Mie 16-6 force field predicts viscosity with faster-than-exponential pressure dependence for 2,2,4-trimethylhexane

Richard A. Messerly

Thermodynamics Research Center, National Institute of Standards and Technology, Boulder, Colorado, 80305

Michelle C. Anderson

Thermodynamics Research Center, National Institute of Standards and Technology, Boulder, Colorado, 80305

S. Mostafa Razavi

Department of Chemical and Biomolecular Engineering, The University of Akron, Akron, Ohio, 44325-3906

J. Richard Elliott

Department of Chemical and Biomolecular Engineering, The University of Akron, Akron, Ohio, 44325-3906

Abstract

In response to the $10^{\rm th}$ Industrial Fluid Properties Simulation Challenge, we report viscosity (η) estimates obtained with equilibrium molecular dynamics for 2,2,4-trimethylhexane at 293 K and over a range of pressures (P) from 0.1 MPa to 1000 MPa. The Mie Potentials for Phase Equilibria (MiPPE) force field is utilized in this study, as a previous study demonstrated that it provides reliable estimates of η with respect to P. Whereas most studies report only the uncertainties associated with random fluctuations in the simulation output and subsequent data analysis, we investigate the effect of uncertainties arising from the force field non-bonded and torsional parameters. The pressure-viscosity coefficient as a function of pressure is reported for several different empirical models, namely, McEwen-Paluch, Roelands, Roelands-Modified, and Barus. Although the uncertainties increase substantially with increasing pressure, cross-validation model

Email addresses: richard.messerly@nist.gov (Richard A. Messerly), michelle.anderson@nist.gov (Michelle C. Anderson), sr87@zips.uakron.edu (S. Mostafa Razavi), elliot1@uakron.edu (J. Richard Elliott)

selection provides quantitative evidence supporting faster-than-exponential, a.k.a. super-Arrhenius, behavior with an apparent inflection point in a $\log_{10}(\eta)$ -P plot around 200 MPa. Near-quantitative agreement between simulation results and new experimental data is achieved for $P \leq 600$ MPa, followed by significant over prediction at higher pressures.

Keywords:

Pressure-viscosity coefficient, Industrial Fluid Properties Simulation Challenge, Uncertainty Quantification, Molecular Dynamics Simulation, Green-Kubo

1. Introduction

The Industrial Fluid Properties Simulation Challenge (IFPSC) is an open international competition aimed at aligning the molecular simulation community, which is primarily academic, with the goals of industrial research. The present work is a submission to the $10^{\rm th}$ Industrial Fluid Properties Simulation Challenge (IFPSC10)¹, which challenges modelers to predict the viscosity (η) of 2,2,4-trimethylhexane (224TMH) over a wide range of pressures (P), specifically, from 0.1 MPa (atmospheric) to 1000 MPa, at a constant temperature (T) of 293 K.

The practical application of IFPSC10 is elastohydrodynamic lubrication (EHL), where knowledge of the pressure-viscosity relationship is paramount. The challenge compound was chosen as an ideal lubricating oil candidate for which no published experimental viscosity data are available above ambient pressure. New experimental measurements have been performed by Scott Bair of Georgia Tech but were not made available prior to the challenge to avoid the experimental data biasing the simulations and to fairly test the molecular modeling approaches in a predictive manner. The sample purity is greater than 98 % and the estimated experimental uncertainties for η , T, and P are, respectively, 3 %, 0.3 K, and the greater of 1 MPa and 0.4 %.

Classical film thickness formulas rely heavily on the so-called pressure-viscosity co-

¹http://fluidproperties.org/10th

efficient (α) , which is essentially an Arrhenius-like activation parameter that is obtained from the slope of a $\log_{10}(\eta)$ -P plot (see Section 2.4). However, faster-than-exponential, a.k.a. super-Arrhenius, dependence on pressure has been observed through experimental viscometry measurements for nearly a century [1]. This super-Arrhenius trend is typically manifest by an inflection point in the $\log_{10}(\eta)$ -P plot at high pressures. While this behavior is common in experimental measurements, we are not aware of any rheological molecular simulation studies that have addressed this topic, as most simply assume a linear relationship between $\log_{10}(\eta)$ and P [2, 3, 4]. IFPSC10 is an ideal opportunity to demonstrate whether or not molecular simulation can provide evidence supporting or opposing the existence of super-Arrhenius behavior.

In a previous study [5], we investigated four different united-atom (UA) Mie λ -6 (generalized Lennard-Jones, LJ) force fields, namely, the Transferable Potentials for Phase Equilibria (TraPPE-UA [6, 7, 8]), Transferable Anisotropic Mie (TAMie) [9, 10], Mie Potentials for Phase Equilibria (MiPPE, referred to simply as "Potoff" in our previous study) [11, 12], and fourth generation anisotropic-united-atom (AUA4) [13, 14, 15]. Molecular simulation results were compared with experimental data for saturated liquid viscosity ($\eta_{\rm liq}^{\rm comp}$) over a wide temperature range and compressed liquid viscosity ($\eta_{\rm liq}^{\rm comp}$) at 293 K from atmospheric pressure to 1000 MPa, i.e., the same temperature and pressure range as the challenge. The compounds in question were seven n-alkanes ranging in length from ethane to n-docosane and seven branched alkanes ranging in size from 2-methylpropane to 2,2,4-trimethylpentane (224TMP). The 224TMP results at high pressures are especially useful as this compound is a close analogue to the challenge compound and, in contrast with 224TMH, 224TMP has been well studied experimentally.

While TraPPE and AUA4 (LJ 12-6 based potentials) under predict $\eta_{\rm liq}^{\rm sat}$ by 20 % to 50 % for all compounds studied, TAMie (Mie 14-6) and MiPPE (Mie 16-6) predict $\eta_{\rm liq}^{\rm sat}$ within 10 % for most compounds [5]. For $\eta_{\rm liq}^{\rm comp}$, TAMie is the most reliable at predicting the viscosity-density dependence, while MiPPE significantly over estimates $\eta_{\rm liq}^{\rm comp}$ with respect to density. However, since MiPPE also over estimates pressure at high densities

[16], the pressure-viscosity trend for MiPPE is remarkably accurate even at pressures approaching 1000 MPa. In particular, the MiPPE force field predicts the pressure-viscosity trend for 224TMP to within 10 %. For this reason, we implement the MiPPE (Mie 16-6) force field to predict η and α for the challenge compound. We should note, however, that our previous study did not provide any definitive evidence that the MiPPE force field could predict a super-Arrhenius trend for any compound studied, including for 224TMP.

One of the entry guidelines for IFPSC is "an analysis of the uncertainty in the calculated results." Traditionally, simulation uncertainties are limited to the random fluctuations of simulation output and/or the uncertainty related to data post-processing. This class of uncertainty is referred to as "numerical uncertainty" (also frequently referred to as "statistical uncertainty") [17, 18, 19, 20]. Two other classes of uncertainty, namely, "parameter uncertainty" and "functional form uncertainty" (also referred to as "model uncertainty") are typically ignored in uncertainty quantification (UQ) due to the increased computational cost [17, 18, 19, 20]. The latter refers to the uncertainty associated with the choice of force field functional form, while the former refers to the uncertainty in the force field parameters for a given force field functional form.

Quantifying the functional form uncertainty is an extremely difficult task, as it often requires testing numerous force field functional forms. For this reason, we focus on numerical and parameter uncertainties without addressing functional form uncertainties. Specifically, we apply bootstrap re-sampling [21] and Bayesian inference Markov Chain Monte Carlo (MCMC) [16, 17, 18, 19] to quantify numerical and parameter uncertainties, respectively. The chosen functional form is the same as the MiPPE force field, namely, a united-atom, fixed bond length, harmonic angular potential, Fourier series torsional potential, and a Mie 16-6 non-bonded potential (see Section 2.1 for details). As viscosity is highly sensitive to the non-bonded [5, 22] and torsional [14, 23] potentials, we limit our parameter uncertainty investigation to the non-bonded and torsional parameters.

