Relaxation and Recovery in Hydrogel Friction on Smooth Surfaces

B. Wu · J. S. Méndez Harper · J. C. Burton

Received: 3 December 2020 / Accepted: 9 June 2021 / Published online: 22 June 2021 © Society for Experimental Mechanics 2021

Abstract

Background Hydrogels are crosslinked polymer networks that can absorb and retain a large fraction of liquid. Near a critical sliding velocity, hydrogels pressed against smooth surfaces exhibit time-dependent frictional behavior occurring over multiple timescales. The origin of these dynamics is unresolved.

Objective Here, we characterize this time-dependent regime and show that it is consistent with two distinct molecular processes: sliding-induced relaxation and quiescent recovery.

Methods Our experiments use a custom pin-on-disk tribometer to examine poly(acrylic acid) hydrogels on smooth poly(methyl methacrylate) surfaces over a variety of sliding conditions, from minutes to hours.

Results We show that at a fixed sliding velocity, the friction coefficient decays exponentially and reaches a steady-state value. The time constant associated with this decay varies exponentially with the sliding velocity, and is sensitive to any precedent frictional shearing of the interface. This process is reversible; upon cessation of sliding, the friction coefficient recovers to its original state. We also show that the initial direction of shear can be imprinted as an observable “memory”, and is visible after 24 hrs of repeated frictional shearing.

Conclusions We attribute this behavior to nanoscale extension and relaxation dynamics of the near-surface polymer network, leading to a model of frictional relaxation and recovery with two parallel timescales.

Keywords Hydrogel · Friction · Polymer · Tribology

Introduction

A hydrogel consists of a solvent-saturated, crosslinked polymer network that exhibits unique frictional properties due to its ambiguous nature. When swollen with liquid, osmotic pressure forces from the hydrophilic nature of the polymer network give rise to macroscale elasticity. However, hydrogels can be over 90% liquid by weight, leading to a self-lubricating slippery surface [1–3]. When exposed to a sliding interface, this dichotomy results in non-trivial and time-dependent frictional behavior that is important for numerous applications including biomaterials [4–11], biomechanics [12–15], soft robotics [16–22], and industrial water management [18, 23–29]. The friction coefficient associated with a given interface generally depends on contact geometry [30–35], sliding velocity [36–39], liquid viscosity [40], hydration [41, 42], polymer density [29, 40, 43], and any physicochemical absorption of polymers to the sliding layer [1, 44].

Although hydrogels exhibit interesting bulk rheologies, sliding interfaces expose the near-surface polymers in the network to shear forces that are orders of magnitude larger than those in the bulk. Such strong forces can alter the polymers’ entropic configurations and preferentially stretch them in the sliding direction [36, 37, 39–41]. This deformation can result in time-dependent frictional behavior during a single experiment at constant velocity [38–40, 45]. Although the polymers near the interface are generally sparse, forming a crosslinked network with a “mesh size” [46] of order 1-200 nm [29, 40, 47, 48], local entanglements and crosslinking constraints may increase the extension and relaxation rates. In Kim et al. [38, 39, 45] and Cuccia et al. [40] time-dependent frictional behavior was observed over periods of minutes or longer. Furthermore, Cuccia et al. [40] showed that for hydrogels on smooth surfaces, the friction coefficient evolves over multiple timescales which are not directly proportional to liquid viscosity, suggesting that the frictional evolution is not solely due to re-hydration of the interface. The authors suggested that topological polymer entanglements may give rise to a free energy landscape with sufficient complexity to explain the timescales of frictional relaxation and recovery.
Here, we show how the friction between a hydrogel and a smooth surface evolves under a broad set of sliding conditions by rapidly varying the velocity and pre-shearing of the interface. During some experiments, the hydrogel interface is allowed to rest in order to examine the recovery of friction after sliding. For polyacrylic acid (PAA) hydrogels on poly(methyl methacrylate) (PMMA) surfaces, we find that the friction coefficient, $\mu$, only displays time dependent behavior between $v \approx 0.3 - 3 \text{ cm/s}$, where $v$ is the sliding velocity. For a given sliding velocity in this regime, $\mu$ decays exponentially and approaches a steady-state value, independent of the sliding history of the hydrogel. However, the time constant and initial value of the friction do depend on the sliding history. In particular, the time constant decreases exponentially with sliding velocity. Similarly, the recovery of $\mu$ is exponential in the waiting time elapsed since the last experiment. We propose a simple, phenomenological model that captures the salient features of this behavior. The model consists of two parallel dynamical processes (relaxation and recovery), with timescales that can vary with the sliding velocity. Finally, and surprisingly, we show how the initial sliding direction can be imprinted as a “memory” into the time evolution of $\mu$, even after 24 hrs of continuous shearing.

