Single Polymer Dynamics in an Elongational Flow

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The stretching of individual polymers in a spatially homogeneous velocity gradient was observed through use of fluorescently labeled DNA molecules. The probability distribution of molecular extension was determined as a function of time and strain rate. Although some molecules reached steady state, the average extension did not, even after a ~300-fold distortion of the underlying fluid element. At the highest strain rates, distinct conformational shapes with differing dynamics were observed. There was considerable variation in the onset of stretching, and chains with a dumbbell shape stretched more rapidly than folded ones. As the strain rate was increased, chains did not deform with the fluid element. The steady-state extension can be described by a model consisting of two beads connected by a spring representing the entropic elasticity of a worm-like chain, but the average dynamics cannot.

The behavior of dilute polymers in elongational flow has been an outstanding problem in polymer science for several decades (1, 2). In elongational flows, a velocity gradient along the direction of flow can stretch polymers far from equilibrium. Extended polymers exert a force back on the solvent that leads to the important, non-Newtonian properties of dilute polymer solutions, such as viscosity enhancement and turbulent drag reduction.

A homogeneous elongational flow is defined by a linear velocity gradient along the direction of flow such that \( \mathbf{v} = \hat{x} \hat{\theta} \), where \( \hat{\theta} = \frac{\partial v}{\partial y} \), the strain rate, is constant. Theory suggests that the onset of polymer stretching occurs at a critical velocity gradient or strain rate of \( \dot{\epsilon}_c \) of

\[
\dot{\epsilon}_c \sim \left( \frac{\eta}{\tau_c} \right)^{0.5}
\]

where \( \tau_c \) is the longest relaxation time of the polymer (3). For \( \dot{\epsilon} < \dot{\epsilon}_c \), the molecules are in a “coiled” state. But as \( \dot{\epsilon} \) is increased above \( \dot{\epsilon}_c \), the hydrodynamic force exerted across the polymer just exceeds the linear portion of the polymer’s entropic elasticity, and the polymer stretches until its nonlinear elasticity limits the further extension of this “stretched” state. De Gennes predicted that this “coil-stretch transition” would be sharpened by an increase in the hydrodynamic drag of the stretched state relative to the drag of the coiled state (1).

In many types of elongational flows, such as flow through a pipe per se, the residence time \( t_{res} \) of the polymers in the velocity gradient is limited. To increase \( t_{res} \), flows in which there is a stagnation point are often used. As molecular trajectories approach the stagnation point, \( t_{res} \) diverges. The classical techniques for inferring the degree of polymer deformation have been light scattering (4, 5) and birefringence (6–9). For example, Keller and Odell reported a rapid increase in the birefringence for \( \dot{\epsilon} \) above \( \dot{\epsilon}_c \), followed by a saturation (6). Such saturation was interpreted as an indication that the polymers had reached equilibrium in a highly extended state (10). Molecular weight analysis showed some chains are fractured in half, further supporting the hypothesis that the polymers reached full extension (8, 11). However, light-scattering experiments imply deformations of only two to four times the equilibrium size (4, 5). But, these “bulk” measurements average over a macroscopic number of molecules with a broad range of \( t_{res} \). Moreover, only recent experiments have been dilute enough to prevent the polymers from altering the flow field (9).

Many rheological effects also remain unexplained. James and Saringer measured a pressure drop in a converging flow that was significantly greater than that predicted by simple models (12). Recently, Tirtaaatmadja and Sridhar measured extensional viscosities \( \eta_{ext} \) in filament stretching experiments that were several thousand times greater than the shear viscosities (13). At large deformations, \( \eta_{ext} \) saturated, suggesting again that the polymers were fully extended. However, the measured stress was significantly lower than expected for fully extended polymers, implying that full extension had not actually been achieved (14). Also, the stress relaxation in such experiments contained both a strain-rate independent “elastic” and a strain-rate dependent “dissipative” component. The molecular origin of the dissipative component is uncertain (15). Examples such as these indicate that, even after a tremendous amount of study, the deformation of polymers in elongational flows is still poorly understood (14, 16).

We report the direct visualization of individual polymers in an elongational flow. The conformation and extension of each molecule was measured as a function of \( \dot{\epsilon} \) and \( t_{res} \), thereby eliminating the ambiguities in conformation and \( t_{res} \). We further eliminated polymer-polymer interactions and polymer-induced alterations of the flow field by working with single isolated molecules. The inherent uniformity...
in size of lambda bacteriophage DNA [λ-DNA, \( L_{\text{stretched}} \approx 22 \mu m \) (17)] also eliminated complications due to polydispersity and enabled accurate calculation of ensemble averages (18). Thus, we determined the probability distribution of molecular extension rather than just an average or a moment of that distribution.

