The role of lubricant molecular shape in microscopic friction

Oleg M. Braun
Institute of Physics, National Academy of Sciences of Ukraine, 03028 Kiev, Ukraine

Nicola Manini
Dipartimento di Fisica, Università di Milano, Via Celoria 16, 20133 Milano, Italy

Erio Tosatti
International School for Advanced Studies (SISSA), Via Beirut 2-4, I-34014 Trieste, Italy
CNR-INFM Democritos National Simulation Center, Via Beirut 2-4, I-34014 Trieste, Italy and
International Centre for Theoretical Physics (ICTP), P.O. Box 586, I-34014 Trieste, Italy

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With the help of a simple two-dimensional model we simulate the tribological properties of a thin lubricant film consisting of linear (chain) molecules in the ordinary soft-lubricant regime. We find that friction generally increases with chain length, in agreement with their larger bulk viscosity. When comparing the tribological properties of molecules which stick bodily to the substrates with others carrying a single sticking termination, we find that the latter generally yield a larger friction than the former.

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I. INTRODUCTION

Understanding sliding friction between substrates separated by a thin lubricant film is technologically crucial but also rich physically [1, 2]. Without lubricant, it is known that the lowest friction can be achieved for a contact of two hard crystalline solids, when the atomically flat surfaces are incommensurate (e.g., see [2] and references therein). When the surfaces are separated by a lubricant film, the lowest friction is achieved if the film is solid and crystalline but incommensurate with the substrate surface. Such a situation appears in the so-called “hard” lubricant system [2], where the intermolecular interaction strength within the lubricant, $V_{ll}$, is stronger than the lubricant-substrate interaction, $V_{sl}$. In that case, the shape of lubricant molecules is not too relevant, because friction is determined uniquely by the substrate-lubricant interface structure.

Conventional lubricants, however, belong typically to the opposite “soft” lubricant type, with $V_{ll} < V_{sl}$. Here each substrate is covered by a glued lubricant monolayer, protecting it from wear. Sliding occurs deeper inside the lubricant film, where below the film melting point a typical melting-freezing mechanism operates: the lubricant locally melts at slip (i.e., during sliding) and re-solidifies at stick [3]. For a thick lubricant film above its melting point, e.g., and for thicknesses larger than say ten molecular diameters, friction is proportional to the bulk viscosity of the lubricant. Therefore, thick fluid lubricants consisting of simple spherical molecules or individual atoms, characterized by a low bulk viscosity, would provide the best frictional properties. Indeed, experiments show that the bulk viscosity for linear molecules grows with the molecule length as was shown in Ref. [4]. One could then expect that the friction coefficient of chain lubricant molecules should grow with their molecular length.

However, it is well known that long molecules often make better lubricants. The reason for this is the squeeze-out effect: at high load, when the regime of boundary lubrication is approached and the film is locally extremely thin, at stick the lubricant may be squeezed out in correspondence of the substrate asperities. Direct contact of un lubricated asperities leads to a high static friction force $f_s$ and generally to substrate wear. Sivebaek et al. [4] showed that with the increase of the alkane chain length, transitions from $N_l$ to $N_l - 1$ lubricant layers occurs at higher pressure. Longer lubricant molecules are harder to squeeze out, and this is why they provide better lubrication properties. Following that line, Ref. [4] studied the “totally-glued” case, where all atoms of the lubricant molecule interact strongly with the substrates. Longer molecules adhere more to the substrates, making it harder for the squeeze-out of the last lubricant layer to nucleate. In agreement with that, longer alkanes perform better as boundary lubricants than shorter ones, because they prevent more effectively the appearance of cold-welded junctions and the resulting wear.

Tribological properties of lubricants constructed of linear (chain) molecules of length $L$, again for the totally-glued system, were also studied with the help of molecular dynamics (MD) simulation by He and Robbins [5]. They found that at the small coverage $\theta = M/N_s = 1/8$ (the number of lubricant molecules $M$ divided by the number of substrate surface atoms $N_s$) the dependence of the static friction on chain length (for $L = 1, 3, 6$) is small, while for a higher coverage, $M/N_s = 1/2$, chains with $L = 3$ and $6$ show nearly the same behavior, while “monoatomic molecules” ($L = 1$) display a dramatically
The paper is organized as follows: Sec. II introduces our minimal model; Sec. III provides the mathematical and numerical details of the model’s solution; Sec. IV contains a detailed analysis of the results, which are finally discussed in Sec. V.