### Experimental Methods

We performed experiments using the custom pin-on-disk, bi-directional tribometer, illustrated schematically in Fig. 1(a). A similar device was used in Cuccia et al. [40]. A spherical sample – PAA or agarose – of radius $R \approx 7.5 \text{ mm}$ was held stationary to the end of a low-force strain sensor (Strain Measurement Devices S256). This cantilevered spherical sample rested upon a horizontal PMMA substrate fixed to a circular, rotating frame. Using atomic force microscopy, we determined that the root-mean-square roughness of the PMMA to be 3.8 nm over a 1 $\mu\text{m}^2$ area. The strain sensor measured the tangential force ($F_t$) on the sphere and was calibrated prior to use. A normal force $F_n = 0.2 \text{ N}$ was applied above the point of contact using a fixed load, $F_n$. A macroscopic layer of water over the substrate kept the contact hydrated during the experiment. We varied the thickness of the surrounding water from 1-5 mm and found no difference in the frictional force measurements, so we used a consistent thickness of 3 mm for all experiments reported here. Any bulk drag on the sphere as it moved this layer of fluid would result from velocity gradients over a length scale $R$, as in Stokes drag. This drag is much smaller than the frictional drag under the contact region, where velocity gradients occur over a length scale $d$ (Fig. 1(c)). All experiments were conducted at $22 \pm 2^\circ\text{C}$.

The experimental setup was automated using a LabVIEW program. To achieve a broad range of rotational velocities at the point of contact between the spherical sample and the substrate, we drove the substrate with a stepper motor operating in a “microstepping” mode together a set of timing belts. This arrangement allowed us to investigate frictional behaviors at velocities ranging from 0.03 m/s and 60 m/s. The analog voltage at the output of the strain sensor was digitized using a USB data acquisition unit (NI USB-6008) at a rate of 24 kSamples per second. To reduce noise, 12,000 data points were

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**Fig. 1** (a) Diagram of the experimental setup showing the spherical sample pressed against the rotating surface. The hydrogel is sheared in the $x$-direction. (b) The characteristic diameter of the circular contact area is $2a$. (c) The polymer network adjacent to the surface experiences shear as fluid is dragged through it. (d) Plot of $\mu$ vs. $v$ for a PAA particle with $F_n = 0.2 \text{ N}$ on an PMMA surface for three different values of the experimental running time, $t_p$, at each data point. Also shown is the effective decay length of the velocity into the hydrogel, as calculated from (equation (3)) using the viscosity of water ($8.9 \times 10^{-3} \text{ Pa.s}$). Error bars are not shown for clarity, but noise from the measurement apparatus is comparable to the point size.
averaged every 0.5 seconds. Thus, the friction coefficient was effectively measured two times per second. Additionally, we were able to visually assess the point of contact through a milled hole in the structure underlying the spinning substrate.

Individual experiments had durations ranging from 30 s < t_p < 3600 s and the data from each trial was averaged to provide a single value of the friction coefficient µ_avg. For some experiments (where indicated), the experiment was paused for 5 s, and then rotated in the opposite direction for the same time t_p in order to average over both directions and calibrate the zero-point of the force sensor. At some velocities, the data displayed an exponential-like decay towards a steady-state value, leading to a large variation in the friction coefficient. We particularly observe these effects within the transition regime when the friction decreased rapidly with sliding velocity. We also observe that this decaying behavior is far more sensitive to changes in speed than changes in direction.

We used two types of hydrogels for our experiments: poly(acrylic acid) (PAA) and agarose. Agarose samples were fabricated in the lab, and the PAA spherical particles were purchased from JRM Chemical (Cleveland, OH). The PAA particles were composed of approximately 70% polyacrylic acid (PAA) and 30% acrylamide monomer, as reported by the manufacturer. Spherical agarose samples were made by mixing agarose (0.5-2.0 g) into 100 ml of distilled water and heating the solution to 60 °C. The solution was then pipetted into a spherical silicone mold and left aside at room temperature for 30 mins. The hydrogel was then immersed in water for at least 2 hrs. All chemicals and solvents used for fabrication were purchased from Millipore Sigma. We measured the ultimate swelling ratio of the gels by immersing them in water and letting them equilibrate. Dry commercial PAA particles, when immersed in water, would swell from an initial radius of ≈ 1 mm to a final radius of ≈ 7.5 mm. Once swollen, the commercial PAA particles consisted of 99.2% water.