The outline for the present work is the following. Section 2 explains the force field, parameter uncertainties, simulation methodology, data analysis, and empirical models.

Section 3 presents the simulation results, with an emphasis on uncertainty quantification. Section 4 discusses some important observations and limitations. Section 5 recaps the primary conclusions from this work.

2. Methods

2.1. Force field

We utilize the MiPPE force field, as it provides reliable estimates of the η -P dependence for normal and branched alkanes that are similar to the challenge compound [5]. In addition, we quantify the uncertainty in η that is inherited from uncertainties in the non-bonded Mie 16-6 and torsional parameters. The parameter uncertainties are obtained using Bayesian inference Markov Chain Monte Carlo (MCMC). This UQ analysis is performed sequentially. First, we account for only the non-bonded uncertainties (referred to as MCMC-nb). Then, we include both the non-bonded and torsional uncertainties (MCMC-nb-tors). This sequential approach provides insight into which source of uncertainty has a greater impact on η .

2.1.1. MiPPE force field

The MiPPE (Mie λ -6) force field utilizes united-atom (UA) sites, where 2,2,4-trimethylhexane is represented with CH₃, CH₂, CH, and C UA sites (see Figure 1). Neighboring UA sites are separated by a fixed 0.154 nm bond length. Note that we observed in our previous study that η increases by several percent when flexible bonds are employed instead of fixed bonds [5]. Therefore, the choice of fixed bonds was not arbitrary and is a possible source of uncertainty for which we did not rigorously account. The primary reason we utilize fixed bonds is to reduce the fluctuations in the stress tensor and, thereby, decrease the numerical uncertainty of the viscosity estimate.

The angular contribution to internal energy is computed using a harmonic potential:

$$u^{\text{bend}} = \frac{k_{\theta}}{2} \left(\theta - \theta_0\right)^2 \tag{1}$$

Figure 1: United-atom representation of the challenge compound, 2,2,4-trimethylhexane. All bond lengths are 0.154 nm.

where $u^{\rm bend}$ is the bending energy, θ is the instantaneous bond angle, θ_0 is the equilibrium bond angle (see Table 1), and k_{θ} is the harmonic force constant with $k_{\theta}/k_{\rm B}=62500~{\rm K/rad^2}$ for all bonding angles, where $k_{\rm B}$ is the Boltzmann constant.

Table 1: MiPPE equilibrium bond angles (θ_0) [7, 12]. CH_i and CH_j represent CH_3 , CH_2 , CH, or C sites.

Bending sites	θ_0 (degrees)
CH _i -CH ₂ -CH _j	114.0
CH _i -CH-CH _j	112.0
CH_i - C - CH_j	109.5

Dihedral torsional interactions are determined using a modified cosine series:

$$u^{\text{tors}} = c_0 + c_1 [1 + \cos \phi] + c_2 [1 - \cos 2\phi] + c_3 [1 + \cos 3\phi] + A_s \sin^2 \left[\frac{3}{2} (\phi + 180^\circ) \right]$$
$$= (c_0 - A_s) + c_1 [1 + \cos \phi] + c_2 [1 - \cos 2\phi] + \left(c_3 + \frac{A_s}{2} \right) [1 + \cos 3\phi] \quad (2)$$

where $u^{\rm tors}$ is the torsional energy, ϕ is the dihedral angle, c_n are the Fourier constants used in the MiPPE force field and listed in Table 2, and $A_{\rm s} \sin^2\left[\frac{3}{2}(\phi+180^\circ)\right]$ is an additional term proposed by Nieto-Draghi et al. to shift the torsional barrier heights for normal and branched alkanes [14, 15]. We follow a convention similar to that of the International Union of Pure and Applied Chemistry (IUPAC) such that $\phi=180^\circ$ for the

trans conformation [7], whereas Nieto-Draghi et al. define the trans conformation as 0° or 360° [14, 15], hence the $\phi + 180^{\circ}$ term in Equation 2. As $\sin^2\left[\frac{3}{2}(\phi + 180^{\circ})\right]$ has a maximum value of 1 at 0° , 120° , 240° , and 360° , the torsional barriers located at these dihedral angles increase by $A_{\rm s}$. By contrast, this additional term does not shift $u^{\rm tors}$ for dihedral angles of 60° , 180° , and 300° , which correspond to the equilibrium conformations of $gauche^-$, trans, and $gauche^+$, respectively.

Table 2: MiPPE fourier constants (c_n/k_B) and shifting parameter (A_s/k_B) in units of K [7, 12]. CH_i and CH_j represent CH₃, CH₂, CH, or C sites.

Torsion sites	$c_0/k_{\rm B}$	$c_1/k_{ m B}$	$c_2/k_{ m B}$	$c_3/k_{\rm B}$	$A_{ m s}/k_{ m B}$
CH _i -CH ₂ -CH-CH _j	-251.06	428.73	-111.85	441.27	0.0
CH _i -CH ₂ -C-CH _j	0.0	0.0	0.0	461.29	0.0

Note that by expressing the $\sin^2\left[\frac{3}{2}(\phi+180^\circ)\right]$ term as $\cos3\phi$, this additional term does not actually modify the torsional functional form. Instead, it only affects c_0 and c_3 according to $c_0'=c_0-A_{\rm s}$ and $c_3'=c_3+\frac{A_{\rm s}}{2}$, where c_0' and c_3' are the shifted Fourier constants. Clearly, the non-shifted MiPPE torsional potential is obtained only when $A_{\rm s}=0$. The real reason we include this additional torsion term is to provide a simple method for quantifying the torsional parameter uncertainties (see Section 2.1.2).

Non-bonded interactions between sites located in two different molecules or separated by more than three bonds within the same molecule are calculated using a Mie λ -6 potential (of which the traditional Lennard-Jones, LJ, 12-6 is a subclass) [24]:

$$u^{\text{vdw}}(\epsilon, \sigma, \lambda; r) = \left(\frac{\lambda}{\lambda - 6}\right) \left(\frac{\lambda}{6}\right)^{\frac{6}{\lambda - 6}} \epsilon \left[\left(\frac{\sigma}{r}\right)^{\lambda} - \left(\frac{\sigma}{r}\right)^{6}\right]$$
(3)

where u^{vdw} is the van der Waals interaction, σ is the distance (r) where $u^{\mathrm{vdw}}=0$, $-\epsilon$ is the energy of the potential at the minimum (i.e., $u^{\mathrm{vdw}}=-\epsilon$ and $\frac{\partial u^{\mathrm{vdw}}}{\partial r}=0$ for $r=r_{\min}$), and λ is the repulsive exponent.

The MiPPE non-bonded parameters are provided in Table 3. Note that MiPPE provides a "generalized" and "short/long" (S/L) CH and C parameter set. The "general-

ized" CH and C parameter set is an attempt at a completely transferable force field, while the "short" and "long" parameters are implemented when the number of carbons in the backbone is ≤ 4 and > 4, respectively. As the longest continuous carbon backbone for 2,2,4-trimethylhexane consists of six carbons, the MiPPE results presented in this study are obtained with the "long" parameters.

Table 3: MiPPE non-bonded (Mie λ -6) parameters [11, 12]. The CH and C values are the "long" MiPPE parameters.

	MiPPE			
United-atom	$\epsilon/k_{ m B}$ (K)	σ (nm)	λ	
CH ₃	121.25	0.3783	16	
CH_2	61	0.399	16	
СН	14	0.47	16	
С	1.2	0.62	16	

Non-bonded parameters between two different site types (i.e., cross-interactions) are determined using Lorentz-Berthelot combining rules for ϵ and σ [25]:

$$\epsilon_{ij} = \sqrt{\epsilon_{ii}\epsilon_{jj}} \tag{4}$$

$$\sigma_{ij} = \frac{\sigma_{ii} + \sigma_{jj}}{2} \tag{5}$$

where the ij subscript refers to cross-interactions and the subscripts ii and jj refer to same-site interactions.