II. THE MODEL

We use a 2D model, where point particles, representing individual atoms or monomer units of larger molecules, can move in two dimensions \( x \) and \( z \), where \( x \) is the sliding direction and \( z \) is perpendicular to the substrates. The substrates are modeled as two rigid chains of \( N_s \) atoms and lattice spacing \( R_s \), so that the total mass of each substrate is \( N_s m_s \) and the system size in the sliding direction is \( L_z = N_s R_s \). We apply periodic boundary condition along the \( x \) direction. The bottom substrate is fixed at \( x = z = 0 \), while the top one is free to move in both \( x \) and \( z \) directions. The top substrate is pressed toward the bottom one by a \((-z)\)-directed constant load force \( F_l = N_s f_l \), and is driven in the \( x \) direction at velocity \( v_s \) through an attached spring of elastic constant \( N_s k_s \), as sketched in Fig. 1. The spring force \( F \), which works against the friction force, is monitored during simulation (throughout the paper we report the force per substrate atom \( f = F/N_s \)). Thus, our model is a 2D variant of a typical experimental setup used in tribology [1, 2].

Between the substrates we insert \( N \) lubricant atoms of mass \( m_l \). How eventually these \( N \) atoms are to be lumped together to form linear molecules will be detailed below. Independently of that, all atoms interact via pairwise 12-6 Lennard-Jones (LJ) potential

\[
V_{\text{LJ}}(r) = V_{\alpha\alpha'} \left[ \left( \frac{R_{\alpha\alpha'}}{r} \right)^{12} - 2 \left( \frac{R_{\alpha\alpha'}}{r} \right)^{6} \right].
\]

Here \( \alpha, \alpha' = s \) or \( l \) for the substrate or lubricant atoms respectively. Thus, the lubricant-lubricant interaction is characterized by the parameters \( V_{\text{ll}} \) and \( R_{\text{ll}} \), while the lubricant-substrate interaction by \( V_{\text{sl}} \) and \( R_{\text{sl}} \) (direct interaction between the top and bottom substrates is omitted). Throughout the paper we will use dimensionless units, where \( m_s = m_l = 1 \), \( R_{\text{ll}} = 1 \), and the energy parameters \( V_{\alpha\alpha'} \) takes values around \( V_{\alpha\alpha'} \sim 1 \). The relative strength of the energy parameters \( V_{\text{ll}} \) and \( V_{\text{sl}} \) determines the low-temperature lubricant behavior [2]. The lubricant is “hard” (i.e., at low temperature it remains solid at slip) when \( V_{\text{ll}} \gtrsim V_{\text{sl}} \), and “soft” (i.e., the film melts at least locally during slips) for \( V_{\text{ll}} \ll V_{\text{sl}} \). Because a 2D model cannot reproduce even qualitatively the true phonon spectrum of a 3D system, and because moreover frictional kinetics is generally diffusional rather than inertial, we use Langevin equations of motion with a Gaussian random force corresponding to a given temperature \( T \), and a damping force

\[
\begin{align*}
  f_{\eta,x} &= -m_l \eta(z) \dot{z} - m_l \eta(z - x) (\dot{x} - \dot{X}), \\
  f_{\eta,z} &= -m_l \eta(z) \dot{z} - m_l \eta(z - x) (\dot{z} - \dot{Z}).
\end{align*}
\]

Here \( x, z \) are the coordinates of a generic lubricant atom and \( X, Z \) is the center-of-mass coordinate of the top substrate. The viscous damping coefficient is assumed to decrease exponentially with the distance from the corre-
sponding substrate,
\[ \eta(z) = \eta_0 \left(1 - \tanh \left(\frac{z}{z_d}\right)\right), \]
where we typically use \( \eta_0 = 1 \) and \( z_d = 0.1 \). This form is meant to mimic the Joule heat dissipation, which can only take place through the substrates, and therefore is mostly effective at the interfaces. The Gaussian width of the Langevin random force equals \( 2\eta(z) m_1 T^{1/2} \).

We compare the main tribological properties of lubricants consisting of molecules of different lengths, from individual (atom-like) monomers to linear chain molecules composed of \( L \) atomic/monomeric units, so that the number of lubricant molecules is \( M = N/L \). To build chain molecules, we follow the procedure due to Robbins et al.\(^2\)\(^3\)\(^8\). The lubricant is described with the Kremer-Grest bead-spring model\(^9\), which yields realistic dynamics for polymer melts\(^10\). While all monomers within a molecule still interact with each other via the LJ potential, Eq. (1), adjacent monomers inside a given molecule (chain) interact via an additional (“FENE”) potential\(^2\)\(^3\),
\[ V_{\text{CH}}(r) = \begin{cases} -\tilde{V}_{\text{CH}} \ln \left[1 - \left(\frac{r}{R_{\text{CH}}}\right)^2\right] & \text{for } r < R_{\text{CH}} \\ +\infty & \text{for } r \geq R_{\text{CH}}, \end{cases} \]
where \( R_{\text{CH}} = 1.336 R_{ll} \) and \( \tilde{V}_{\text{CH}} = 33.75 V_{ll} \) (these parameters are the same as in Ref.\(^9\)). This additional potential has the main effect of preventing molecular breaking. Besides that, it is fairly small at the equilibrium distance \( r = R_{ll} \) of \( V_{ll} \), and causes a bond-length contraction to \( R_{\text{mol}} \approx 0.856 R_{ll} \).