Results

Regimes of Frictional Behavior

First it is important to note the distinct regimes of frictional behavior we observe in hydrogels. At low velocities, the hydrogel is pressed firmly against the PMMA surface, making a circular contact area of radius a ≈ 3 mm (Fig. 1(b)). By varying the normal load and imaging the corresponding contact area using optical microscopy, we confirmed that the hydrogels obeyed a Hertzian relationship [40]. For a sphere pressed against a flat surface, the Hertz theory predicts that

\[ d^3 = \frac{3F_n R}{4E^*}, \]

where \( E^* \) is the reduced average modulus for the sphere and the substrate:

\[ \frac{1}{E^*} = \frac{1 - \nu_g^2}{E_g} + \frac{1 - \nu_s^2}{E_s}. \]

Here \( E \) and \( \nu \) are the Young’s modulus and Poisson’s ratio for the gel (g) and substrate (s). The effective modulus for the PAA particles was measured as \( E^* \approx 45 \) kPa, and for 2% wt agarose it was measured as \( E^* \approx 47 \) kPa. The contact area was observed from below for some experiments, as was found to be independent of time and velocity [40].

As shown in Fig. 1(c), when the hydrogel is adjacent to the surface, fluid is dragged through the porous polymer network. The characteristic decay length of the velocity is of order the mesh size [40], so that the frictional force is proportional to the viscous shear stress at the interface, as described by Newton’s law of viscosity:

\[ \mu = \frac{A \eta}{d} \frac{v}{d}. \]

Here \( \eta \) is the dynamic viscosity of the solvent, \( A = \pi a^2 \) is the area of contact, and the velocity gradient has been replaced with \( v/d \), where \( v \) is the sliding velocity and \( d \) is the effective decay length of the velocity into the hydrogel surface layer (Fig. 1(a)), which is proportional to the pore or “mesh” size of the hydrogel [40]. At low velocities, this leads to a friction coefficient that increases with velocity due to viscous drag (Fig. 1). The friction grows slower than linear with \( v \) since \( d \) can vary slightly with shear stress [40]. We can solve (equation (3)) for \( d \), and plot the result as a function of velocity, also shown in Fig. 1(d). In the low-velocity regime, the friction coefficient is independent of the amount of time spent measuring it at a given data point (t_p). For the PAA particles used in this study, \( d \approx 10-30 \) nm, and for the agarose particles, \( d \approx 4-8 \) nm [40]. At high velocities, a bulk fluid layer of thickness ≈ 1 μm develops between the hydrogel and the PMMA surface. In this regime, the friction coefficient is determined by elastohydrodynamic lubrication theory (EHL), and is also independent of \( t_p \). Both the low velocity and high velocity regimes are described in detail in Cuccia et al. [40].

Relaxation of Friction During Sliding

At intermediate velocities (0.3 cm/s < v < 3.0 cm/s), we observe time-dependent dynamics. This can be seen in Fig. 1(d), where the sharp drop in friction and transition to the EHL regime is sensitive to \( t_p \). The variation of \( \mu_{avg} \) with \( t_p \) seen in Fig. 1(d) occurs because the friction decays
exponentially with time during the course of an experiment at a single velocity [40]. At each new data point, $\mu_{\text{avg}}$ depends on the new velocity, but also on the history of shearing it has experienced. In order to tease apart this history dependence, we performed a series of experiments with a hydrogel particle that has been allowed to “rest” for 2 hrs prior to sliding. Figure 2(a) shows the time evolution of $\mu$ after immediately shearing the hydrogel at different velocities for $t_p = 1800$ s. To compare the data, we normalize the data for each experiment by dividing by the maximum $\mu$ at $t = 0$ s, denoted as $\mu_{\text{max}}$. For low velocities (0.23 cm/s), there is no variation in $\mu$ with time. However, at higher velocities, $\mu$ decreases and approaches a steady-state value. The rate of decay increases dramatically with velocity. The data at each sliding velocity is fit remarkably well with a single exponential form, as shown by the solid line in Fig. 2(a).

$$\mu = \Delta \mu \times e^{-\frac{t}{\tau}} + \mu_f,$$

(4)
as shown by the solid line in Fig. 2(a). Here, $\mu_f$ is the final, steady state friction coefficient, $\Delta \mu = \mu_{\text{max}} - \mu_f$ is the change in friction, and $\tau$ is the decay time constant. Each of these fitting parameters varied with the sliding velocity, as shown in Fig. 2(b-d). Both $\Delta \mu$ and $\mu_f$ reached constant values at high velocities, 0.03 and 0.005, respectively. However, the decay time $\tau$ decreased rapidly in this range of sliding velocities, and follows an exponential dependence on $v$, as shown in Fig. 2(b).