2.1.2. Parameter uncertainty

Nieto-Draghi et al. set $A_{\rm s}$ equal to 40% and 15% of the maximum dihedral barrier (with the *cis* conformation) for the terminal and internal torsions, respectively [14, 15]. For example, this corresponds to $A_{\rm s}/k_{\rm B}\approx 1000$ K and ≈ 375 K for the CH₃-CH₂-CH₂-CH₂ and CH₂-CH₂-CH₂ torsional potentials, respectively. The primary reason Nieto-Draghi

et al. increase the torsional barrier, i.e., $A_{\rm s}>0$, is because AUA4 under predicts η by approximately 20 % to 40 %. However, despite the relatively large increase in the torsional barriers, the modified force field (AUA4m) typically provides only marginal improvement of 5 % to 15 % compared to AUA4 (see Tables 4 and 5 of Reference 15).

As the MiPPE potential is already quite reliable for predicting viscosity, we would expect significant over prediction of viscosity if we coupled the MiPPE (Mie 16-6) potential with $A_{\rm s}/k_{\rm B}\gg 0$. Thus, unlike Nieto-Draghi et al., we do not propose that the torsional barriers must be increased unilaterally. Instead, we assume that $A_{\rm s}$ follows a skewed normal distribution with a mean value near zero and lower and upper 95 % confidence intervals corresponding to -15 % and +40 % of the maximum barrier height for the non-shifted MiPPE torsional potential. The rationale for the $A_{\rm s}$ distribution is presented in Section S.V of Supplementary Material. The MCMC-nb-tors parameters $(A_{\rm s})$ sampled from this distribution are reported in Section S.VI of Supplementary Material.

Figure 2 compares the non-shifted MiPPE torsional potential, ± 40 % shift in barrier heights, and the MCMC-nb-tors potentials. The insets also depict the skewed distributions and the randomly sampled MCMC $A_{\rm s}$ sets. Note that the challenge compound consists of four ${\rm CH}_i{-}{\rm CH}_2{-}{\rm CH}{-}{\rm CH}_j$ torsions and three ${\rm CH}_i{-}{\rm CH}_2{-}{\rm C}{-}{\rm CH}_j$ torsions. Note that, unlike Nieto-Draghi et al., we make no distinction between internal and terminal torsions.

Figure 3 depicts the MCMC non-bonded parameters for CH₃, CH₂, CH, and C unitedatom sites (ϵ_{CH_3} , σ_{CH_2} , ϵ_{CH_2} , σ_{CH_2} , ϵ_{CH} , σ_{CH} , ϵ_{C} , and σ_{C}) which are used for MCMC-nb and MCMC-nb-tors. Note that $\lambda_{\text{CH}_3} = \lambda_{\text{CH}_2} = \lambda_{\text{CH}} = \lambda_{\text{C}} = 16$. Parameters are assumed to be transferable, e.g., the CH₂ MCMC parameters are sampled independent of the CH₃ MCMC parameters. An important observation from Figure 3 is that the ranges of MCMC sampled CH and C non-bonded parameters are considerably larger (on a percent basis) than those for CH₃ and CH₂. In fact, the MCMC ranges for CH and C encompass both the "long" and "generalized" parameters, suggesting that these are actually indistinguishable parameter sets.

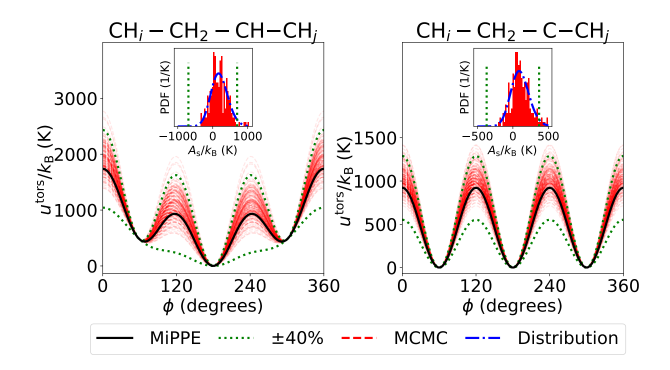


Figure 2: Comparison of MiPPE (black solid lines), ± 40 % (green dotted lines), and MCMC-nb-tors (red dashed lines) torsional potentials. The transparency of the red dashed lines is proportional to the probability density function (PDF). Insets show the distribution for $A_{\rm s}$ as blue dash-dotted lines. Left and right panels correspond to ${\rm CH}_i-{\rm CH}_2-{\rm CH}-{\rm CH}_j$ and ${\rm CH}_i-{\rm CH}_2-{\rm C-CH}_j$ torsions, respectively. Both $u^{\rm tors}/k_{\rm B}$ and $A_{\rm s}/k_{\rm B}$ are expressed in units of K.

The MCMC non-bonded parameters for CH₃ and CH₂ sites were determined in our previous study using a likelihood function based on experimental saturated liquid density and saturated vapor pressure data for ethane, propane, n-butane, and n-octane [16]. By contrast, the MCMC parameters for CH and C sites are obtained from the scoring function reported by Mick et al. [12] that depends on several vapor-liquid coexistence properties for a diverse set of branched alkanes. Details regarding the generation of MCMC parameter sets from the scoring function are found in Section S.III of Supplementary Material. Tabulated non-bonded MCMC parameter sets (ϵ and σ) are provided in Section S.VI of Supplementary Material.

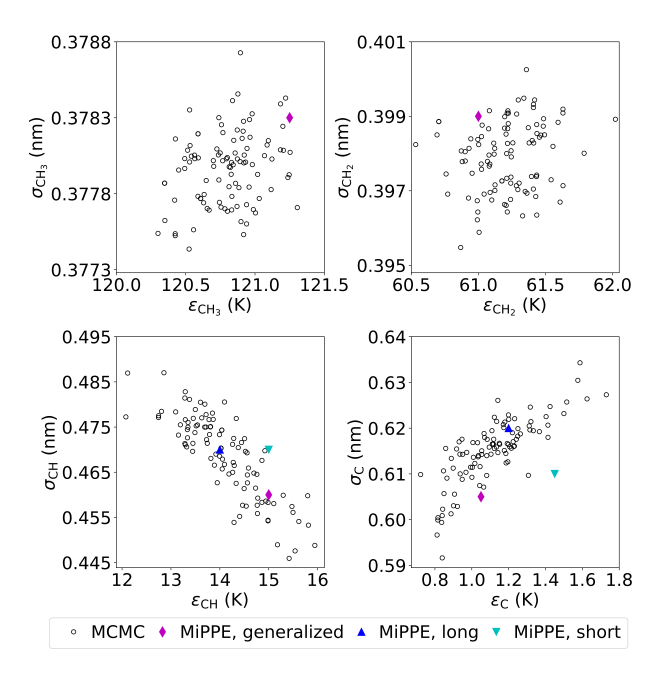


Figure 3: Uncertainty in non-bonded parameters determined with Markov Chain Monte Carlo (MCMC). The MiPPE generalized and S/L parameters are also included as a reference [11, 12]. Top left, top right, bottom left, and bottom right panels correspond to CH₃, CH₂, CH, and C parameters, respectively. Note that a small discrepancy exists between the generalized C parameters in Table 1 of Reference 12 and the optimal region depicted in Figure 1 of Reference 12. The generalized C parameters shown here were obtained from private communication of tabulated scoring function values and are, thus, slightly different than those reported by Mick et al.[12]

2.2. Simulation set-up

Historically, non-equilibrium molecular dynamics (NEMD) has been preferred for highly viscous systems [3, 4]. However, in our recent publication we successfully predicted the viscosity of 2,2,4-trimethylpentane at 293 K and 1000 MPa (the highest pressure required for the challenge) with equilibrium molecular dynamics (EMD) [5]. Consistent with our previous study, we perform EMD simulations using GROMACS version 2018 with "mixed" (single and double) precision [26]. Example GROMACS input files (.top, .gro. and .mdp) with corresponding shell and Python scripts for preparing, running, and analyzing simulations are provided in Section S.II of Supplementary Material.