Another advantage is that the entropic elasticity (19, 20) and hydrodynamic drag (17, 21, 22) of single DNA molecules have been previously characterized.

Using a microfabricated flow cell, we generated a planar elongational flow with a cross-slot geometry adapted for fluorescence microscopy (23). The main design consideration of the flow cell was to ensure that we studied dynamics of polymers unwinding from equilibrium (24). Our imaging area was 100 μm by 94 μm with the stagnation point 15 μm from the center of one side. The onset of the elongational flow, where \( t_{\text{res}} = 0 \), was 960 μm up the inlet channel from the stagnation point.

By tracking individual molecules, we measured the extension \( x \) and \( t_{\text{res}} \) of each molecule in our imaging area. Some molecules deformed only slightly, whereas others rapidly reached a steady-state extension (Fig. 1A). This large and previously unobservable heterogeneity was perhaps unexpected because these molecules were identical in size and had experienced the same ε and \( t_{\text{res}} \). From an ensemble of individual measurements, we calculated the average extension \( \langle x(t_{\text{res}}) \rangle \) as well as the time evolution of the probability distribution for molecular extension (Fig. 1B).

We characterized the conformation of each polymer in the ensemble. In general, the molecules were found in one of seven conformations which we refer to as dumbbell, half-dumbbell, folded, uniform, kinked, coiled, or extended. The first three types were dominant at \( \epsilon = 0.86 \) s⁻¹. As shown in Fig. 2A, these are highly nonequilibrium conformations, and they occurred only at higher strain rates (\( \epsilon > 0.5 \) s⁻¹) (25). In this case, the molecules were subject to a \( \epsilon \) significantly greater than the inverse relaxation time \( \tau_{\text{relax}} \approx 0.26 \); \( \tau_{\text{relax}} = 3.89 \) s (26). For λ-DNA (~400 persistence lengths), we saw only single folds at the highest \( \epsilon \) investigated. However, for longer molecules, we observed multiple folds (27).

There were clear differences in dynamics for the three dominant conformations. To highlight these differences, we plotted data, using only those molecules that best typified each conformational class. Molecules in a dumbbell configuration stretched significantly faster than folded ones (Fig. 2B). In addition, the residency time \( t_{\text{res}} \) at which significant stretching begins for any particular molecule was highly variable (28).

An analysis of the rate of stretching \( \dot{x} \) as function of \( x \) shows that once a molecule in a dumbbell configuration starts to stretch, its dynamics follows a specific time evolution (Fig. 2C, inset). This result indicates that the data would approximately collapse onto a single “master curve” by sliding the individual curves along the time axis. Up to \( \epsilon L = 0.6 \), we observed a linear increase in \( \langle x(t_{\text{res}}) \rangle \) with \( x = \pm 12 \mu m \) at \( \epsilon = 0.86 \) s⁻¹. When integrated, this yields an initial exponential growth of the master curve. We show three such master curves generated from the molecules that best typify each of the dominant conformations (Fig. 2C). For comparison, we show \( \langle x(t_{\text{res}}) \rangle \) for the full data set as well as for several of the different conformational classes arising from the first, general classification (Fig. 2D).

Because of the large variation in \( t_{\text{res}} \), the master curve better represents the unwinding dynamics of individual molecules and is different in shape than \( \langle x(t_{\text{res}}) \rangle \).

We plotted the fractional average extension \( \langle x/L \rangle \) as a function of the accumulated fluid strain or “Henky” strain \( \langle \epsilon = \epsilon^* t_{\text{res}} \rangle \) (Fig. 3A). By analyzing the subset of molecules that reached steady state (Fig. 3B, inset), we determined the steady-state extension \( x_{\text{steady}} \) as a function of the dimensionless strain rate or “Deborah number” \( \epsilon^* \tau_{\text{relax}} \) (Fig. 3B).

Note that \( x_{\text{steady}} \) rises sharply at a critical strain rate of \( \epsilon^* \tau_{\text{relax}} \approx 0.4 \) and that for \( \epsilon^* \approx 0.9 \), the fractional size of fluctuations is large (\( \sigma / L_{\text{steady}} \approx 0.4 \)). Similar behavior is often seen at phase transitions. In comparison with classical bulk measurements, we also plotted a spatio-temporal average \( x_\text{bulk} \) of all our data (Fig. 3B).