This dependence suggests that the imposed shear may serve as an excitation mechanism to induce relaxation in the polymer network adjacent to the surface. A classical analogy is the $\alpha$-relaxation time for strong glass formers, which decreases exponentially with inverse temperature [49]. Using (equation (4)), we can also compute the average friction coefficient over the time interval $t_p$, as was plotted in Fig. 1(d):

$$\mu_{\text{avg}} = \mu_f + \Delta \mu \frac{\tau}{t_p} \left(1 - e^{-\frac{t_p}{\tau}}\right).$$

(5)

From this it can be seen why $\mu_{\text{avg}}$ drops so sharply in Fig. 1(d), since $\tau$ appears in the exponent, and also decreases exponentially with velocity.

**Recovery of Friction During Rest**

In addition to exponential relaxation at sliding velocities in the intermediate regime, the friction coefficient also experiences a “recovery” phase upon cessation of sliding. If the shear induced by sliding is able to preferentially align the near-surface polymers and modify the local entanglement

$$\mu_{\text{avg}} = \frac{1}{t_p} \int_0^{t_p} \mu(t) \, dt,$$

fitting each data set to a single exponential form (equation (4)). Both $\mu_f$ and $\Delta \mu$ approach constant values at high velocities, while $\tau$ decays exponentially.
structure of the network, then thermal fluctuations should facilitate a reversible transition to an equilibrium state in the absence of shear. Figure 3 shows a series of experiments where a fresh PAA hydrogel sphere is sheared for 1 hr, and then allowed to rest for a given waiting time $t_w$. At the end of each experiment, as in Fig. 2(a), the hydrogel is able to reach its final friction coefficient ($\mu_f$). After waiting $t_w = 60$ s, there is a small amount initial increase in the friction before it rapidly decays again. For larger values of $t_w$, both $\mu_{max}$ and $\tau$ increase so that the friction coefficient recovers more and it subsequently takes longer to fully decay again when sliding is initiated. Finally, after waiting 1800 s, the frictional evolution has almost fully recovered to that of a fully-rested hydrogel.

By fitting the time series of $\mu$ to the single exponential form (equation (4)) for different values of $t_w$, we find that $\mu_{avg}$, $\tau$, and $\Delta \mu$ all recover to their initial values after waiting for a sufficiently long time. This is shown in Fig. 4(a-c). The dashed lines show the values associated with experiments from fresh, rested hydrogels. Importantly, even the decay time, $\tau$, seems to depend on the pre-shearing and $t_w$, changing by roughly a factor of 4 from short to long waiting times. This suggests that during recovery, the polymer network evolves over multiple timescales back to equilibrium. This is reasonable since structural evolution in disordered systems typically contain a variety of timescales due to the ruggedness of the free energy landscape [50]. More specifically, in the case of polymer brushes, the timescale for polymer retraction can be $10^8$ or larger than the Rouse time, and can depend on the degree of local entanglement and polymer chain length [51].

All of the data shown in Fig. 4(a-c) are consistent with a generic, single exponential recovery: $y = c_1(1 - e^{-t_w/e_2}) + c_3$, where $c_1$, $c_2$, and $c_3$ are fitting parameters. These fits are shown by the solid lines. At the moment, we do not assign physical meaning to these parameters, but the origin of the fitting form will be discussed in Sec. 3.5. We cannot be certain what detailed molecular mechanism accounts for recovery in our experiments without more direct measurements. However, the data necessitates a mechanism that accommodates a range of timescales that are history-dependent. It is unclear if

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**Fig. 3** The friction coefficient $\mu$ versus time for a spherical PAA hydrogel being shear at $v = 0.85$ cm/s. “New experiment” corresponds to a hydrogel that has rested for 2 hrs, whereas $t_w$ refers to the elapsed time after shearing that the hydrogel was allowed to wait prior to subsequent shearing.

**Fig. 4** (a) $\mu_{avg}$ versus $t_w$ for a PAA hydrogel sphere on PMMA. After initially shearing the hydrogel at $v = 1.27$ cm/s until it has reached a steady state value ($\mu_f$), the hydrogel is allowed to rest for a time $t_w$. Then data is acquired for $t_p = 600$ s at $v = 1.27$ cm/s for each data point. (b) The decay time, $\tau$, versus $t_w$, obtained from fitting each 600 s time series of $\mu$ to (equation (4)). (c) The change in friction coefficient, $\Delta \mu$, versus $t_w$, also obtained from fitting the data to (equation (4)). The solid lines represent exponential fits to the form $y = c_1(1 - e^{-t_w/e_2}) + c_3$, as described in the text. Error bars represent the standard deviations from three separate runs. The dashed lines in each panel represent the average values for a hydrogel which has not experienced prior shearing.
hydrodynamic models based on lubrication and poroelasticity alone could explain these results. This will be discussed further in Sec. 4.