### III. SIMULATION TECHNIQUE

Our frictional simulations use periodic boundary conditions throughout, a choice which as usual minimizes size effects. Results are generally dependent on the total number of lubricant atoms \( N \). If \( N \) does not match exactly the exact number of atoms required for an integer number of closely packed layers, then the extra interstitial atoms or vacancies constitute structural defects. In real life the presence of these defects is of course the rule rather than the exception, but in the large interface size of an actual experiment their importance is more marginal than in our finite-size simulation. To reduce errors due to these misfit defects, we use a geometry with a slightly curved top substrate, whose z-coordinate varies along the x direction by
\[ z = Z + \frac{1}{2} R_{sl} \left[1 - \cos \frac{2\pi(x - X)}{L_x}\right], \]
as illustrated in Fig. \( \mathbf{1} \).

In the simulations to be presented below, each substrate contains \( N_s = 62 \) rigid atoms and the substrate lattice constant is chosen as \( R_s = 2/3 \), moderately

\begin{align*}
\text{“incommensurate” with the lubrication equilibrium interatomic distance } R_{ll} = 1, \text{ a feature not atypical for lubricant/substrate interfaces.}
\end{align*}

Accordingly, the system size is \( L_x = 41.3 \). This fits up to \( N \lesssim 50 \) atoms (in the atomic lubricant case) in one monolayer film under a standard applied load \( f_l = 1 \). The \( N = 100 \) atoms therefore complete two monolayers (one glued to the bottom substrate, the other to the top substrate). Atoms in excess of 100 pile up in between, giving rise to a third, then fourth, etc., layer, consisting of up to 50 atoms, for the typical load conditions of the simulations.

To guarantee an efficient melting-freezing of the soft lubricant in correspondence to the stick-slip motion for the “incommensurate” lubricant-substrate interface, we require a rather strong inequality \( V_{sl} \gg V_{ll} \). Specifically, we choose \( V_{ll} = 1/9 \) and \( V_{sl} = 5 \). In detail, we set the lubricant-substrate interaction energies to be \( V_{sl} = 5 \) in the “totally-glued” case, while for the “head-glued” system, the large interaction \( V_{sl} = 5 \) applies exclusively to the “head” molecular atom, while other atoms in the rest of the molecule (the “tail”) interact with the substrates with the much weaker \( V'_{ll} = 1/9 \). With these parameters, the melting temperature of the lubricant is of the order of \( T \approx 0.1 \); it is a generally decreasing function of the lubricant layer thickness, but an increasing function of the chain length and applied load \( \mathbf{1} \).

The load force used is typically \( f_l = 1 \) per substrate atom. Assuming an energy scale \( 9V_{ll} \approx 0.1 \, \text{eV} \) and interatomic distances \( R_{ll} \approx 0.3 \, \text{nm} \) (typical of Van der Waals intermolecular interactions), in a hypothetical 3D geometry such load would correspond to a pressure of almost 1 GPa.

The top substrate is driven at an average velocity \( v_s \) through a relatively stiff spring of elastic constant \( k_s = 0.1 \). For the remaining parameters, we take \( R_{sl} = R_s \) and \( z_d = R_s \). We run all simulations starting from well-relaxed and annealed initial configurations, and exclude from averaging any initial transient leading to the steady dynamical state. In more detail, we solve the Langevin equations of motion using a fourth-order Runge-Kutta method with a time step \( \Delta t = \tau_0/128 \), where \( \tau_0 = 2\pi R_{ll}(72 \cdot 9 V_{ll}/m_l)^{-1/2} \) is an average characteristic period of the LJ interatomic potentials. The initial configuration is realized by a careful annealing cycle up to the high temperature \( T = 2 \), followed by a “relaxation” run, typically of 4096 \( \tau_0 \) or longer, at the appropriate Langevin temperature \( T \) and sliding velocity \( v_s \), before measuring the friction force by averaging over a run typically of the same duration. The final configuration of every run is saved: this allows us to restart the run when we need to improve accuracy. To estimate error bars of various quantities, we split the measuring trajectory into 30 pieces of equal duration, and then estimate the variance of the global average through the standard deviation of the averages carried out over each individual piece. To verify the results, for some parameters we also made runs for total integration times and system
that the method allows us to evaluate $\zeta < f$ indeed, during sticks the force grows as $\zeta$.

