Molecular Dynamics Simulation Studies of Viscosity and Diffusion of *n*-Alkane Oligomers at High Temperatures

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In this paper we have carried out molecular dynamics simulations (MD) for model systems of liquid *n*-alkane oligomers (C_{12} - C_{80}) at high temperatures (~2300 K) in a canonical ensemble to calculate viscosity η , self-diffusion constants D, and monomeric friction constant ζ . We found that the long chains of these *n*-alkanes at high temperatures show an abnormality in density and in monomeric friction constant. The behavior of both activation energies, E_{η} and E_{D} , and the mass and temperature dependence of η , D, and ζ are discussed.

Key Words : MD simulation, Diffusion, Viscosity, Friction, n-Alkane

Introduction

A number of studies have been reported on the relationship between the zero shear viscosity (η_0) and molecular weight. For linear polymers, η_0 increases linearly with the molecular weight M.¹ At values of molecular weight greater than a critical molecular weight, M_c , η_0 becomes proportional to M^{3.4} proposed in 1951 by Fox and Flory² on the basis of measurements on narrow distribution fractions of polystyrene and polyisobutylene. Self-diffusion constant (D) of polymer melts have also shown a similar trend: $D \sim M^{-1}$ for $M < M_c$ and $D \sim M^{-2}$ for $M > M_c$. M_c is called the entanglement coupling molecular weight. Below and above M_c , the polymer chain dynamics of unentangled and entangled chains are commonly described by the Rouse model and the reptation model.^{1,3,4}

At the molecular weight below the Rouse regime, D and η of *n*-alkanes also show power law behaviors. For example, D of *n*-alkanes from *n*-octane to polyethylene of the molecular weight of several thousands was reported to follow the relation, $D \sim M^{-\alpha}$ in which the exponents α is in the range of 2.72-1.85 depending on temperature.⁵⁻⁷ Although the power law dependence has the same functional form as in the Rouse and reptation regimes, the origin is totally different. The molecular weight dependence of D and η in the Rouse and reptation regimes is attributed to the topological entanglement effect not to the local segmental friction while the exponent found below the Rouse regime reflects the molecular weight dependence of the local friction rather than the topological effect.

While the Rouse-reptation crossover has received considerable attention in recent years,^{8,9} the *n*-alkane-Rouse crossover has received relatively little attention. The equilibrium (EMD) and non-equilibrium molecular dynamics (NEMD) simulations¹⁰⁻¹² on the system are useful for the study on the transport properties of liquid *n*-alkanes of moderate molecular sizes (C_{10} - C_{40}), which can shed light on the polymer dynamics in the molecular level.

In this study, we use the Green-Kubo and Einstein formulas

for the calculation of the viscosities and diffusion constants of liquid n-C₁₂H₂₆ to n-C₈₀H₁₆₂ using EMD simulation. The main purpose of this study is to characterize the viscosity and self-diffusion dynamics of small *n*-alkanes as a function of chain length n (or molecular weight M) especially at high temperatures.

Molecular Models and MD Simulation Methods

For liquid *n*-alkanes, we have chosen 7 systems - *n*-dodecane($C_{12}H_{26}$), *n*-eicosane($C_{20}H_{42}$), *n*-octaconsane($C_{28}H_{58}$), *n*-hexatriacontane($C_{36}H_{74}$), and *n*-tetratetracontane($C_{44}H_{90}$), *n*-hexacontane($C_{60}H_{122}$), and *n*-octacontane($C_{80}H_{162}$). Each simulation was carried out in the NpT ensemble to determine the volume of each system at given temperatures, and after the equilibrium density and hence the length of cubic simulation box were obtained, new NVT MD simulation was performed for each system to store the configurations of *n*-alkanes for later analyses. The usual periodic boundary condition in the x-, y-, and z-directions and the minimum image convention for pair potential were applied. Gaussian isokinetics was used to keep the temperature of the system constant.^{13,14}

We used a united atom (UA) model for *n*-alkanes, that is, methyl and methylene groups are considered as spherical interaction sites centered at each carbon atom. This model have been used in the previous simulation studies.¹⁰⁻¹² Here, we briefly describe the salient features of the model. The interaction between the sites on different *n*-alkane molecules and between the sites separated by more than three bonds in the same *n*-alkane molecule is described by a Lennard-Jones (LJ) potential. All the sites in a chain have the same LJ size parameter $\sigma_i \equiv \sigma_{ii} \equiv 3.93$ A, the well depth parameters were $\epsilon_i \equiv \epsilon_{ii} \equiv 0.94784 \text{ kJ/mol}$ for interactions between the end sites and $\epsilon_i \!\!=\!\! 0.39078$ kJ/mol for interactions between the internal sites. The Lorentz-Berthelot combining rules $[\varepsilon_{ij} = (\varepsilon_i \varepsilon_j)^{1/2}]$ $\sigma_{ii} \equiv (\sigma_i + \sigma_i)/2$ were used for interactions between an end site and an internal site. A cut-off distance of $2.5\sigma_i$ was used for all the LJ interactions.