We utilize the same simulation specifications as our previous study [5]. The general simulation specifications are provided in Table 4. Good performance is achieved with a relatively long time-step of 2 fs because there are no fast vibrating modes. Specifically, all bond lengths are fixed and the UA model does not include explicit hydrogen sites. Bonds are constrained using LINear Constraint Solver (LINCS) with a LINCS-order of eight [27, 28].

Note that we employ a 1.4 nm non-bonded cut-off with analytical tail corrections (where GROMACS only includes the contribution from the r^{-6} term) instead of the 1.0 nm cut-off utilized to develop the MiPPE potential [11, 12]. Although a 1.0 nm cut-off is reliable for most compounds, it can lead to spurious viscosity estimates and even unstable simulations for large molecules, e.g., n-hexadecane and n-docosane [5]. For this reason, we incur the additional computational cost with the longer cut-off so as to avoid any unforeseen simulation anomalies [29].

Finite size effects for viscosity are typically negligible with a 200 or 400 molecule system [5, 30]. The MiPPE simulations utilize the larger system size only for $P \leq 500$ MPa, while the smaller system size is favored at higher pressures due to the significant increase in simulation time (see Table 6). To reduce the computational cost, the MCMC-nb and MCMC-nb-tors simulations use 200 molecules at each pressure. Note that because viscosity is a collective property the uncertainty does not depend on the number of molecules.

Table 4: General simulation specifications.

Time-step (fs)	2
Cut-off length (nm)	1.4
Tail-corrections	U and P
Constrained bonds	LINCS
LINCS-order	8
Number of molecules	200 or 400

We perform a sequence of six simulation stages: energy minimization, NPT equilibration, NPT production, energy minimization, NVT equilibration, and NVT production. The average box size from the NPT production stage is utilized in the second energy minimization and subsequent NVT stages. Table 5 lists the integrators, thermostats, barostats, and simulation time used for each NPT and NVT equilibration and production stage. These specifications are also the same as our previous study, with the exception of the NVT production simulation times, which are state point dependent. The specific production times for the NVT production stage are provided in Table 6. Note in Table 6 that we include simulations at an additional pressure of 975 MPa to reduce the uncertainties in our high pressure viscosity estimates.

A large number of replicate simulations are required at each state point to quantify and reduce the numerical uncertainty [39, 40]. We utilize between 40 and 80 independent replicates, where a greater number of replicates are needed for more viscous systems, i.e., at higher pressures. Each MiPPE replicate simulation utilizes the same force field parameters, whereas MCMC-nb and MCMC-nb-tors utilize different parameter sets for each replicate (see Figures 2 and 3). To ensure independence between replicates, the entire series of simulation stages is repeated, the energy minimization stages start with a different pseudo-random configuration, and the initial velocities are randomized for each *NPT* and *NVT* equilibration stage. The replicates, therefore, sample from a distribution

Table 5: Simulation specifications for equilibration (Equil.) and production (Prod.) stages. $t_{\rm sim}$ is the simulation time, τ_T is the thermostat time-constant, τ_P is the barostat time-constant, and ζ_P is the barostat compressibility.

	NPT Equil.	NPT Prod.	NVT Equil.	NVT Prod.
$t_{ m sim}$ (ns)	1	1	1	1 to 96
Integrator	Leap frog [31]	Leap frog	Velocity Verlet [32]	Velocity Verlet
Thermostat	Velocity rescale [33]	Nosé-Hoover [34, 35]	Nosé-Hoover	Nosé-Hoover
$ au_T$ (ps)	1.0	1.0	1.0	1.0
Barostat	Berendsen [36]	Parrinello-Rahman [37, 38]	N/A	N/A
τ_P (ps)	1.0	5.0	N/A	N/A
ζ_P (1/bar)	4.5×10^{-5}	4.5×10^{-5}	N/A	N/A

of densities (i.e., box sizes) for a given pressure.

2.3. Data analysis

The post-simulation data analysis is identical to that prescribed in our previous study [5]. In brief, we implement the Green-Kubo "time-decomposition" analysis [39, 40]

$$\eta(t) = \frac{V}{k_{\rm B}T N_{\rm reps}} \sum_{n=1}^{N_{\rm reps}} \int_0^t dt' \left\langle \tau_{\alpha\beta,n}(t') \tau_{\alpha\beta,n}(0) \right\rangle_{t_0,\alpha\beta} \tag{6}$$

where t is time, V is volume, N_{reps} is the number of independent replicate simulations, α and β are x, y, or z Cartesian coordinates, $\tau_{\alpha\beta,n}$ is the α - β off-diagonal atomic stress tensor element for the n^{th} replicate, and $\langle \cdots \rangle_{t_0,\alpha\beta}$ denotes an average over twelve different time origins (t_0) and all three unique off-diagonal $\tau_{\alpha\beta}$ components.

For an accurate integration of Equation 6, $\tau_{\alpha\beta,n}$ is recorded every 6 fs (3 time-steps) for $P \leq 700$ MPa. Less-frequent recording of $\tau_{\alpha\beta,n}$ is required to avoid unmanageable file sizes for the three highest pressures, as these systems require significantly longer simulations. Specifically, we record $\tau_{\alpha\beta,n}$ every 18 fs (9 time-steps) for P=800 MPa, every 24 fs (12 time-steps) for P=900 MPa, and every 36 fs (18 time-steps) for P=975 and 1000 MPa. We validate the results at high pressures with less-frequent recording of $\tau_{\alpha\beta,n}$ in Section S.VIII of Supplementary Material.

Table 6: State point specific production times. Pressure is prescribed only in NPT equilibration and production stages.

Pressure (MPa)	NVT Prod. time (ns)
0.1	1
25	1
50	1
100	1
150	1
250	2
400	4
500	8
600	8
700	16
800	64
900	72
975	96
1000	96

The force field viscosity is obtained by evaluating Equation 6 as $t \to \infty$, i.e., the infinite-time-limit viscosity (η^{∞}) . As the long-time tail does not converge, we fit the "running integral" to a double-exponential function

$$\eta(t) = A\alpha \tau_1 \left(1 - \exp\left(-t/\tau_1\right) \right) + A(1 - \alpha)\tau_2 \left(1 - \exp\left(-t/\tau_2\right) \right) \tag{7}$$

where A, α, τ_1 , and τ_2 are fitting parameters and $\eta^{\infty} = A\alpha\tau_1 + A(1-\alpha)\tau_2$. Our previous work provides details regarding the fitting procedure [5].

The uncertainty in η is obtained by bootstrap re-sampling and reported at the 95 % confidence level. Specifically, the fitting of Equation 7 is repeated thousands of times using randomly selected subsets of replicate simulations from which we obtain a distri-

bution for η^{∞} . Section S.IV of Supplementary Material validates the methodology for quantifying the MCMC-nb and MCMC-nb-tors uncertainties, namely, by bootstrap resampling replicate results obtained with different MCMC parameter sets.

2.4. Pressure-viscosity coefficient

The simulated η values with respect to P are fit to four different empirical models from which the pressure-viscosity coefficient (α) is calculated

$$\alpha \equiv \frac{d\ln(\eta)}{dP} \tag{8}$$

As viscosity ranges over several orders of magnitude, the objective function for fitting is the sum-squared-error of $\log_{10}(\eta)$. Only the MiPPE results are included in the fit.