Figures 3a–c show the simulation results for $T = 0$, reporting the time-averaged kinetic friction force $f_k$ as a function of the total number of lubricant atoms, which is basically proportional to the lubricant layer thickness, for different molecular lengths $L$.

The monoatomic lubricant (diamonds in Fig. 3) forms two complete monolayers for $N = 100$ and three monolayers for $N = 150$. For $N \geq 150$, one ordered layer sticks to the bottom substrate and another ordered monolayer to the top substrate, while the layer(s) in between are liquid during sliding. For the monatomic lubricant, the static friction force (not shown) is larger than the kinetic friction by a factor of 2 to 4; the transition from stick-slip to smooth sliding occurs at $v_s \approx 0.03$. Friction generally decreases as the number of lubricant atoms $N$ and thus the lubricant thickness $-\zeta$ grows, but the dependence $f_k(N)$ cannot be described by the simple laws of viscous-friction flow. At lower lubricant thickness, the kinetic friction force (not shown) is larger than for the monomer lubricant. In general, regardless of $L$, friction decreases roughly monotonically with increasing $N$, the lubricant thickness. In Fig. 3 the lubricant amount decreases further, $70 < N \leq 100$, there remain only two, generally incomplete, lubricant layers, each glued to the nearest substrate (e.g., $N = 100$ atoms form exactly two complete lubricant layers, which slide slowly, thanks to incommensurability, over the substrates to which they are attached). At even smaller values of $N$ we have one monolayer or eventually just a monolayer island in the narrow gap between substrates, and at this point both the friction and the critical velocity $v_c$ grow rapidly when $N$ decreases.

For all chain molecules (polymer lubricant) $L > 1$, friction is always larger than for the monomer lubricant. The kinetic friction force increases with the chain length $L$, if the total number of atoms $N$ is kept fixed. In general, regardless of $L$, friction decreases roughly monotonically with increasing $N$, the lubricant thickness. In Fig. 3 we re-organize the kinetic friction $f_k$ data of Fig. 3 but as a function of the number of molecules $M$ (only points corresponding to three or more molecular layers are shown). In the standard viscous flow of a fluid with a viscosity $\eta$, the friction force $f_k$ should depend on the thickness $d = \langle Z \rangle$ of the liquid layer $(d \propto N \propto M)$ as $f_k = \eta v_c R_s^2/d$. Unlike expectations for a fluid lubricant at high temperature, at $T = 0$ the log-log plot of the function $f_k(M)$ suggests that the simulation data may be (crudely) fit by power laws $f_k \approx 0.18 (M/N_s)^{-1/2}$ for the totally-glued case, and $f_k \approx 0.26 (M/N_s)^{-2/3}$ for the head-glued system. Accordingly, the function $f_k(M/N_s)$ displays a nontrivial apparent "universal scaling" with thickness, at least within the nanoscale thickness range

FIG. 2: (Color online) Comparison of the stick-slip motion [$v_s = 0.03$, panels (a,c,e)] and the smooth sliding [$v_s = 0.1$, panels (b,d,f)] for $M = 40$ totally-glued three-atomic ($L = 3$) lubricant molecules at $T = 0$. Panels (a) and (b): typical time dependence of the friction force $f(t)$; panels (c) and (d): histogram $P(f)$ of the friction force $f(t)$; panels (e) and (f): depth distribution $P(\zeta)$. The top substrate is curved, see Fig. 1 and the parameters are the following: $V_\text{int} = 1/9$ and $R_{\text{sl}} = 1$, $V_{\text{sl}} = 5$ and $R_{\text{sl}} = R_s$, $f_i = 1$, $k_s = 0.1$, $R_s = 2/3$, and $N_s = 62$.

sizes $2 \times 4$ times larger to check that the results do not change.

