5 \text{ mV K}^{-1}$, highlighting differences in the TP response of these two polar fluids. We also note that $|S_{TP}|$ increases with density along the $T = 700 \text{ K}$ isotherm by $\sim 200\%$ from 0.333 $\text{ g cm}^{-3}$ to 0.770 $\text{ g cm}^{-3}$.

For all the systems investigated in this work, the thermal polarization coefficient of acetonitrile is negative, $S_{TP} < 0$. This is in contrast with the TP effect in water, where a field inversion is observed. This inversion phenomenon in water is connected to the balance of the dipolar and higher order contributions to the electrostatic field, with the latter changing significantly with the thermodynamic conditions. We report in Fig. 5 the dipolar and higher order contributions to the total thermal polarization field of acetonitrile. At high densities, $\rho = 0.770 \text{ g cm}^{-3}$, the dipolar term is negative for all temperatures investigated and is the dominant contribution ($\sim 70\%$--$90\%$, noting the high uncertainties of $\sim 10\%$--$30\%$ associated with these percentages) to the total field. $E_{TP} < 0$ indicates that the molecules orient preferentially with the dipole pointing toward the hot region. The small impact of higher order terms in defining the field explains the lack of inversion in the TP field at this high density. Along the critical isochore, the situation is more complex. In all cases, $E_{TP} > 0$ (or straddles 0 given the uncertainty), indicating that the molecules orient preferentially with the dipole pointing toward the cold region. Our results indicate that at high temperatures, the dipolar and higher order term contributions have similar values, but opposite signs, while the higher order terms dominate the TP response upon approaching the critical temperature. This result follows previous observations for water, where contributions beyond the dipolar term became dominant near critical conditions, mirroring the increase in the thermal expansion coefficient.

Next, we briefly consider the relative importance of the parameters given in Eq. (5) to $S_{TP}$. As shown in Table II, $\epsilon_s$ features only a weak dependence with temperature: at a density of $\rho = 0.770 \text{ g cm}^{-3}$ ($\rho = 0.249 \text{ g cm}^{-3}$), $\epsilon_s$ decreases by $\sim 8\%$ ($\sim 1\%$) from
300 K to 700 K (610 K–670 K), while $S_{TP}$ changes by ~60% (~70%). These small changes in $\varepsilon_r$ correspond to even smaller differences of <1% in $(1 - 1/\varepsilon_r)$. Since $TS_{TP}$ is not constant with $T$, $S_{TP}$ must be interpreted in terms of changes in both $T$ and $L_{pp}/L_{pp}$. We address the latter in the following, by calculating $L_{pp}$ and $L_{pp}$.

The phenomenological coefficients can be obtained using Eqs. (5) and (7). To calculate $L_{pp}$, we computed the static dielectric constant $\varepsilon_r$ and Debye relaxation time $\tau$ of acetonitrile. $\varepsilon_r$ was calculated from the analysis of the total dipolar fluctuations in the simulation box,

$$\varepsilon_r = 1 - \frac{\langle \mathbf{P}^2 \rangle - \langle \mathbf{P} \rangle^2}{3k_B T_0},$$

(11)

where $\mathbf{P} = \sum_{i=1}^{N} q_i \mathbf{r}_i$, $V$ is the volume of the simulation box, and the brackets $\langle \rangle$ indicate a time average. The Debye relaxation time $\tau$ was calculated from the normalized total dipole moment correlation functions: $\langle \mathbf{P}(t) \cdot \mathbf{P}(0) \rangle / \langle \mathbf{P}(0) \cdot \mathbf{P}(0) \rangle$ and $\frac{1}{N} \sum_{i=1}^{N} \langle \mathbf{P}(t) \cdot \mathbf{p}(0) \rangle / \langle \mathbf{P}(0) \cdot \mathbf{p}(0) \rangle$, respectively, where $\mathbf{p}$ is the dipole moment vector of a single molecule. The dielectric constants were computed using trajectories spanning 9 ns [see Fig. 6(a)], with 6–11 statistically independent replicas to calculate averages and associated uncertainties. The dipole–dipole total and molecular correlation functions were computed using the same trajectories. The resulting correlation functions feature a typical exponential decay. We fitted the initial decay of the correlation functions to a single exponential curve $f(t) = A \exp(-t/\tau_{pp})$, with an upper cutoff for the fitting of 5 ps and 2 ps, for high (0.770 g cm$^{-3}$) and low (0.249 g cm$^{-3}$) density states, respectively. The exception to this is the ($\rho$, $T$) = (0.770 g cm$^{-3}$, 697 K) point, with a cutoff of 3.5 ps. This model assumes the simplest treatment considering a single relaxation time, consistent with Eq. (6).