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Initially in NpT MD simulations, the bond-stretching was described by a harmonic potential, with an equilibrium bond distance of 1.54 A and a force constant of 1882.8 kJ/mol/A². The bond bending interaction was also described by a harmonic potential with an equilibrium angle of 114° and a force constant of 0.079187 kJ/mol/degree.² The torsional interaction was described by the potential developed by Jorgensen *et al.*¹⁵

$$U_{\text{torsion}}(\phi) = a_0 + a_1 \cos\phi + a_2 \cos^2\phi + a_3 \cos^3\phi, \quad (1)$$

where ϕ is the dihedral angle, and $a_0 = 8.3973 \text{ kJ/mol}$, $a_1 =$ 16.7862 kJ/mol, $a_2 = 1.1339$ kJ/mol, and $a_3 = -26.3174$ kJ/ mol. For the time integration of the equations of motion, we adopted Gear's fifth-order predictor-corrector algorithm¹⁶ with a time step of 2 femto-second for all the systems. Later in NVT MD simulations, the bond-stretching was switched to a constraint force which keeps intramolecular nearest neighbors at a fixed distance. The advantage for this change is to increase the time step as 5 femto-seconds with the use of RATTLE algorithm¹⁷ and velocity Verlet algorithm.¹⁸ After a total of 1,000,000 time steps (5 nano-seconds) for equilibration, the equilibrium properties were then averaged over 5 blocks of 200,000 time steps (1 nano-seconds). The configurations of all the molecules for further analyses were stored every 10 time steps (0.05 pico second) which is small enough for the tick of any time auto-correlation functions.

Self-diffusion constant is obtained from the Einstein relation of the mean square displacement (MSD):¹⁹

$$\mathbf{D} = \frac{1}{6} \lim_{t \to \infty} \frac{d}{dt} < [\mathbf{r}_i(t) - \mathbf{r}_i(0)]^2 >, \qquad (2)$$

The Green-Kubo relation for the viscosity is given in terms of the pressure auto-correlation function by

$$\eta = \frac{V}{kT} \int_{0}^{\infty} dt < P_{\alpha\beta}(0) P_{\alpha\beta}(t) >, \qquad (3)$$

where $\alpha\beta = xy$, xz, yx, yz, zx, or zy and for a system of N molecules, the molecular formalism for the pressure tensor is given by

$$P_{\alpha\beta} = \frac{1}{V} \left[\sum_{i} m_{i} v_{i\alpha} v_{i\beta} + \sum_{i} \sum_{j>i} (r_{i\alpha} - r_{j\alpha}) F_{ij\beta} \right] \quad , \qquad (4)$$

where the indices i and j refer to molecules i and j, and F_{ij} is the force between the centers of mass of molecules i and j.

Friction constant is obtained from the time integral of the force auto-correlation (FAC) function^{20,21}:

$$\zeta = \frac{1}{3kT} \int_0^\tau dt \langle f_i(0) \cdot f_i(t) \rangle$$
(5)

where $\mathbf{f}_i(t) = \mathbf{F}_i(t) - \langle \mathbf{F}_i(t) \rangle$ and $\mathbf{F}_i(t)$ is the total force exerted on molecule *i*. Molecular friction force is defined by $\zeta_M = 1/\sigma_r = \zeta/M = \zeta/nm$ with M molecular mass of *n*-alkane, n number of carbons, m mass of monomer and τ_r the macroscopic relaxation time of the FAC.²¹ **Results and Discussion**

Density. In Figure 1, we plot the density of *n*-alkanes as a function of temperature. The values of the density were obtained from a long time of NpT MD simulations as discussed above. Generally the density decreases linearly with increasing temperature as expected, but an abnormality is found in the long *n*-alkanes at high temperatures. For the short *n*-alkanes, we observed that the volumes of the systems increase infinitely as the temperature increases. This indicates "evaporation" of the systems. However, for the long chain *n*-alkane, the density loses the linear decrement as temperature increases further and becomes flat at high temperatures. This may indicates the existence of a boundary between the small molecular melts and the Rouse regime. In this study, the crossover is between C_{44} and C_{60} .