As in typical tribological systems, we observe a transition from stick-slip to smooth sliding with increasing driving velocity. An example is shown in Fig. 2. In the smooth sliding regime, the kinetic friction force can easily be measured as the average of the spring force $f(t)$. To evaluate the static friction force $f_s$ in the stick-slip regime, one can calculate the histogram $P(f)$ of $f(t)$ and then select $f_s$ as the point where $P(f)$ decays below a given level, e.g., $P(f_s) = 0.5 \times \max P(f)$ (see Fig. 2). A harder problem is that of determining the precise transition from stick-slip to smooth sliding and its critical velocity $v_c$, since this transition is often a smooth crossover. For this purpose, we calculate the distribution (histogram) $P(\zeta)$ of an auxiliary variable $\zeta$, defined as $\zeta(t) = [df(t)/dt] / (k_s v_s)$. In smooth sliding the function $P(\zeta)$ decays monotonically from a maximum at $\zeta \approx 0$ although this transition is often a smooth crossover. For this purpose, we calculate the distribution (histogram) $P(\zeta)$ of an auxiliary variable $\zeta$, defined as $\zeta(t) = [df(t)/dt] / (k_s v_s)$. In smooth sliding the function $P(\zeta)$ decays monotonically from a maximum at $\zeta \approx 0$ regime (see Fig. 2). In contrast, for stick-slip, $P(\zeta)$ should show a maximum at some $0 < \zeta \leq 1$ (Fig. 2). Indeed, during sticks the force grows as $f(t) \approx k_s v_s t$ (so that $P(\zeta) = \delta(\zeta - 1)$), and it drops quickly to substantially smaller values during short slip events. Thus, this method allows us to evaluate $v_c$ fairly accurately.
FIG. 3: (Color online) Smooth sliding ($T = 0$, $v_s = 1$) time-averaged kinetic friction force $f_k$ as a function of dimensionless coverage $N/N_s$ (bottom scale), as well as total number of atoms $N = ML$ (top scale) for linear lubricant molecules of different lengths $L = 1$ (black diamonds), 3 (red circles), 5 (magenta up triangles), and 7 (blue down triangles), for either the totally-glued (open symbols and dotted lines) or head-glued (solid symbols and curves) cases. The top substrate is curved according to Eq. (5) (see Fig. 1), and the other parameters are the following: $V_{sl} = 1/9$ and $R_{ll} = 1$, $V_{sl} = 5$ or $1/9$ and $R_{ll} = R_s$, $f_i = 1$, $k_s = 0.1$, $R_s = 2/3$, and $N_s = 62$. Considered here. The slower decay of friction compared to $d^{-1}$ may be attributed to a non-uniformity of local heating due to sliding-induced local shear. At fixed driving velocity, the local shear decreases as thickness increases, whence the local viscosity involved increases, and so does frictional dissipation, causing a positive deviation from the fluid-like $M^{-1}$ behavior. In other words, the higher average kinetic energy, and therefore higher effective temperature, produces a smaller viscosity $\eta$ [3], thus a smaller friction for a thinner lubricant layer than for a thicker one, moderating the power-law decay of $f_k$ in Fig. 3. For a much thicker lubricant, a $T = 0$ simulation would eventually show lubricant solidification with slip movements associated to local melting, even at the very high sliding velocity considered here.

Figure 3 reports similar calculations carried out at a finite temperature $T = 0.2$, compared to which the extra kinetic energy induced by shearing at sliding velocity $v_s = 1$ becomes negligible when the lubricant thickness exceeds about $100R_{ll}$. When the lubricant thickness exceeds such values we do observe a crossover to the expected scaling $f_k \propto (M/N_s)^{-1} \propto d^{-1}$, as soon as the shearing kinetic energy becomes smaller than the thermal fluctuations imposed by the finite $T = 0.2$. Note that small finite-size effects induced by the periodic simulation box tend to delay slightly the onset of the $f_k \propto d^{-1}$ constant-viscosity regime.

Analyzing the simulation movies for the totally-glued lubricant with $L > 1$ and large enough $N$ we observe two complete layers sticking to the corresponding substrates and a liquid lubricant between these layers. However, not all boundary molecules are glued completely, some of them are glued partially, i.e. only a part of a molecule sticks to the substrate while another part is detached.
and points into the lubricant bulk. It is precisely these “free tails” that make the friction higher as compared with the monoatomic case. At lower lubricant thickness ($M \lesssim 50$ for the $L = 3$ case, $M \lesssim 35$ for the $L = 5$ case, and $M \lesssim 25$ for the $L = 7$ system) the middle layer is incomplete: its molecules group together at the narrowest intersubstrate gap and just ahead of it (jamming/plowing effect). With a further decrease of lubricant thickness, sliding takes place at a single interface only, namely between the lubricant layers which are attached to bottom and top substrates. In this case, we observe a slow “caterpillar” motion of glued molecules relative the substrate, but this effect is weak because of the strong lubricant-substrate interaction. At even lower lubricant thickness occasional molecules may be glued to both surfaces simultaneously, and in such cases the molecular end sticking with fewer atoms slides along the corresponding substrate.