**Asymmetry of Approach to $\mu_f$**

The approach to a steady-state friction coefficient, $\mu_f$, depends on the shearing history of the hydrogel, and can be asymmetric. Figure 5 shows the decay in $\mu$ from two separate experiments at two different velocities. If the hydrogel is initially rested for 2 hrs before starting an experiment, $\mu_{avg}$ starts higher and decreases exponentially to $\mu_f$, as in Fig. 2. However, if the hydrogel is instead sheared at a high velocity, e.g. 3.0 cm/s, for 1800 s, the polymer network is presumably in the “fully sheared” state, far from its equilibrium, rested value. In this regime, $\mu_f$ has plateaued with velocity (Fig. 2(c)). By immediately switching back to a lower velocity, the friction coefficient will then increase with time, and approach the same value of $\mu_f$.

For both $v = 0.48$ cm/s and $v = 0.58$ cm/s in Fig. 5, the initial value of $\mu$ and the decay constant $\tau$ is different, but the data for a given velocity tends to the same asymptotic value at long times. This behavior is reminiscent of the Kovacs effect in glassy, disordered materials [52, 53]. The effect is nonlinear in nature since perturbations away from equilibrium follow different relaxation mechanisms. Nevertheless, the fact that $\mu_f$ remains the same regardless of the shearing history suggests that the asymptotic sheared state of the system only depends on the sliding velocity.

**Model for Frictional Relaxation and Recovery**

The various timescales associated with relaxation and recovery can at first seem complex. However, a simple phenomenological model with two parallel timescales can capture the qualitative and some of the quantitative features of nearly all data for PAA hydrogel friction in the intermediate regime. We start by introducing a state parameter, $\gamma$, which has a range between 0 and 1. This parameterizes the sheared state of the hydrogel network at a given moment. For example, the fully “rested” state with no shearing, $\gamma = 0$, characterizes the state of the polymer network in the low velocity regime, where fluid is dragged through a mostly-isotropic polymer network, and $\mu$ is determined by (equation (3)).

In order to be as quantitative as possible, we take into account that $\mu_{avg}$ is not perfectly linear in $v$ at low velocities (Fig. 1(d)). Thus, we use a power-law relationship to describe $\mu$ in this regime:

$$\mu = K_1 v^\alpha,$$  \hspace{1cm} (6)

where $K_1$ is a fitting constant and $\alpha$ an exponent close to 1. In our model, we denote the friction coefficient of the hydrogel when $\gamma = 1$ as $\mu_1 = K_1 v^\alpha$. On the other hand, $\gamma = 0$ parameterizes the state where the hydrogel is fully sheared. We define a constant $\mu_0$ to be the lowest friction coefficient that the hydrogel can attain in the time dependent regime. For example, $\mu_f \approx 0.005$ at $v = 1.9$ cm/s in Fig. 2(c). For simplicity, we assume that $\gamma$ varies linearly between $\mu_1$ and $\mu_0$, or

$$\gamma = \frac{\mu - \mu_0}{\mu_1 - \mu_0}.$$  \hspace{1cm} (7)

We now consider the relaxation and recovery processes separately. First, when perturbed by shear, thermal fluctuations will drive the polymers near the surface back to equilibrium ($\gamma = 1$). As shown in Fig. 4(a), this process is well-captured by an exponential fit with a single time constant of order 1000-3000 s. We define this recovery time constant as $\tau_1$ as it refers to recovery to the $\gamma = 1$ state. The evolution of $\mu$ is then given by:

$$\frac{d\mu}{dt} = \frac{\mu_1 - \mu}{\tau_1}.$$  \hspace{1cm} (8)

We note that the recovery time $\tau_1$ should be independent of sliding velocity, although it may depend on the fluid properties and temperature.

In the absence of recovery processes, the application of a sliding shear will lead to relaxation of $\mu$ to the minimum possible value, $\mu_0$. The evolution of $\mu$ is then given by
\[ \frac{d\mu}{dt} = \frac{\mu_0 - \mu}{\tau_0}, \]  

where \( \tau_0 \) is the time constant associated with relaxation to the \( \gamma = 0 \) state. As suggested by Fig. 2(b), we assume \( \tau_0 \) depends on the sliding velocity \( v \):

\[ \tau_0 = K_0 e^{-v/v_c}, \]

where the fitting constants \( K_0 \) and \( v_c \) can be determined by fitting the data in Fig. 2(b).