Our results for the dielectric constant and dipolar correlation times are reported in Table II. At 303 K and 0.770 g cm$^{-3}$, the simulated dielectric constant is close to the experimental value, with a relative deviation of ~13%, which is acceptable considering the simplicity of the model and slight difference in thermodynamic conditions. The Debye relaxation time is also consistent with experimental estimates of 3.5 ps and 3.7 ps. Similar relaxation times are obtained using the total ($\tau_r$) or molecular

<table>
<thead>
<tr>
<th>$\rho$ [g cm$^{-3}$]</th>
<th>$T$ [K]</th>
<th>$\varepsilon_r$</th>
<th>$\tau_p$ [ps]</th>
<th>$\tau_p$ [ps]</th>
<th>$L_{pp} \times 10^{-4}$ [C$^2$/K/(J m)]</th>
<th>$L_{pp} \times 10^{-4}$ [C/m s]</th>
<th>$L_{pp}/L_{pp}$</th>
<th>$\lambda$ [W/(K m)]</th>
<th>$\lambda_{\alpha=0}$ [W/(K m)]</th>
<th>$\delta\lambda$ [%]</th>
<th>$\delta\lambda$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.770</td>
<td>303</td>
<td>31.3(1)</td>
<td>3.061(2)</td>
<td>3.13(2)</td>
<td>2.59(2)</td>
<td>0.45(5)</td>
<td>0.18(2)</td>
<td>0.239(7)</td>
<td>0.248(7)</td>
<td>-0.009(2)</td>
<td>-3.6(8)</td>
</tr>
<tr>
<td>0.770</td>
<td>498</td>
<td>30.0(1)</td>
<td>1.462(1)</td>
<td>1.51(3)</td>
<td>8.5(1)</td>
<td>1.5(3)</td>
<td>0.17(3)</td>
<td>0.236(5)</td>
<td>0.246(6)</td>
<td>-0.010(4)</td>
<td>-4(2)</td>
</tr>
<tr>
<td>0.770</td>
<td>697</td>
<td>28.94(9)</td>
<td>0.929(1)</td>
<td>0.98(2)</td>
<td>17.6(3)</td>
<td>2.6(7)</td>
<td>0.15(4)</td>
<td>0.234(9)</td>
<td>0.24(1)</td>
<td>-0.008(4)</td>
<td>-3(2)</td>
</tr>
<tr>
<td>0.249</td>
<td>607</td>
<td>8.78(1)</td>
<td>0.367(1)</td>
<td>0.47(1)</td>
<td>8.9(2)</td>
<td>-2.0(8)</td>
<td>-0.23(9)</td>
<td>0.26(2)</td>
<td>0.27(2)</td>
<td>-0.012(9)</td>
<td>-5(4)</td>
</tr>
<tr>
<td>0.249</td>
<td>638</td>
<td>8.81(2)</td>
<td>0.348(1)</td>
<td>0.42(2)</td>
<td>10.5(4)</td>
<td>-1.2(4)</td>
<td>-0.12(4)</td>
<td>0.121(5)</td>
<td>0.125(6)</td>
<td>-0.004(2)</td>
<td>-3(2)</td>
</tr>
<tr>
<td>0.249</td>
<td>667</td>
<td>8.84(2)</td>
<td>0.325(1)</td>
<td>0.40(1)</td>
<td>11.6(4)</td>
<td>-0.9(4)</td>
<td>-0.07(3)</td>
<td>0.088(4)</td>
<td>0.090(5)</td>
<td>-0.001(1)</td>
<td>-2(1)</td>
</tr>
</tbody>
</table>
The phenomenological coefficients can be used to assess the impact of the TP effect on the thermal conductivity. It is well established that coupling effects impact transport properties, for example, the Soret effect, where heat-mass flux coupling decreases the thermal conductivity.\textsuperscript{25,29} However, the impact of TP remains unexplored. In this case, the thermal conductivity can be written in terms of the phenomenological coefficients,