We come now to analyze the head-glued lubricant, comparing its behavior to the totally-glued one described above, all other parameters being the same. Firstly, the head-glued system (solid symbols in Figs. 3 and 4) display a systematically higher friction than the totally-glued lubricant (open symbols). As in the totally glued case, the surfaces are covered by lubricant layers. Again, the layers are orientationally ordered: in this case the molecular heads are attached to the corresponding surface, while the tails are directed outside the surface, and inclined in the sliding direction, like combed hair. Also, the sliding interface is more “irregular”, than in the totally-glued system. Therefore, one may expect a smoother frictional motion even in the stick-slip regime at low driving velocities. It is important to note that contrary to the fully glued case, in the head-glued system we find essentially no jamming effects even at low lubricant thickness.

Figure 6 depicts the different atomic density $\rho(z)$ and velocity distributions $v_x(z)$ and $|v(z)|$ across the lubricant for the totally- and head-glued molecular lubricants. In both cases, one layer sticks to the substrate, but the thickness of this layer is $\sim R_{ll}$ in the totally-glued system.
and much thicker, $\lesssim L R_{\text{mol}}$, similar to the molecular gyration radius, in the head-glued case. Accordingly, while the sliding profile $v_s(z)$ is fairly linear in the totally-glued case, in the head-glued case the shearing dissipation concentrates mostly in a bulk lubricant region remote from the substrates.

To elucidate the different tribological properties of the two kinds of molecular lubricants, in Fig. 7 we analyze the distribution of energy losses (dissipated power) due to sliding, with the method described in Ref. [2]. The losses occur mainly within the first (glued) lubricant layer, as a consequence of the model Eqs. (2) and (3), mimicking the capability of different lubricant layers to transfer energy into phononic/electronic degrees of freedom of the substrates. Importantly, we find that the main losses come from atomic vibrations in the transverse ($z$) direction, by far dominant over those coming from the motion along the driving ($x$) direction. This observation explains why, even though head-glued molecules displace sliding away from the substrates surfaces where most dissipation takes place, the head-glued system leads to a higher friction: transverse vibrations of larger amplitude penetrate better near the substrates (where most dissipation occurs) through head-glued molecules than through totally-glued ones.

Our results suggest that a totally-glued lubricant is more effective than a head-glued one, because it provides lower kinetic friction. According to Sivebaek et al. [3], totally-glued molecules with their stronger sticking to the substrate, should also be more stable against squeezeout of the contact area at high load. Unfortunately, as was said in the beginning, in our simplified 2D model we cannot study quantitatively the squeezeout process. Squeezout involves nucleation of a small $(N-1)$ negative island or “crater” and its subsequent expansion, a process which also involves elasticity of the substrates, and which is very characteristically three-dimensional [4]. We can, however, analyze in detail the dependence of frictional forces on the number of lubricant molecules $M$. Results for the $L = 5$ lubricant molecules are presented in Fig. 8.

It is useful to define the maximum number of molecules $M_2$ which can arrange themselves exactly to form the two interface molecular layers (one attached to the bottom substrate and the other to the top substrate), so that sliding takes place in the remaining lubricant just in between these two layers. For the totally-glued lubricant $M_2 = 2 L_x/R_{\text{Hil}}$, and for the head-glued lubricant $M_2 < 2 L_x/R_{\text{Hil}}$. $M_2$ depends on $L$ and on the load force, so that it takes an analysis of the MD trajectories to define its precise values. In the current model (with $L_x = 41.3$, $N_s = 62$, $L = 5$) we find $M_2 = 20$ for the totally-glued system and $M_2 = 58$ for the head-glued case. Based on the definition of $M_2$ we can introduce the dimensionless coverage $\theta_M = M/M_2$, so that $\theta_M = 2$ corresponds to precisely two molecular layers. Figure 8 shows that, as expected, both the friction force and the critical velocity $v_c$ increase when the coverage decreases below the two-layer value. The rise of friction below $M_2$
heat-glued ones. Note also that the static friction of the totally-glued chains is much higher than for the head-glued ones. In the totally-glued case, the velocity depends only weakly on the molecular length $L$.

We come now to discuss in some detail how the tribological properties vary with the system parameters. Consider first the role of temperature. Figure 10 reports the temperature dependence of the kinetic friction force $f_k$ of the monoatomic lubricant $(L = 1, N = 357, M_2 = 100, \text{diamonds})$ and of head-glued (full circles) and totally-glued (open triangles) chain molecules of length $L = 7$ $(M = 51, N = 357, M_2 = 14)$, at fixed driving velocity $v_\text{d} = 0.316$. The inset shows the effective thermal expansion of the mean lubricant thickness. Error bars are smaller than the symbol size. Other simulation parameters same as in Fig. 9.