Self-diffusion Constant. Self-diffusion constants of n-alkanes, D, obtained from the Einstein relation, Eq. (2), are plotted in the inset of Figure 1 as a function of T. The dependence of D on T in the small molecular melts is obviously different from that in the Rouse regime.

The temperature dependence of the calculated diffusion constants of liquid *n*-alkanes over the whole temperatures considered is suitably described by an Arrhenius plot as shown in Figure 2:

$$D = D_0 \exp(-E_D/RT), \tag{6}$$

where D_0 is the pre-exponential factor, RT has the usual meaning, and E_D is the activation energy of *n*-alkane diffusion. The value of the activation energy is a direct measure of the temperature dependence of self-diffusion constant. The activation energies obtained from the slope of the least square fit are 2.83, 3.41, 3.84, 4.49, 4.74, 4.87, and 4.95 kcal/mol for C₁₂, C₂₀, C₂₈, C₃₆, C₄₄, C₆₀, and C₈₀, respectively. The previous MD simulation study²² reported this value as 3.98 kcal/mol for the UA model of *n*-tetracosane (C₂₄) which





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Figure 2. Arrhenius plot of D (10^{-5} cm²/sec) vs. 1/T.

is compared with a value of 4.04 kcal/mol obtained from viscosity measurements²³ and a value of 4.36 from PFG NMR results.²⁴ The values of E_D are plotted in Figure 3. As chain length n increases the increment of E_D decreases, and it is expected to approach an asymptotic value as n increases further. It was reported that E_D increases linearly with log M from 2.32 kcal/mol for *n*-heptane to 5.81 kcal/mol for *n*-hexacontane (C₆₀).^{7,25} Fleisher⁵ also determined E_D to be about 4.8 kcal/mol for several polyethylene independent of molecular weight from 9 to 52.7 kg/mol, from which we can deduce that E_D of *n*-alkane reaches an asymptotic limit over C₁₀₀.

We also show the log-log plot of diffusion constant versus molecular mass in Figure 4. The obtained exponents are between -2.36 and -1.44. Recent experimental study report-



Figure 3. Activation energies of D (circle) and η (square) vs. chain length n.



Figure 4. A log-log plot of D (10^{-5} cm²/sec) vs. M (g/mol).

ed that $D \sim M^{-\alpha}$, with a changing approximately linearly from 2.72 to 1.85 as T increases.⁷ Thus the apparent activation energy also rises linearly with log M. In the absence of molecular entanglements, Rouse kinetics predicts $\alpha = -1$, but Cohen-Tumbull-Bueche free-volume effects due to molecular chain ends add a further nonpower-law term,²⁶ enhancing D increasingly at low M.

Viscosity. Viscosities of *n*-alkanes, η , are obtained from the Green-Kubo relation. It is worth note that the pressure auto-correlation functions of long chains of *n*-alkanes at low temperatures have the non-decaying long-time tails (not shown) and that it is impossible to get the plateau values in the running time integrals of the pressure auto-correlation functions. For example, in this study, C₆₀ at 318 K and C₈₀ at 318 K and 418 K. The temperature dependence of the calculated viscosities of liquid *n*-alkanes over the temper-



Figure 5. Arrhenius plot of η (cP) vs. 1/T.



Figure 6. A log-log plot of η (cP) vs. M (g/mol).

atures is also suitably described by an Arrhenius plot except very high temperatures as shown in Figure 5:

$$\eta = C_0 \exp(E_{\eta}/RT) \tag{7}$$

where C_0 is the pre-exponential factor and E_{η} is the activation energy of *n*-alkane viscosity. But the experimental result for the linear polyethylene⁶ showed that the lines are slightly curved, indicating that the temperature dependence, as in most polymers, does not follow the Arrhenius equation. The activation energies obtained from the slope of the least square fit except T over 1318 K are 3.71, 4.77, 6.29, 6.72, 7.17, 7.36, and 7.49 kcal/mol for C12, C20, C28, C36, C44, C60, and C₈₀, respectively and are plotted in Figure 3. It was experimentally reported for n-alkanes and linear polyethylene⁶ that the activation energy increases with chain length and at the highest molecular weight tested (M \approx 4.4 kg/mol) the activation energy reaches 6.6 kcal/mol, which is similar to the average value found for NBS 1482-4 (6.7 kcal/mol) and the values reported by others^{27,28} for high molecular weight linear polyethylene (6.1-6.9 kcal/mol).