In a linear velocity-gradient flow \( \langle \epsilon_\text{fluid} = \epsilon = \partial v / \partial y \rangle \), the distance between two fluid elements grows as \( y \sim \exp(\epsilon_\text{fluid} t_{\text{res}}) \). There was a similar but slower exponential growth in the master curves of molecular extension for \( \epsilon > 0.21 \) s⁻¹. We defined a molecular strain rate \( \epsilon_\text{mol} \) from a fit of \( \langle x(t_{\text{res}}) \rangle \) to \( \epsilon_\text{mol} x + b \) over the region where \( \langle x(t_{\text{res}}) \rangle \) is a linear function of \( x \) (Fig. 4, inset) and compared \( \epsilon_\text{mol} \) to \( \epsilon_\text{fluid} = \epsilon \) (Fig. 4). This analysis averages over the conformation-dependent dynamics shown in Fig. 2. To single out the most rapid stretching conformation, we also plotted \( \epsilon_\text{mol} \) for the dumbbell configuration at \( \epsilon = 0.86 \) s⁻¹.

![Fig. 1. (A) The extension x as a function of the polymer’s interaction or residency time \( t_{\text{res}} \) in the elongational flow at \( \epsilon = 0.86 \) s⁻¹ for 992 molecules. Several individual traces are highlighted. We also plotted the average extension \( \langle x(t_{\text{res}}) \rangle \) as open circles. Notice in particular, the large heterogeneity in the dynamics of these molecules. To slow down the dynamics, we used an aqueous sugar solution (\( \eta = 41 \) cP). We imaged those molecules whose center of mass started within 22 μm of the center line of the inlet channel. Within this region, the measured velocity gradient along the full length of the inlet \( (\partial v / \partial x = -\epsilon) \) was linear and within 2% of the measured velocity gradient along the outgoing axis \( (\partial v / \partial y = \epsilon) \). Because the fluid is incompressible \( (\nu = 0) \) and it is a planar flow \( (\partial v / \partial z = 0) \), the molecules experienced a constant strain rate independent of position. Furthermore, the calibrated strain rate measured by tracking fluorescent beads agreed within 2% with the strain rate calculated from the motion of the center of intensity of individual DNA molecules. The data starts at \( t_{\text{res}} > 0 \), because the onset of elongational flow \( x_{\text{screen}} \) is 960 μm upstream from the stagnation point, whereas the edge of the imaging area \( x_{\text{screen}} \), is 86 μm upstream. The interaction time of the polymer with the applied velocity gradient before imaging is given by \( t = \ln(x_{\text{screen}}/x_{\text{screen}}) / \epsilon \). The raw extension data was smoothed by weighted averages with its nearest neighbor of \( x = 0.21 x_{\text{res}} + 0.59 x + 0.21 x_{\text{res}} \). (B) Time evolution of the probability distribution of molecular extension calculated from an ensemble of at least 40 individual molecules at \( \epsilon = 0.86 \) s⁻¹. The secondary peak arises from molecules in a folded configuration (Fig. 2). Keunings’ simulations of a bead-spring-bead model generated similar broadly shaped histograms (33), though such simulations are incapable of producing the secondary peak associated with folded configuration seen in the experimental data.](www.sciencemag.org)
A polymer is said to “affinely” deform with the fluid if the molecular deformation equals the deformation of the surrounding fluid element. It has been suggested that when \( \dot{\epsilon} \gg 1/\tau_{\text{relax}} \), affine deformation becomes an increasingly valid approximation (29). In the simplest analysis, we note that \((x(t_{\text{res}})) \) did not reach \( x_{\text{steady}} \) even after an accumulated fluid strain of \( \epsilon = -\dot{\epsilon}t_{\text{res}} \approx 5.7 \), which corresponds to an \( e^{5.7} \) or \( \approx 300\)-fold distortion of the fluid element (Fig. 3). For comparison, the required molecular distortion to fully extend stained \( \lambda \)-DNA is \( L/R_{G} \approx 30 \) while \( R_{G} \), the radius of gyration, is \( 0.73 \mu \text{m} \) (21).