The $L$-dependence of friction forces is illustrated in Fig. 9 for a configuration with exactly two molecular layers $(\theta_M = 2)$. In detail, the critical velocity $v_c$ marks the crossover illustrated in Fig. 2 and the frictional forces $f_s$ and $f_k$ are evaluated immediately below and above $v_c$, in the stick-slip motion and smooth sliding regimes respectively. Note that for the head-glued lubricant, the critical velocity depends only weakly on the molecular length $L$, while in the totally-glued case, $v_c$ increases rapidly with $L$, so that for long molecules, $L > 5$, the critical velocity of the totally-glued chains is much higher than for the head-glued ones. Note also that the static friction $f_s$ for the stick-slip motion at $v_s \lesssim v_c$ is approximately twice as large as the kinetic friction $f_k$ in the smooth-sliding regime at $v_s \gtrsim v_c$.

We come now to discuss in some detail how the tribological properties vary with the system parameters. Consider first the role of temperature. Figure 10 reports the $T$-dependence of kinetic friction at a fixed driving velocity $v_\text{d} = 0.316$, in the smooth sliding regime. Friction generally decreases when temperature grows, as is to be expected of the viscosity decrease of a fluid. General trends for different lubricant molecules are similar: the head-glued lubricant provides the largest friction, the monoatomic lubricant the lowest friction. It is interesting that the temperature dependence of kinetic friction $f_k$ for the totally-glued lubricant is significantly weaker than for the head-glued and monoatomic lubricants. The same type of dependence was observed for different driving velocities. The monoatomic lubricant friction actually increases with $T$, at low temperature, where the lubricant structure is crystalline. This is expected for an incommensurate lubricant/substrate interface. The thermal expansion of the simulated lubricant is rather large, especially for the monoatomic lubricant, as shown in the inset of Fig. 10.

Next, Fig. 11 shows a typical load dependence of the kinetic friction. The friction force increases with load, but not quite linearly, i.e., the friction coefficient $\mu_k = f_k / f_1$ is not constant (unlike Amontons first law): $\mu_k$ decreases gradually the load $f_1$ increases. The reason is most likely associated with the increasing effective shearing temperature associated with a fixed shearing velocity under increasing lubricant compression, the lower viscosity in turn reducing friction. Note that friction of the head-glued lubricant is approximately twice that of the totally-glued system.

Figure 12 reports the dependence of the friction forces $f_s$ and $f_k$ on the driving velocity $v_\text{d}$ at a nonzero temperature $T = 0.1$. The friction forces generally rise with velocity, but very gently so. By comparison with Fig. 9 we note that the transition from stick-slip to smooth sliding at a nonzero temperature and for more than two layers occurs earlier than at $T = 0$ and for exactly two layers. The transition shows a nonmonotonic dependence on the molecular length $L$, stick-slip persisting to the highest
and $N = 3$ (between the lubricant molecules. At small
and head-glued (solid curve and open diamonds) lubricant.

Data refer to $M = 70$ chains of $L = 5$ atoms, other parameters
same as in Fig. 3. Note the semilog scale.

Critical velocity for $L = 5$. In the low-$v_s$ regime the friction
data tend to become noisy, announcing the well-known difficulty of MD to address the low shear rates of realistic lab experiments. For an extension of the present study to that regime, a transient-time correlation function approach\cite{12} would be of substantial help.

Figure 13 demonstrates the dependence of the kinetic friction on the strength of the interaction $V_{ll}$ between the lubricant molecules. At small $V_{ll}$, e.g., for $V_{ll} = 0.1$, the surfaces are covered by monolayers of lubricant molecules, and the sliding interface lies somewhere near the middle of the lubricant film. As $V_{ll}$ increases, sliding becomes more and more viscous, thus friction increases reaching a maximum near $V_{ll} = V_{ll}^* \lesssim V_{si} = 5$. At this point, the sliding mechanism changes: the lubricant becomes rigid enough to move as a whole body, and frictional sliding is now shifted to the substrate/lubricant interfaces. If $V_{ll}$ increases further beyond this point, sliding becomes easier and easier because of increasing lubricant rigidity; this regime corresponds to the sliding of two hard incommensurate surfaces\cite{1, 2}. We found the same type of crossover for other numbers of lubricant molecules.