Figure 6 shows the log-log plot of viscosity versus molecular mass. The obtained exponents are between 2.41-3.32 except C₁₂ (4.32). The experimental results for *n*-alkanes and linear polyethylene⁶ show that η is well described by the power law $\eta \sim M^{1.8}$ at low molecular weight (M < 5 kg/mol) and $\eta \sim M^{3.64}$ at high molecular weight (M > 5 kg/mol) at 448 K. The molecular weight of the *n*-alkanes tested in this study is much lower than this experimental range of molecular weight, but the lowest exponent gives 2.41 at 818 K.

Friction Constant. It is well known that the calculation of friction constant from the force auto-correlation function is fairly hard due to the non-decaying long-time tails. As Kubo pointed out in his "fluctuation-dissipation theorem",¹⁹ the correlation function of random force R will decay in a time interval of τ_c (microscopic time or collision duration time), whereas that of the total force F has two parts, the short time

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Figure 7. Friction constants, ζ (10³ g/ps·mol) *vs.* T. The inset shows plot of ζ/n *vs.* T.



Figure 8. Friction constants, ζ (10³ g/ps·mol) *vs.* chain length *n*. The inset shows plot of ζ/n *vs. n*. From top, T = 318, 418, 518, 618, 718, 818, 918, 1118 and 1318 K, respectively.

or the fast part similar to that of the random force and the slow part which should just cancel the fast part in the time integration.²⁹ This means that the time integral of Eq. (4) up to $\tau = \infty$ is equal to zero.

The time integral in Eq. (4) attains a plateau value for τ satisfying $\tau_c \ll \tau \ll \tau_r$, if the upper limit of the time integral, Eq. (4), is chosen that $\tau_c \ll \tau \ll \tau_r$ because the slow tail of the correlation function is cut off. However, we were unable to get the plateau value in the running time integral of the force auto-correlation function. Kubo suggested that the friction constants should be obtained from the random FAC function not from the total FAC and that there exists a difficulty to separate the random force part from the

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total force. We could obtain the friction constants by the time integral of the total FAC with choosing the upper limit of τ as the time that the FAC has the first negative value by assuming that the fast random force correlation ends at that time.

Figure 7 shows the plot of ζvs . T for given *n*-alkanes. The calculated friction constants give a correct qualitative trend: ζ decreases with increasing T. However, the abnormality in density of the long *n*-alkanes at high temperatures is recovered in the friction constant. For the short *n*-alkanes, ζ is inversely proportional to T but for the long chain *n*-alkane, the density loses the inverse decrement as temperature increases further and becomes flat at high temperatures. When ζ/n is plotted as a function of T as shown in the inset of Figure 7, the overall behavior of ζ/n for all *n*-alkanes is rather coincided except at high temperatures.

We plot ζ *vs. n* in Figure 8 in which ζ increases linearly with increasing *n* for a given T. The slope of ζ *vs. n* decreases with increasing T. An atomistic MD simulation of linear polyethylene melts³⁰ reported that the friction constant extracted from D by applying the Rouse model is seen to increase from value of short *n*-alkanes to a chain-lengthindependent plateau one, reached in a region of n = 60-80. The linear increase of ζ with increasing *n* as shown in Figure 8 might indicate that the liquid *n*-alkanes over the whole temperatures considered are far away from the Rouse regime. When ζ/n is plotted as a function of *n* as shown in the inset of Figure 8, ζ/n is almost independent of *n* for given temperatures, which indicates the friction is governed by the local segmental friction not by the topological entanglement effect.

Conclusion

We have presented the results for viscosity and selfdiffusion constants of model systems for liquid *n*-alkane oligomers (C_{12} ~ C_{80}) at high temperatures (~2300 K) using molecular dynamics (MD) simulations in a canonical ensemble. The long chains of these *n*-alkanes at high temperatures show an abnormality in density. Calculated viscosity η and self-diffusion constants D are comparable with experimental results and the temperature dependence of both η and D is suitably described by the Arrhenius plot. The behavior of both activation energies, E_n and E_D , indicates that the activation energies approach asymptotic values with increasing chain length, which is consistent with the experimental observation. The calculated monomeric friction constant ζ shows an abnormality again as in density. The slope of ζ vs. *n* decreases with increasing T but does not provide a chain-length-independent plateau.

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