\begin{equation}
\lambda = \frac{1}{T^2} \left( L_{qq} - L_{pq} \frac{L_{pp}}{L_{pp}} \right),
\end{equation}

where \(L_{qq}\) is the phenomenological coefficient associated with the heat flux and \(L_{pq} \equiv L_{qp}\) are the cross-coupling coefficients. Following non-equilibrium thermodynamics, \(L_{pq}^{\text{non}} \equiv L_{pp} - L_{pq}^{\text{eq}} \geq 0\); hence, \(L_{pq}^{\text{eq}} \neq 0\) implies that the TP coupling effect should always decrease \(\lambda\). The thermal conductivity values were obtained directly from our NEMD simulations using Fourier’s law and local thermal gradients, calculated for the densities and temperatures of interest (see Table I). Hence, the NEMD thermal conductivities include all possible non-equilibrium effects, in our case the TP coupling. Equation (13) thus provides a route to estimate the reduction in the thermal conductivity associated with the cross-coupling of heat and polarization fluxes. When \(L_{pq}^{\text{eq}} = 0\), such coupling does not exist, and we obtain the corresponding thermal conductivity, \(\lambda_{\text{non}}\). Thus, one can write \(\lambda = \lambda_{\text{non}} + \delta \lambda\), where we compute \(\delta \lambda = -\frac{1}{T^2} \frac{L_{pq}}{L_{pp}}\) directly from variables \(T, S_{TP}, T_p, \) and \(\varepsilon_r\), according to

\begin{equation}
\delta \lambda = \frac{S_{TP}^2 T^2}{\tau} \left( \frac{\varepsilon_r}{\varepsilon_r - 1} \right).
\end{equation}

Our results for \(\delta \lambda\) indicate that the TP effect reduces the thermal conductivity by \(-2\%–5\%\) for the thermodynamics states investigated in this work. In all cases, \(S_{TP}\) carries the largest relative error. This result confirms the notion that fluids that become polarized under a thermal gradient become worse heat conductors.

**IV. CONCLUSION**

We have shown that thermal gradients applied to acetonitrile in the liquid and supercritical states induce thermal polarization.
This is reflected in the build up of an electrostatic field between hot and cold regions. The strength of this thermal polarization field increases with the static dielectric constant and the reduction in temperature, showing that the effect can be significant at near standard conditions, with a TP coefficient \( S_{TP} \sim -0.6 \, \text{mV K}^{-1} \). Large temperature gradients, \( 10^{-5} \, \text{K m}^{-1} \), can be obtained using plasmonic systems;\(^6\) hence, the effect could lead to measurable fields, \( 10^{-5} \, \text{V m}^{-1} \), opening a route to experimental observations of the TP effect at standard conditions. The TP effect might be relevant in thermo-plasmonics, where local polarization effects and the refractive index of the surrounding medium influence the plasmonic response.

We have shown that electrostatic fields of similar magnitude are obtained at near standard and supercritical conditions. The analysis of the phenomenological coefficients defining \( S_{TP} \) shows that at near standard conditions, large dielectric constants and the low temperatures are major factors in contributing to the field strength. In contrast, the TP strength at supercritical conditions is connected to the enhanced rotational diffusion and the stronger cross-coupling between the heat and polarization fluxes. At standard conditions, the TP electrostatic field is dominated by dipolar contributions, while higher order terms become the largest contributors for the supercritical fluid (as is the case for water). Unlike the physical behavior observed in water, the TP field of acetonitrile does not feature a reversal in sign.

We have demonstrated that the TP effect has a measurable impact on the thermal conductivity of polar fluids. We find that the coupling of heat and polarization fluxes leads to a reduction in thermal conductivity of \(-2\%–5\%\), both at near standard and supercritical conditions. While differences in thermal conductivities obtained from non-equilibrium and equilibrium simulation techniques might be masked by statistical uncertainties, these thermal conductivities must be different due to the coupling of polarization and heat fluxes. Equilibrium approaches relying on linear response theory and Green–Kubo equations would thus need to include the cross coefficients to obtain reliable thermal conductivities in a similar fashion to what has been done before\(^{25,26}\) in equilibrium simulations of the Soret effect.

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DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

REFERENCES