The first empirical model we implement is the traditional Barus expression [1, 41]

$$\eta = \eta_0 \exp(\alpha P) \tag{9}$$

where the fitting parameters η_0 and α correspond to the zero-pressure viscosity and the pressure-viscosity coefficient, respectively. The second model is the popular Roelands equation [1, 42]

$$\eta = \eta_p \left(\frac{\eta_0}{\eta_p}\right)^{\left(\frac{P_p - P}{P_p}\right)^Z} \tag{10}$$

where η_0 and Z are fitting parameters and $\eta_p = 6.31 \times 10^{-5}$ Pa-s and $P_p = -0.196$ GPa. The third model is an alternative form of Equation 10 where η_p and P_p are additional fitting parameters (rather than fixed constants). This four parameter Roelands equation, which we refer to as "Roelands-Modified," is more flexible than the standard Roelands model and can predict super-Arrhenius behavior. The fourth model is a hybrid McEwen-Paluch expression [1]

$$\eta = \eta_0 \left(1 + \frac{a_0}{q} P \right)^q \exp\left(\frac{C_{\rm F} P}{P_{\infty} - P} \right) \tag{11}$$

where η_0 , a_0 , q, $C_{\rm F}$, and P_{∞} are fitting parameters. Note that, although the Roelands-Modified equation can predict super-Arrhenius behavior, only the McEwen-Paluch model

is capable of representing an inflection point, i.e., a transition from Arrhenius to super-Arrhenius behavior. By contrast, the Barus and Roelands models are only capable of fitting sub-Arrhenius and Arrhenius-like data.

3. Results

Table 7 provides tabulated values for viscosity (η) and density (ρ) for MiPPE, MCMC-nb, and MCMC-nb-tors at the prescribed temperature (T) and pressure (P). The uncertainties (expressed at the 95 % confidence level) for T, ρ , and P are computed with the standard deviation of the replicate simulation averages, while those for η are obtained from bootstrap re-sampling. Note that the average temperature slightly decreases with increasing pressure, but T is always within the 0.3 K uncertainty stipulated by the IF-PSC10 experimental measurements. Also, the average pressures tend to be lower than the prescribed pressure. For most state points, however, the combined uncertainty in pressure from experiment (the greater of 1 MPa and 0.4 %) and simulation is sufficient to account for this discrepancy.

The values from Table 7 are also depicted in Figure 4 along with the model fits to the MiPPE values and the IFPSC10 experimental data (see Section S.I of Supplementary Material). Recall that these experimental data were made available only after submitting our simulation results to IFPSC10. Prior to the challenge, the Thermodynamic Research Center database contained only a single experimental viscosity value (at saturation pressure) for this compound [43].

The top panel of Figure 4 demonstrates that an inflection point, suggesting super-Arrhenius behavior, is observed in the simulation results around 200 MPa. However, the inflection point for the IFPSC10 experimental data is much less pronounced, manifested by large systematic deviations between simulation and experimental values for $P \geq 700$ MPa.

The bottom panel of Figure 4 presents the percent deviation between the McEwen-Paluch empirical model fit ($\eta_{\rm MP}$) and the simulation results, IFPSC10 experimental data,

Table 7: Simulation results for MiPPE, MCMC-nb, and MCMC-nb-tors. Subscripts and superscripts (when present) correspond to the lower and upper 95 % confidence intervals, respectively. If only a subscript is present the lower and upper uncertainties are approximately equal (i.e., the bootstrap distribution is essentially symmetric) and only the average uncertainty is reported.

T (K)	ρ (kg/m ³)	P (MPa)	$\eta \ (10^{-3} \ \text{Pa-s})$		
			MiPPE	MCMC-nb	MCMC-nb-tors
292.854 _{0.095}	716.46 _{0.46}	-2.08 _{0.83}	$0.605_{0.045}$	$0.565_{0.038}^{0.048}$	$0.578_{0.043}^{0.059}$
292.851 _{0.085}	734.67 _{0.44}	22.9 _{1.1}	$0.803^{0.078}_{0.098}$	$0.786^{0.079}_{0.062}$	$0.763^{0.087}_{0.056}$
292.847 _{0.089}	749.22 _{0.45}	47.9 _{1.2}	$0.957^{0.076}_{0.061}$	$0.965^{0.099}_{0.070}$	$1.014_{0.091}$
292.85 _{0.10}	772.24 _{0.37}	98.3 _{1.3}	$1.60_{0.14}$	1.50 _{0.11}	$1.59_{0.13}$
292.838 _{0.088}	790.25 _{0.35}	148.3 _{1.6}	$2.18_{0.19}^{0.24}$	$2.10^{0.21}_{0.17}$	$2.18_{0.21}^{0.28}$
292.819 _{0.096}	818.22 _{0.31}	248.5 _{2.0}	$4.56_{0.45}$	$4.25_{0.34}$	$4.36_{0.38}$
292.816 _{0.090}	849.17 _{0.38}	398.7 _{2.7}	13.5 _{1.2}	12.5 _{1.2}	$14.1_{1.2}$
292.790 _{0.099}	865.64 _{0.49}	498.4 _{4.6}	$26.6_{3.1}^{3.8}$	$28.5_{2.7}$	28.6 _{3.1}
292.774 _{0.055}	879.9 _{1.2}	598.5 _{9.5}	$65.0_{9.6}$	66.2 _{7.5}	$61.8^{6.8}_{4.8}$
292.763 _{0.041}	892.6 _{1.2}	698 ₁₀	175 ₂₉	_	_
292.752 _{0.026}	903.5 _{1.1}	806 ₁₄	523 ₆₈	_	_
292.738 _{0.022}	914.4 _{1.0}	899 ₁₁	1480_{206}	_	_
292.725 _{0.020}	921.2 _{1.1}	973 ₁₆	3528 ₃₇₂	_	_
292.719 _{0.013}	923.7 _{1.2}	1000 ₁₇	4948_{554}	_	_

and the other three empirical model fits. This panel demonstrates that near-quantitative agreement is achieved between the simulation results and the IFPSC10 experimental data (to within the combined uncertainties) for pressures ranging from 0.1 MPa to 600 MPa. Furthermore, this panel is useful for comparing the MiPPE, MCMC-nb, and MCMC-nb-tors uncertainties and for visualizing the goodness of fit with the various empirical models. For example, note that both the McEwen-Paluch and Roelands-Modified fits agree with nearly all of the simulation values to within their corresponding uncertainties.

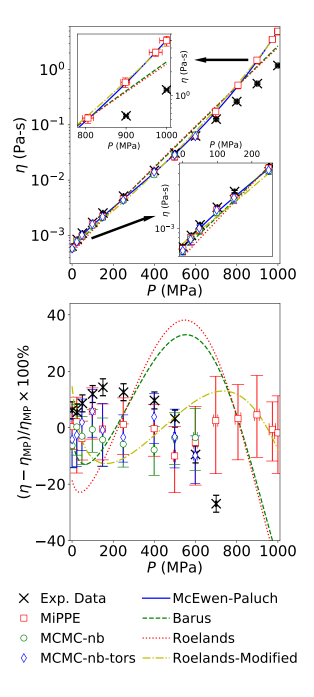


Figure 4: Pressure-viscosity results for MiPPE (red squares), MCMC-nb (green circles), and MCMC-nb-tors (blue diamonds). Top panel plots $\log_{10}(\eta)$ -P where different line colors and styles represent different empirical model fits (Equations 9, 10, and 11) to MiPPE values. Bottom panel is a percent deviation plot relative to the McEwen-Paluch fit ($\eta_{\rm MP}$). Experimental viscosity data above saturation pressure were not available prior to the challenge. Error bars represent 95 % confidence intervals obtained from bootstrap re-sampling. If not depicted, uncertainties are smaller than the symbol size.

Recall that the MiPPE results only account for numerical uncertainties, while MCMC-nb accounts for numerical and non-bonded parameter uncertainties, and MCMC-nb-tors accounts for numerical, non-bonded and torsional parameter uncertainties. Notice in Table 7 and Figure 4 that the MiPPE, MCMC-nb, and MCMC-nb-tors uncertainties are approximately equal. This somewhat surprising result supports the conclusion that the non-bonded and torsional uncertainties are negligible compared to the numerical uncertainties in the Green-Kubo viscosity.