In part, this lack of affine deformation in \((x(t_{\text{res}})) \) arises from the large variation in \( t_{\text{res}} \). Notwithstanding this variation which is intrinsically nonaffine, we wanted to know if molecules deform affinely once they start to stretch. To do so, we analyzed the dynamics of the master curve because it suppresses the variation in \( t_{\text{res}} \) by computing \((\hat{x}(\hat{x})) \) instead of \((x(t_{\text{res}})) \). At moderate strain rates, affine deformation is not expected, because there must be some slip between the polymer and the fluid to create the hydrodynamic force necessary to overcome the native elasticity of the polymer. Because there is no deformation for \( \dot{\epsilon} < \dot{\epsilon}_{\text{c}} \), we plotted \( x_{\text{max}} \) versus \( x_{\text{fluid}} \), where \( \dot{\epsilon}_{\text{c}} \approx 0.4/\tau_{\text{relax}} \). At lower \( \dot{\epsilon} \), the molecules are stretching the theoretically expected limit (Fig. 4). At higher \( \dot{\epsilon} \), the data shows a marked departure, and it is clear that the affine deformation approximation breaks down. Furthermore, when plotted as \( x_{\text{max}}/(\dot{\epsilon}_{\text{fluid}} - \dot{\epsilon}_{\text{c}}) \) versus \( (\dot{\epsilon}_{\text{fluid}} - \dot{\epsilon}) \), the data is decreasing at \( 0.86 \text{ s}^{-1} \). Thus, the data shows neither an absolute nor a fractional approach toward affine deformation at higher \( \dot{\epsilon} \) even after eliminating the large variation in \( t_{\text{res}} \). This failure arises from the introduction of intramolecular constraints (folds) which dramatically slow down the average dynamics. On the other hand, the subset of molecules in a dumbbell configuration stretched almost as fast as can be theoretically expected.

Our steady-state results are approximately characterized by a simple “dumbbell” model consisting of two beads connected by a spring based on the Marko-Siggia force law (Fig. 3B, solid line) (20). Previously, the steady-state extension of a tethered polymer in a uniform flow was well described by this model (17), and we developed a molecular understanding of the origin of this agreement based on simulations (22). An extrapolation of the model to \( \dot{\epsilon} = 0 \) gives a critical strain rate of \( \dot{\epsilon}_{\text{c}} \tau_{\text{relax}} \approx 0.4 \), which is near the theoretical value of 0.5 calculated from the Zimm model and by the numerical calculation of Larson and Magda (3). This value of \( \dot{\epsilon}_{\text{c}} \tau_{\text{relax}} \approx 0.4 \) is less than the values of 3 to 8 seen in recent birefringence measurements of polystyrene solutions by Nguyen et al. (9).

To see if this model could self-consistently describe the dynamics of the master curve, we calculated the expected dynamics, using parameters determined from the steady-state
results. Whereas the predicted rate of extension is close to the measured dynamics for the dumbbell configuration, it overestimates the average measured dynamics (Fig. 4, inset). So, although this dumbbell model describes the steady-state extension, it fails to describe these simplified dynamics in which the large variation in $t_{\text{rise}}$ is suppressed. Therefore, we expect difficulty in trying to predict the transient stress in the fluid by constitutive equations based on a simple dumbbell model (2). Given the nonaffine deformation, a term proportional to $-\dot{x}$ which can describe an “internal viscosity” might be added (2). Such a term is suggested by the measurements of $\eta_{\text{A}}$ because it leads to a dissipative component of the stress relaxation (15). Although a term proportional to $-\dot{x}$ can approximately compensate for the slower average dynamics, our data show that, in part, these slower dynamics arise from folded configurations which are meta-stable rather than arising from the monomer-monomer friction typically associated with internal viscosity. We note that there are additional terms besides $-\dot{x}$ that can lead to dissipative stresses (30, 31).

Given our measurements of the dynamic, steady-state, and ensemble-averaged properties of polymers in an elongational flow, we now compare our data to previous experimental and theoretical results. Atkins and Taylor measured the birefringence of $\lambda$-DNA in a similar planar elongational flow (Fig. 3B) (8). Our ability to select only those molecules that have reached steady-state extensions reveals a much sharper transition occurring at a lower $\dot{\varepsilon}_{\text{relax}}$. As discussed above, the higher value of $\dot{\varepsilon}_{\text{relax}}$ seen by birefringence occurred for synthetic polymers as well as for DNA (9). Evidently there is no direct correspondence between either $x_{\text{steady}}$ or $x_{\text{bulk}}$ and the birefringence at the stagnation point. Because birefringence measures orientation rather than extension, some disagreement would be expected based on the observed conformational features such as folds. However, folds would cause a premature saturation in birefringence with respect to $x_{\text{steady}}$. Our results highlight the difficulties in interpreting birefringence and other bulk measurements and suggest that this difficulty may be even greater for synthetic polymers, for which the larger ratio of $L/R_G$ requires an ever larger accumulated fluid strain than is needed to extend $\lambda$-DNA.