During an experiment, these two processes are occurring in parallel. The full evolution of \( \mu(t) \) is given by combining (equations (8) and (9)):

\[ \frac{d\mu}{dt} = \frac{\mu_1 - \mu}{\tau_1} + \frac{\mu_0 - \mu}{\tau_0}, \]  

The measured dynamics of \( \mu \) for a given experiment will depend on its initial shear state \( \gamma \) and the sliding velocity \( v \). Since \( \mu_1 \) and \( \tau_0 \) don’t explicitly depend on time (only on \( v \)), we can solve the equation analytically with the initial condition \( \mu(t = 0) = \mu_i \):

\[ \mu = (\mu_i - \mu_f)e^{-t/\tau} + \mu_f. \]  

The solution is a single exponential decay where

\[ \mu_f = \frac{\tau_0 \mu_1 + \tau_1 \mu_0}{\tau_1 + \tau_0}, \]

\[ \tau = \frac{\tau_1 \tau_0}{\tau_1 + \tau_0}. \]

With this analytic form, we can recreate much of the data from the experiments. The empirical components of the model are the velocity-dependent forms of \( \mu_1 \) and \( \tau_0 \) (equations (6) and (10)). Figure 6 shows analytic predictions which quantitatively capture many of our results. Panels (a-c) correspond to Figs. 1(d), 3, and 5. Panels (d-f) correspond to Fig. 4. By choosing \( \mu(t = 0) \) and the sliding velocity \( v \), an analytic time series can be created over a time \( t_w \). This data then serves at the starting condition for the next data point in a simulated experiment shown in Fig. 6.

One of the most important insights from this model is that the effective decay time \( \tau \) observed during an experiment depends

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**Fig. 6** Results from the analytic model (equation (12)) for different experimental protocols with \( \tau_1 = 1910 \) s, \( K_0 = 8887 \) s, \( v_c = 0.276 \) cm/s, \( \mu_i = 0.006 \), \( a = 3/4 \), and \( K_1 = 0.050 \) (s/cm)^1/2. (a) Starting from an initial, unsheared state (\( \gamma = 1 \)), \( \mu_{avg} \) is computed over a time \( t_w \). The value of \( \mu \) at the end of each period is used as the initial condition for the next sliding velocity. (b) Recovery of friction with \( v = 0.85 \) cm/s. After an initial evolution for 1800 s, the \( \mu \) evolves with \( v = 0 \) cm/s for a time \( t_w \). Then sliding proceeds at the same velocity. The value of \( \mu \) after the recovery period is used as the initial condition for each time series. (c) The green downward triangles and black squares show the evolution of \( \mu \) at the given sliding velocities with no pre-shear. After evolving \( \mu \) at \( v = 3.0 \) cm/s for 1800 s, \( \mu \) is then allowed to evolve and approach the same steady-state value, demonstrating an asymmetry of approach. (d-e) \( \mu_{avg}, \Delta \mu, \) and \( \tau \) plotted as a function of waiting time \( t_w \). Starting from an unsheared state, \( \mu \) evolves at \( v = 1.27 \) cm/s for 600 s, then is allowed to recover for \( t_w \). The value of \( \mu \) at the end of each times series is used as the initial condition for the next time series.
on a parallel combination of two independent time constants (equation (14)). Thus, $\tau$ will be smaller than either $\tau_1$ or $\tau_0$. This explains why we observe a variety of timescales for frictional dynamics in our experiments which depend on sliding velocity. Since Fig. 2(b) is technically reporting $\tau$ and not $\tau_0$, we convert it to $\tau$ using (equation (14)) and then apply (equation (10)) for curve-fitting for all the simulation results reported above.

An important limitation of the model is evident in Fig. 6(f), which shows $\tau$ versus $t_{\text{av}}$. Since $\tau$ only depends on sliding velocity in our model and not on $t_{\text{av}}$, it is constant throughout the modeled experiment. However, as Fig. 4(b) shows, $35 \leq \tau \leq 175$ s throughout the physical experiment, implying that $\tau_1$ may depend somewhat on the state of the system, $\gamma$. This is reasonable since the perturbations from equilibrium can be quite large, and there can simultaneously exist slow and fast timescales due to the large deformation of the polymer network.

**Agarose Hydrogels**

Although most of our experiments were performed with PAA hydrogel, we also tested agarose hydrogel spheres. In addition to acrylic acid and acrylamide hydrogels, Cuccia et al. [40] reported a similar rise in friction at low velocities for agarose spheres, followed by a sharp drop in friction in the time-dependent regime. However, the dynamics of $\mu$ versus $t$ for agarose hydrogels is quite distinct from PAA, and is not exponential. Figure 7(a) shows that $\mu$ often starts at rather large values, near 0.4, then drops precipitously with an increasing rate until it reaches values near 0.01 after 100 s of sliding. For these experiments, $\Delta \mu$ can be quite large from the beginning to the end of a time series, more than 10 times the final value $\mu_f$.