Figure 5 presents the predicted pressure-viscosity coefficient (α), as determined by fitting the MiPPE results to Equations 9, 10, and 11. The uncertainties in α are obtained with bootstrap re-sampling for the empirical model fits.

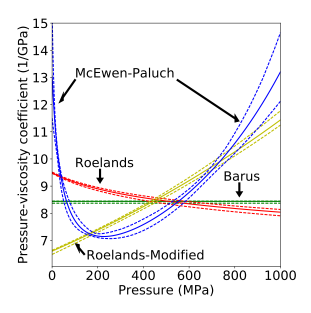


Figure 5: Pressure-viscosity coefficient predicted with empirical model fits (Equations 9, 10, and 11). Dashed lines represent 95 % confidence intervals obtained from bootstrap re-sampling.

As expected, the traditional Barus α value is constant with respect to pressure. By contrast, the Roelands α value actually decreases with increasing pressure, while the Roelands-Modified α value increases with respect to pressure but without a change from negative to positive slope. Only the McEwen-Paluch α value shows the marked change in slope which corresponds to an inflection point in the $\log_{10}(\eta)$ -P plot.

Note that the α magnitudes for all empirical models are reasonable (i.e., similar in magnitude to other lubricants [2, 3, 4]) over the entire range of pressures. In comparison with the IFPSC10 experimental value at 900 MPa (α = 7.52), however, the MiPPE simulation results overestimate the pressure-viscosity dependence at high pressures.

Although the hybrid McEwen-Paluch model clearly reproduces the simulation results with lower deviations than those of the Roelands and Barus models (see Figures 4 and 6), this should be anticipated considering the McEwen-Paluch model has five fitting parameters while the Barus and Roelands models only have two. Note that the four parameter Roelands-Modified model also has lower deviations than the Roelands and Barus models. Therefore, it is possible that the McEwen-Paluch model actually over fits our simulation results.

To assess this possibility, Figure 6 presents the cross-validation results for each model. Specifically, we implement a Monte Carlo cross-validation scheme where thousands of random sub-samples are selected for the training and testing set. Approximately 70 % of the MiPPE simulation results (10 pressures) are included in the training set while 30 % (4 pressures) are excluded for the testing set. The η value at a given pressure is randomly sampled from the bootstrapped simulation uncertainties, such that the complete set of 14 η values varies for each round of cross-validation.

The left panel of Figure 6 demonstrates that the mean-squared-error (MSE) for the training set is approximately equal to the MSE for the testing set of each model, suggesting that none of the models over fit the data. Note that, although somewhat counter intuitive, $MSE_{test} < MSE_{train}$ typically denotes that the training set is "easier" to fit than the testing set. The right panel shows that only the Roelands-Modified model has a similar MSE to that of the McEwen-Paluch model for the same testing set. As both the Roelands-Modified and McEwen-Paluch models predict super-Arrhenius behavior, Figure 6 provides strong statistical evidence that the MiPPE force field predicts super-Arrhenius behavior. However, as the Roelands-Modified model does not predict that the slope of α with respect to P changes signs, there exists some doubt whether an inflection point pre-

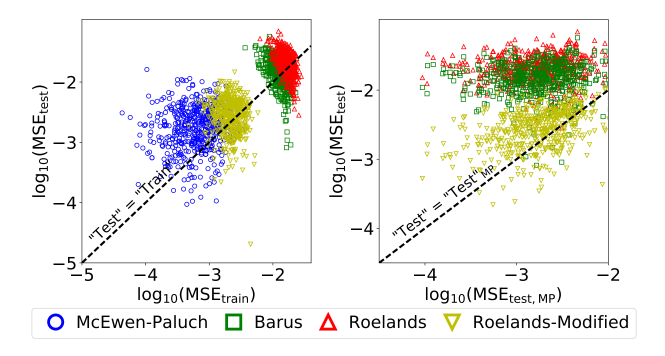


Figure 6: Monte Carlo cross-validation for empirical model fits (Equations 9, 10, and 11). $MSE_{\rm train}$ and $MSE_{\rm test}$ are the mean-squared-error for the "training" and "testing" sets, respectively. Left panel compares $MSE_{\rm train}$ and $MSE_{\rm test}$ for each model, while the right panel compares the McEwen-Paluch $MSE_{\rm test}$ with the $MSE_{\rm test}$ for the other three models.

cedes the super-Arrhenius region.

Table 8 is included to facilitate scoring our entry for the $10^{\rm th}$ Industrial Fluid Properties Simulation Challenge. Table 8 provides "smoothed" η and α values calculated with the McEwen-Paluch fit ($\eta_{\rm MP}$) to the MiPPE simulation results. As our initial fit under estimated the MiPPE η value at atmospheric pressure, the values in Table 8 are obtained by constraining η_0 in Equation 11 to reproduce the simulated viscosity at P=0.1 MPa. Note that, despite performing simulations at both 975 and 1000 MPa, the uncertainty in η at 1000 MPa is considerably larger than at the other pressures.

The uncertainties in Table 8 reflect both the simulation and experimental pressure uncertainties. Specifically, the McEwen-Paluch model is fit thousands of times using bootstrap re-sampling. Each fit is subsequently evaluated at $P^{\rm exp} \pm u_P^{\rm exp}$, where $u_P^{\rm exp}$ is the reported uncertainty in the experimental pressure $(P^{\rm exp})$. The lower and upper 95 % con-

fidence intervals are obtained from the distributions of $\eta_{\rm MP}$ computed at $P^{\rm exp}-u_P^{\rm exp}$ and $P^{\rm exp}+u_P^{\rm exp}$, respectively.

Table 8: Recommended (smoothed) values for the purpose of scoring our entry to the $10^{\rm th}$ Industrial Fluid Properties Simulation Challenge. Uncertainties are expressed at the 95 % confidence level. Pressure uncertainties are those reported for the experimental measurements.

P (MPa)	$\eta \ (10^{-3} \ \text{Pa-s})$	α (1/GPa)
$0.1_{1.0}$	$0.605_{0.045}$	13.1 _{1.8}
25.0 _{1.0}	$0.801_{0.018}$	$10.168_{0.091}$
$50.0_{1.0}$	$1.014_{0.018}$	8.87 _{0.26}
100.0 _{1.0}	$1.528_{0.028}$	$7.74_{0.23}$
150.0 _{1.0}	$2.222_{0.055}$	7.31 _{0.11}
250.0 _{1.0}	$4.56_{0.12}$	7.16 _{0.11}
$400.0_{1.6}$	13.68 _{0.39}	7.59 _{0.21}
500.0 _{2.0}	29.9 _{1.5}	$8.10_{0.22}$
600.0 _{2.4}	69.5 _{4.8}	8.77 _{0.18}
700.0 _{2.8}	174 ₁₅	$9.58_{0.20}$
800.0 _{3.2}	47544	10.57 _{0.41}
900.0 _{3.6}	1449_{177}	11.77 _{0.78}
1000.04.0	5040_{863}	13.2 _{1.1}

4. Discussion

Although the MiPPE viscosities for 224TMH are quantitatively reliable for $P \leq 600$ MPa, the percent deviation relative to experiment is approximately 300% at 1000 MPa. The rapid increase in η with respect to P also results in over prediction of the pressure-viscosity coefficient (α) at 900 MPa (11.77 compared to the IFPSC10 experimental value of 7.52). We suspect that this poor performance at the highest pressures is caused by the Mie

16-6 potential being overly repulsive at short distances [5, 16]. For this reason, MiPPE is more likely to predict super-Arrhenius behavior than traditional Lennard-Jones 12-6 force fields. Note that, as demonstrated in our previous study [16], we also expect the densities reported in Table 7 at high pressures to be significantly lower than experimental densities.

It is surprising that both the non-bonded and torsional parameter uncertainties are negligible compared to the numerical uncertainties in η (see Figure 4). A possible explanation for why the non-bonded parameter uncertainty has a negligible impact on η is that the CH₃ uncertainties are considerably smaller than those for CH₂, CH, and C (see Figure 3). As 224TMH consists primarily of CH₃ sites, the larger uncertainties in CH₂, CH, and C appear to not affect the results significantly.