Finally, Fig. 14 demonstrates the variation of the friction coefficient with the substrate lattice constant $R_s$. In these simulations, we keep the size of the system constant at $L_x = 62 \times 2/3 \approx 41.3$, and vary the number of substrate atoms from $N_s = 41$ to 83, so that $R_s = L_x/N_s$ changes from $\approx 0.5$ to $\approx 1$. The total load force $F_l = f_l N_s$ is also kept constant at the level $F_l = 62$ (so that we have $f_l = 1$ for $N_s = 62$ as in the previous simulations). As expected, there appears to be no systematic dependence of friction on the substrate/lubricant interface $R_s/R_{ll}$ for the head-glued system, where the molecules arrange themselves at frequently varying mutual distance, and no commensurability effect can arise. For the totally glued case, however, the friction coefficient shows a well pronounced minimum at $R_s \lesssim 0.75$, especially for a lower value of the lubricant-substrate interaction $V_{si} = 1$. This is associated to a slow “cater-
some molecules are attached with all caterpillar-type motion indeed takes place. However, as values are shifted upwards [14]. As shown in Fig. 15, a R or R = 1 lubricant molecules of length L = 5 for totally glued (blue open symbols) and head-glued (green solid symbols) chains. The sliding velocity is v_s = 0.316, the interaction radius R_sl = R_s is varied, and in (b) all other parameters are as in Fig. 13 including V_{sl} = 5 or 1/9, while in (a) the gluing interaction is V_{sl} = 1 or 1/9.

![Graph showing friction coefficient vs. R/R_s](image)

**FIG. 15:** (Color online) The central region of five successive snapshots of the motion of M = 70 totally-glued L = 5 lubricant molecules. The time interval between successive snapshots (a) to (e) is Δt = 1. Snapshots b-e display only the atoms nearest to the bottom substrate, for better visibility of the caterpillar motion. To provide a clearer reference frame, one of the static atoms in the bottom-substrate is drawn as empty. The other parameters are as in Fig. 13(a), except for R_{sl} = R_s = 2/3.

pillar” motion of the glued molecules in the periodic substrate potential [14], caused by advancing of kinks (misfit dislocations) at the lubricant-substrate interfaces [15,16,17]. For a rigid linear molecule in a sinusoidal potential, virtually free sliding of the molecule should be observed for ratios R_s/R_{mol} = L/(2L - n), where n = 1, 2, ..., int[L/2]. This gives, e.g., R_s/R_{mol} = 2/3, or R_s ≈ 0.57 for L = 2 and 4, and R_s/R_{mol} = 5/8, or R_s ≈ 0.535 for L = 5. For a nonrigid molecule, these values are shifted upwards [14]. As shown in Fig. 13 a caterpillar-type motion indeed takes place. However, as some molecules are attached with all L atoms but others stick only partially, e.g., with 2 – 4 atoms, while the glued layer should move as a whole (the molecules cannot overtake one another), caterpillar motion is disturbed, and this makes the minimum in Fig. 13(a) broad and shallow. The dependence of friction on the incommensurability between the substrate and molecular spacings was actually observed experimentally [18].

V. DISCUSSION AND CONCLUSION

We presented simulations of a simple 2D model of molecular lubrication. Results indicate that layers of similar thickness of totally-glued lubricant molecules generally produce a smaller friction than lubricant molecules that are head-glued. Totally-glued lubricants also stick better to the substrates, and boundary layers thereof may accordingly be harder to squeeze out under high load. On the other hand, whenever the applied load becomes strong enough to squeeze the lubricant out to less than two molecular layers, then the head-glued kind behaves much better in sustaining smooth sliding and low friction even at very small residual coverage. We predict that when such extreme-load situations are likely to occur in a practical lubricated sliding devices, head-glued molecules could provide the best lubricant, unless the squeeze-out resistance is much worse than for totally glued lubricant. Moreover, when the base lubricant is totally-glued as a typical oil and, therefore, provides low friction properties, then even a small concentration of head-glued additives such as diblock polymers may prevent the substrates from wearing at places (asperities) where the surfaces come close to each other within a thickness of less than two layers. This is a new result, susceptible of experimental test.

Similarly, friction shows a clear increasing trend as a function of molecular length – the number L of monomer units in the molecules. This result is similar to the L dependency of the bulk viscosity, and our results compare well to those of Ref. [4]. This would make monoatomic liquids the best lubricants, if it was not for their much reduced resistance against squeeze-out, which eventually makes polyatomic chains more effective lubricants at loads where boundary lubrication matters.

The detailed search for the best compromise between the necessity of a squeeze-out resistance, calling for large L, and of low-friction characteristics, calling for small L, is a rather intricate task that is probably best undertaken on a case-by-case base by experimental trial and error procedures, or possibly by very extended simulations of sophisticated 3D realistic models. The simple 2D model at hand allows us to understand the main mechanisms at play in the soft lubricant and to predict the general trends. For example, local extra heating induced by the increased lubricant shear velocity at thinnest lubricant regions between substrate corrugations may well act as an important mechanisms producing a considerable overall friction reduction in macroscopic sliding.
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