Nevertheless, the friction experiences similar recovery times as the PAA hydrogel, and is consistent with an exponential recovery process, as shown in Fig. 7(a). Agarose is a physical gel with transient crosslinks, suggesting that its near-surface network structure can be substantially altered due to shear, in contrast to the permanent crosslinks in PAA. We suspect that this may explain the non-exponential behavior and sharp drop in $\mu$ in Fig. 7(a), and subsequently the large change in friction upon recovery (Fig. 7(b)). Rather than being perturbed from equilibrium, the local deformation and structural rearrangement is dramatic. However, if the recovery process is controlled by thermal fluctuations proportional to $k_BT$, or possibly by poroelasticity, then the recovery timescales could be similar to PAA. These possibilities will be discussed further in Sec. 4.

**Directional Memories in PAA Hydrogels**

Finally, we explored the long-time behavior and directional dependence of PAA hydrogel friction. Starting with a fresh hydrogel sphere that did not experience pre-shearing, the frictional force was measured in both the counterclockwise (CCW) and clockwise (CW) direction. For each time series, $\mu_{\text{avg}}$ is computed for $t_w = 600$ s, as in Fig. 4. Error bars represent the standard deviations from three separate runs.

To confirm that this is not an experimental artifact, we performed separate experiments with fresh hydrogels with different initial shear directions. Figure 8(b-c) shows that $\mu$ always experiences recovery when shearing in the initial direction, and relaxation in the opposing direction. However, $\mu_f$ was essentially the same in both directions, indicating that the final frictional state of the interface ($\gamma$ in our model) was independent of direction. We note that this asymmetry cannot be captured by our model in its current form since direction is not explicitly included.

We suspect that this memory is imprinted by some semi-permanent orientational ordering at the hydrogel interface due to the initial sliding. It is unclear if this change would fully recover given enough rest time, and
warrants further investigation. We did not investigate the directional dependence of agarose hydrogels because the decay time of \( \mu \) is generally much shorter (Fig. 7), and because agarose is a physical gel that is easily altered in long-duration experiments and may experience wearing of the surface.

**Discussion and Summary**

The results presented here illustrate unique, time-dependent frictional behavior between a hydrogel and a smooth, hard surface. In a range of intermediate sliding velocities, \( \mu \) experiences a decay toward a new, velocity dependent value with a time constant that is itself velocity dependent. The frictional dynamics are reversible. The frictional state will recover given sufficient resting time, and there is no evidence of permanent wear. This behavior can be modeled quantitatively using two distinct time constants representing shear-induced relaxation and exponential recovery toward equilibrium.

Our semi-quantitative analytic model is similar to other “rate and state” models used recently to describe time-dependent frictional behavior in hydrogels. However, the underlying mechanism leading to time-dependent behavior has not been fully resolved. Similar time-dependent behavior first observed in Kim et al. [39] was modeled using a frictional shear stress that followed two separate power-law behaviors according to a state variable that parameterized the rheological state of the interface [38, 45]. A mechanism involving rehydration of the interface, characterized by a heterogeneous distribution of bulk fluid pockets, was employed to explain the time and velocity dependence. Furthermore, studies examining bovine cartilage at high normal loads showed an exponential decrease in compression and friction vs. time, which was attributed to rehydration [42, 54]. This decrease occurred more rapidly at higher velocities, in analogy with the experiments presented in Fig. 4. However, Meier et al. [55] and Simič et al. [56] showed how the entropic freedom of near-surface polymers alter \( \mu \) by more than an order of magnitude. They also illustrate an exponential decay of \( \mu \) whose time constant (while less than 1 minute) decreases with velocity.

As proposed in Cuccia et al. [40], the extremely long relaxation times observed in the experiments, much longer than the Rouse timescale, could be attributed to shear-induced deformation and entanglements of the near-surface polymer layer. This is schematically shown in Fig. 9. The characteristic relaxation time for an isolated polymer blob can be written as \( \tau \approx \eta \xi^3/\kappa B T \), where \( \xi \) is the blob or mesh size [43, 46]. The shear rate used in our experiments is of order \( \dot{\gamma} = v/d \approx v/\xi \), since we are assuming that the characteristic decay length of the velocity into the hydrogel is of order \( \xi \). The Weissenberg number generically describes the threshold for polymers to deform in a shear flow, and is given by \( W = \dot{\gamma} \tau = \eta \xi^2/\kappa B T \). For water at room temperature (\( \eta \approx 0.0009 \) Pa·s), \( v = 0.5-1.0 \) cm/s, and \( \xi \approx 10-30 \) nm (Fig. 1(d)), we see that \( W = 0.1-2.0 \). This illustrates that the strong shear rates in our experiments, of order \( 10^5 \) s\(^{-1} \), are sufficient to initiate deformation of the polymers near the surface.
During deformation, the polymer entanglements are pushed to a smaller volume, leading to long relaxation times due to the required cooperative motion for disentanglement. As the shear rate is increased, the free energy barriers to disentanglement are reduced, and thus the relaxation timescale sharply decreases with sliding velocity. Upon the cessation of sliding, thermal fluctuations drive the polymers back to their unsheared state, yet this can also take hundreds to thousands of seconds since it involves the cooperative motion of many polymers near the surface (Fig. 3).