By contrast, no clear explanation exists for why the torsional parameter uncertainties do not affect the overall uncertainty in η . Nieto-Draghi et al. suggest that a 15 % to 40 % increase in the torsional barriers should increase the viscosities appreciably for similar compounds [14, 15]. However, we do not observe such a strong dependence. Specifically, the required increase in the torsional barriers is between 80 % and 100 % to achieve an increase of approximately 10 % in viscosity (see Section S.V of Supplementary Material).

Other studies [4] correct for systematic errors in viscosity by normalizing η with respect to an experimental viscosity value at low pressure. This approach would be possible for the challenge compound since a single experimental data point is available at saturation pressure. Although this may provide a more accurate prediction, we prefer not to use an empirical correction, especially from a single data point. Our goal, rather, is to truly test the predictive capabilities of the MiPPE force field.

The slow system dynamics (i.e., long rotational relaxation times) at high pressures require extremely long simulations. Two attractive alternatives exist to enhance the configuration sampling by performing simulations at higher temperatures. The viscosity at the desired temperature is then obtained indirectly.

For example, the so-called time-temperature superposition method (TTS) combines

the shear rate dependent viscosity at different (higher) temperatures into a single master curve for estimating viscosity at any temperature [44]. Unfortunately, TTS is only applicable to NEMD simulations. An even simpler method, which is compatible with both NEMD and EMD, is to fit high-temperature viscosity values with an empirical model, e.g., the Vogel-Fulcher-Tammann-Hesse (VFTH) equation, and to then extrapolate to the desired temperature [4, 44]. Despite some obvious benefits, we are wary of the inordinately large uncertainties that this method can produce (see Figure 11 of Reference 4). Determining the existence of super-Arrhenius behavior necessitates manageable uncertainties at high pressures. Another clear disadvantage of both methods is the need to simulate at multiple temperatures, while IFPSC10 only requires viscosity estimates at 293 K. For these reasons, we choose the more direct brute-force approach.

5. Conclusions

Previous work demonstrated that the MiPPE force field provides reliable viscosities (typically within 10 %) for well-studied *n*-alkane and branched alkanes both at saturation and elevated pressures. For this reason, the MiPPE force field was chosen to predict the pressure-viscosity relationship of 2,2,4-trimethylhexane as part of the 10th Industrial Fluid Properties Simulation Challenge. In addition, we investigate the force field parameter uncertainty in the simulation results with Bayesian inference. Specifically, the non-bonded and torsional parameters are varied from run to run according to Markov Chain Monte Carlo. Surprisingly, the non-bonded and torsional parameter uncertainties are typically negligible compared to the numerical uncertainties, i.e., the fluctuations in simulation output and post-processing. Furthermore, we use cross-validation model selection to verify the existence of faster-than-exponential, a.k.a. super-Arrhenius, behavior at high pressures.

Near-quantitative agreement with experimental viscosity data (to within the combined uncertainties) is observed from atmospheric pressure to 600 MPa. At the highest pressures, however, the MiPPE force field dramatically over predicts the viscosity and

the pressure-viscosity coefficient. This suggests that the overly-repulsive Mie 16-6 non-bonded potential limits the pressure range over which the MiPPE force field is reliable.

Supplementary Material

Section S.I reports the IFPSC10 experimental data. Section S.II provides GROMACS input files. Section S.III describes how the CH and C non-bonded MCMC parameter sets are obtained. Section S.IV validates the MCMC-nb uncertainty quantification approach. Section S.V develops the $A_{\rm s}$ distribution used for the MCMC-nb-tors torsional parameters. Section S.VI provides tabulated values of the MCMC parameter sets. Section S.VII presents the average Green-Kubo integrals for each state point. Section S.VIII investigates the impact of the output frequency at high pressures.

Acknowledgments

We would like to acknowledge Jeffrey J. Potoff and Mohammad Soroush Barhaghi for their invaluable insights regarding the MiPPE force field. We are grateful for the internal review provided by Alta Y. Fang, Ian H. Bell, Andrei F. Kazakov, Daniel G. Friend, and Marcia L. Huber of the National Institute of Standards and Technology (NIST). We also appreciate the useful discussions and recommendations from Edward J. Maginn of The University of Notre Dame, Michael R. Shirts of The University of Colorado, J. Ilja Siepmann of The University of Minnesota, Jarod M. Younker of ExxonMobil, and John D. Chodera of the Memorial Sloan Kettering Cancer Center.

This research was performed while Richard A. Messerly held a National Research Council (NRC) Postdoctoral Research Associateship at NIST and while Michelle C. Anderson held a Summer Undergraduate Research Fellowship (SURF) position at NIST.

Commercial equipment, instruments, or materials are identified only in order to adequately specify certain procedures. In no case does such identification imply recommendation or endorsement by NIST, nor does it imply that the products identified are necessarily the best available for the intended purpose.

Partial contribution of NIST, an agency of the United States government; not subject to copyright in the United States.

References

References

- [1] Scott Bair, Laetitia Martinie, and Philippe Vergne. Classical EHL versus quantitative EHL: A perspective part II-super-Arrhenius piezoviscosity, an essential component of elastohydrodynamic friction missing from classical EHL. *Tribology Letters*, 63(3):37, 2016.
- [2] Christopher J. Mundy, Michael L. Klein, and J. Ilja Siepmann. Determination of the pressure-viscosity coefficient of decane by molecular simulation. *The Journal of Physical Chemistry*, 100(42):16779–16781, 1996.
- [3] Clare McCabe, Shengting Cui, Peter T. Cummings, Peter A. Gordon, and Roland B. Saeger. Examining the rheology of 9-octylheptadecane to giga-pascal pressures. *The Journal of Chemical Physics*, 114(4):1887–1891, 2001.
- [4] Pinzhi Liu, Hualong Yu, Ning Ren, Frances E. Lockwood, and Q. Jane Wang. Pressure-viscosity coefficient of hydrocarbon base oil through molecular dynamics simulations. *Tribology Letters*, 60(3):34, 2015.
- [5] Richard A. Messerly, Michelle C. Anderson, S. Mostafa Razavi, and J. Richard Elliott. Improvements and limitations of Mie λ -6 potential for prediction of saturated and compressed liquid viscosity. *Fluid Phase Equilibria*, Pending publication, 2018.
- [6] M. G. Martin and J. I. Siepmann. Transferable potentials for phase equilibria. 1. United-atom description of *n*-alkanes. *The Journal of Physical Chemistry B*, 102(14):2569–2577, 1998.
- [7] Marcus G. Martin and J. Ilja Siepmann. Novel configurational-bias Monte Carlo method for branched molecules. Transferable potentials for phase equilibria. 2.

- United-atom description of branched alkanes. *The Journal of Physical Chemistry B*, 103(21):4508–4517, 1999.
- [8] Mansi S. Shah, J. Ilja Siepmann, and Michael Tsapatsis. Transferable potentials for phase equilibria. Improved united-atom description of ethane and ethylene. *AIChE Journal*, 63(11):5098–5110, 2017.
- [9] Andrea Hemmen and Joachim Gross. Transferable anisotropic united-atom force field based on the Mie potential for phase equilibrium calculations: *n*-alkanes and *n*-olefins. *The Journal of Physical Chemistry B*, 119(35):11695–11707, 2015.
- [10] Dominik Weidler and Joachim Gross. Transferable anisotropic united-atom force field based on the Mie potential for phase equilibria: Aldehydes, ketones, and small cyclic alkanes. *Industrial & Engineering Chemistry Research*, 55(46):12123–12132, 2016.
- [11] J. J. Potoff and D. A. Bernard-Brunel. Mie potentials for phase equilibria calculations: Applications to alkanes and perfluoroalkanes. *The Journal of Physical Chemistry B*, 113(44):14725–14731, 2009.
- [12] Jason R. Mick, Mohammad Soroush Barhaghi, Brock Jackman, Loren Schwiebert, and Jeffrey J. Potoff. Optimized Mie potentials for phase equilibria: Application to branched alkanes. *Journal of Chemical & Engineering Data*, 62(6):1806–1818, 2017.
- [13] Philippe Ungerer, Christele Beauvais, Jerome Delhommelle, Anne Boutin, Bernard Rousseau, and Alain H. Fuchs. Optimization of the anisotropic united atoms intermolecular potential for *n*-alkanes. *The Journal of Chemical Physics*, 112(12):5499–5510, 2000.
- [14] Carlos Nieto-Draghi, Philippe Ungerer, and Bernard Rousseau. Optimization of the anisotropic united atoms intermolecular potential for *n*-alkanes: Improvement of transport properties. *The Journal of Chemical Physics*, 125(4):044517, 2006.

- [15] Carlos Nieto-Draghi, Anthony Bocahut, Benoît Creton, Pascal Have, Aziz Ghoufi, Aurélie Wender, , Anne Boutin, Bernard Rousseau, and Laurent Normand. Optimisation of the dynamical behaviour of the anisotropic united atom model of branched alkanes: application to the molecular simulation of fuel gasoline. *Molecular Simulation*, 34(2):211–230, 2008.
- [16] Richard A. Messerly, Michael R. Shirts, and Andrei F. Kazakov. Uncertainty quantification confirms unreliable extrapolation toward high pressures for united-atom Mie λ -6 force field. *The Journal of Chemical Physics*, 149(11):114109, 2018.
- [17] F. Rizzi, H. N. Najm, B. J. Debusschere, K. Sargsyan, M. Salloum, H. Adalsteinsson, and O. M. Knio. Uncertainty quantification in MD simulations. Part II: Bayesian inference of force-field parameters. *Multiscale Modeling & Simulation*, 10(4):1460–1492, 2012.
- [18] Panagiotis Angelikopoulos, Costas Papadimitriou, and Petros Koumoutsakos. Bayesian uncertainty quantification and propagation in molecular dynamics simulations: A high performance computing framework. *The Journal of Chemical Physics*, 137(14):144103, 2012.
- [19] Fabien Cailliez and Pascal Pernot. Statistical approaches to forcefield calibration and prediction uncertainty in molecular simulation. *The Journal of Chemical Physics*, 134(5):054124, 2011.
- [20] Richard A. Messerly, Thomas A. Knotts IV, and W. Vincent Wilding. Uncertainty quantification and propagation of errors of the Lennard-Jones 12-6 parameters for *n*-alkanes. *The Journal of Chemical Physics*, 146(19):194110, 2017.
- [21] B. Efron. Bootstrap methods: Another look at the jackknife. *The Annals of Statistics*, 7(1):1–26, 1979.
- [22] Peter A. Gordon. Development of intermolecular potentials for predicting transport properties of hydrocarbons. *The Journal of Chemical Physics*, 125(1):014504, 2006.

- [23] Carlos Braga and Karl P. Travis. Computer simulation of the role of torsional flexibility on mass and momentum transport for a series of linear alkanes. *The Journal of Chemical Physics*, 137(6):064116, 2012.
- [24] Carmelo Herdes, Tim S. Totton, and Erich A. Müller. Coarse grained force field for the molecular simulation of natural gases and condensates. *Fluid Phase Equilibria*, 406:91 100, 2015.
- [25] M. P. Allen and D. J. Tildesley. *Computer simulation of liquids*. Oxford University Press, Oxford England, second edition, 2017.
- [26] M.J. Abraham, D. van der Spoel, E. Lindahl, B.Hess, and the GROMACS development team. *GROMACS User Manual version 2018, www.gromacs.org* (2018).
- [27] Berk Hess, Henk Bekker, Herman J. C. Berendsen, and Johannes G. E. M. Fraaije. LINCS: A linear constraint solver for molecular simulations. *Journal of Computational Chemistry*, 18(12):1463–1472, 1998.
- [28] Berk Hess. P-LINCS: A parallel linear constraint solver for molecular simulation. *Journal of Chemical Theory and Computation*, 4(1):116–122, 2008.
- [29] Alison N. Leonard, Andrew C. Simmonett, Frank C. Pickard, Jing Huang, Richard M. Venable, Jeffery B. Klauda, Bernard R. Brooks, and Richard W. Pastor. Comparison of additive and polarizable models with explicit treatment of long-range Lennard-Jones interactions using alkane simulations. *Journal of Chemical Theory and Computation*, 14(2):948–958, 2018.
- [30] Kang-Sahn Kim, Myung Hoon Han, Changho Kim, Zhen Li, George Em Karniadakis, and Eok Kyun Lee. Nature of intrinsic uncertainties in equilibrium molecular dynamics estimation of shear viscosity for simple and complex fluids. *The Journal of Chemical Physics*, 149(4):044510, 2018.
- [31] R.W Hockney, S.P Goel, and J.W Eastwood. Quiet high-resolution computer models of a plasma. *Journal of Computational Physics*, 14(2):148 158, 1974.

- [32] William C. Swope, Hans C. Andersen, Peter H. Berens, and Kent R. Wilson. A computer simulation method for the calculation of equilibrium constants for the formation of physical clusters of molecules: Application to small water clusters. *The Journal of Chemical Physics*, 76(1):637–649, 1982.
- [33] Giovanni Bussi, Davide Donadio, and Michele Parrinello. Canonical sampling through velocity rescaling. *The Journal of Chemical Physics*, 126(1):014101, 2007.
- [34] William G. Hoover. Canonical dynamics: Equilibrium phase-space distributions. *Physical Review A*, 31:1695–1697, 1985.
- [35] Shuichi Nosé. A unified formulation of the constant temperature molecular dynamics methods. *The Journal of Chemical Physics*, 81(1):511–519, 1984.
- [36] H. J. C. Berendsen, J. P. M. Postma, W. F. van Gunsteren, A. DiNola, and J. R. Haak. Molecular dynamics with coupling to an external bath. *The Journal of Chemical Physics*, 81(8):3684–3690, 1984.
- [37] Shuichi Nosé and M.L. Klein. Constant pressure molecular dynamics for molecular systems. *Molecular Physics*, 50(5):1055–1076, 1983.
- [38] M. Parrinello and A. Rahman. Polymorphic transitions in single crystals: A new molecular dynamics method. *Journal of Applied Physics*, 52(12):7182–7190, 1981.
- [39] Edward J. Maginn, Richard A. Messerly, Daniel J. Carlson, Daniel R. Roe, and J. Richard Elliott. Best practices for computing transport properties 1. Self-diffusivity and viscosity from equilibrium molecular dynamics v1.

 Living Journal of Computational Molecular Science, Pending publication, 2018.

 https://github.com/ejmaginn/TransportCheckList.
- [40] Yong Zhang, Akihito Otani, and Edward J. Maginn. Reliable viscosity calculation from equilibrium molecular dynamics simulations: A time decomposition method. *Journal of Chemical Theory and Computation*, 11(8):3537–3546, 2015.

- [41] C. Barus. Isothermals, isopiestics and isometrics relative to viscosity. *American Journal of Science*, 45:87–96, 1893.
- [42] C.J.A. Roelands. *Correlational Aspects of the Viscosity-Temperature-Pressure Relationships of Lubricating Oils*. PhD thesis, TU Delft, Delft University of Technology, 1966.
- [43] Michael Frenkel, Robert D. Chirico, Vladimir Diky, Xinjian Yan, Qian Dong, and Chris Muzny. ThermoData Engine (TDE): Software implementation of the dynamic data evaluation concept. *Journal of Chemical Information and Modeling*, 45(4):816–838, 2005.
- [44] Fardin Khabaz and Rajesh Khare. Molecular simulations of asphalt rheology: Application of time-temperature superposition principle. *Journal of Rheology*, 62(4):941–954, 2018.