However, long-time, transient behavior could also be expected from fluid flow through the nanoscopic porous polymer network, and thus rehydration of the contact region remains an important consideration. Indeed, a simple estimate using diffusion coefficients for bulk water transport, $D \approx 10^{-9}$ m$^2$/s [57], and a length scale of $a = 3$ mm, one can arrive at a poroelastic timescale of $\approx 9,000$ s. However, in the bulk, moving fluid through the pores requires pressures larger than the osmotic pressure [58, 59], and would lead to contact areas that may vary with time or sliding velocity. According to Hertzian contact theory, the maximum pressure beneath the sliding contact is of order 11 kPa in our experiments, and the osmotic pressure must approach zero near the surface of the hydrogel. Here the polymer surface layer is “brushy” [56], and the width of this transition region can be 10’s of microns thick [57]. Certainly this layer can shrink and swell in response to the addition of water into the contact region when the lubrication pressure of the water can support some of the normal load. Yet the diffusive timescale for transport in and out of this layer is fast (on the order of 1 s) [57], and cannot explain the 30 minute or longer timescales observed in our experiments.

It is still possible that rehydration and swelling of this brushy interface is occurring in the transient regime, but we hypothesize that the long timescales and exponential relaxation would still require a restructuring of polymer entanglements near the surface. Very slow, long-time creep has also been observed in compression experiments [58], and was attributed to the diffusion of network heterogeneities rather than poroelasticity based on similar arguments. Given to preponderance of rehydration as a potential progenitor of time-dependent friction in the literature, it is important to summarize and enumerate why rehydration cannot solely explain our results:

1. The Weissenberg number, $\dot{\gamma} \tau$, is of order unity at the experimentally measured transition velocity, 0.5-1.0 cm/s. This is clear evidence that the polymers near the surface will be affected by the shear flow.

2. The time constant associated with frictional relaxation varies exponentially with velocity (Fig. 4(b), equation (10)). We are unaware of a rehydration-based explanation for this dependence. A long relaxation time, much longer than expected for an isolated polymer chain, can exist in crowded, entangled polymer brushes [61], and a specific exponential dependence of the relaxation time on shear rate can exist in end-absorbed polymer brushes [62]. For polyacrylamide hydrogels, experiments support this “brushy” approximation near the surface [56, 57].

3. For two separate hydrogels with similar elastic modulii (PAA and agarose), the decay dynamics are different, one exponential (PAA, Fig. 4), and one distinctly non-exponential (agarose, Fig. 7). Agarose is a physical gel and we suspect that permanent alteration to the sliding surface occurs. Nevertheless, poroelastic rehydration is unlikely to explain this strong asymmetry between the gels.

4. The recovery (increase) in $\mu$ observed in our experiments after shearing, i.e. Fig. 5, shows that the hydrogel surface has been structurally altered after shearing at higher velocities. The friction is more than a factor of 2 lower than it would be if a fresh, fully-swollen gel had been tested. If this were due to rehydration, the lubrication pressure ($\approx 7$ kPa) would need to significantly “over-swell” the gel surface layer well beyond its equilibrium, unloaded value, and require nearly 1800 s to recover.

5. The directional memories observed for 24 hrs simply cannot be explained by a model that only considers swelling and rehydration from flows perpendicular to the hydrogel’s surface.
The hydrogel interface remains a subject of active research due to its complex nature. Our results for hydrogels on smooth surfaces represent a significant advance in our understanding of the polymer network’s response to high shear rates, yet there remains many unanswered questions. Ultimately, experiments involving a molecular probe of the surface dynamics may resolve many of these issues surrounding the importance of local polymer physics versus poroelasticity in determining the transient frictional response.

Acknowledgements The authors would like to acknowledge support from NSF DMR Grant No. 14550869, and also from the Emory SURE summer program for undergraduate research.

Declarations

Conflicts of Interest The authors declare that they have no conflict of interest.

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