In contrast to previous light scattering results on synthetic polymers (4, 5), our data shows extensions significantly greater than $\sim 2 R_G$, though $R_G$ by definition, is always less than $x/2$. In general, the large difference between $R_G$ and $x_{\text{steady}}/2$ is caused by the broad distribution in $t_{\text{rise}}$ for the population of molecules measured by light scattering. Hence, $x_{\text{steady}}/2$, not $x_{\text{steady}}$, should be used for comparison. In addition, the highly asymmetric mass distribution of the most common conformation (half-dumbbell) would further reduce $R_G$. In particular, we directly calculated $R_G$ from the image data for $\varepsilon = 1.2$. A spatio-temporal average of this data yielded $R_G = 2.2 \mu m$, which is three times the equilibrium coil size [$R_G = 0.73 \mu m$ (21)] but is much smaller than steady-state extension ($x_{\text{steady}} = 14.8 \mu m$) at this $\dot{\varepsilon}$. Thus, our results help explain the apparent discrepancy between light scattering and birefringence measurements.

Our results suggest that midpoint chain fracture in stagnation point flows does not imply that all chains are extended. The large variability in $x$ (Fig. 1A) indicates that a number of molecules rapidly reach steady state. If we extrapolate our results to a $\varepsilon$ of 100 times higher, it is these highly extended, early-stretching molecules that will experience a force large enough to fracture at or near their center. Nonetheless, because of the limited $t_{\text{rise}}$, the number of such chains that are rapidly stretching and start stretching early is relatively small. Thus, only a fraction of the total number of chains will fracture in agreement with the results of bulk experiments (8, 11), but this fracture of some chains does not imply that all chains are extended.

Rheologists often infer molecular deformation from bulk viscoelastic measurements (2). Given the data in Fig. 1, the known elasticity of DNA (20), and classical results in rheology (2), one can calculate the extensional stress $\sigma_{\text{e}} = \eta \dot{\varepsilon} F(x)$ and the extensional viscosity $\eta_{\text{A}} = \sigma_{\text{e}}/\dot{\varepsilon}$ where $\eta$ is density of molecules and $F(x)$ is the steady-state elasticity. However, because these molecules are in highly non-equilibrium configurations (Fig. 2A), it is inaccurate to use the steady-state elasticity for molecules at $\varepsilon \gg 1/\tau_{\text{relax}}$. From this and the lack of a physically significant mean as described below, our results suggest difficulties with inferring an average conformation from bulk rheological measurements. Additionally, our results reveal problems with the use of the Peterlin approximation (32), in which $x^2(t_{\text{rise}})$ is replaced by $x^2(t_{\text{rise}})$, to derive constitutive equations that predict bulk rheological measurements from a micromechanical or kinetic theory (2). The heterogeneity in our data that leads to the breakdown of the Peterlin approximation is also seen in Keunings’ stochastic simulations of the finitely extensible dumbbell model (33). Although this simpli-

![Fig. 3.](image-url)
fied model of polymer dynamics based on kinetic theory yields histograms that are in semi-quantitative agreement with our data (Fig. 1B), simulations with the Peterlin approximation in conjunction with kinetic theory lead to qualitatively different results.

To account for the excess stress measured by James and Saringer (12), Ryskin, Larson, Hinch, and King and James have developed theories based on different hypothesized molecular configurations (16, 31). By direct observation of dumbbell, half-dumbbell, folded, and kinked conformations, we confirm the presence of conformations similar to those proposed. The presence of these conformations provides a qualitative explanation for the dissipative component of stress found in measurements of ηp. However, no one of these theories describes the complete range of observed conformations. Rather, the individual conformations assumed in these theories represent one of the several observed conformations.

From a theoretical point of view, the conformation-dependent dynamics implies that the commonly used approach of developing mean-field theories has an inherent disadvantage (34). The probability distribution is not a narrow distribution about a mean but rather a broad, oddly shaped distribution (Fig. 1B) because of several distinctly different dynamical processes (Fig. 2). Further, the differences in x and t0,exp indicate a sensitive dependence on the polymer’s initial configuration when it enters the velocity gradient. Presumably, these variations arise directly from the multitudes of accessible conformations at equilibrium where thermal fluctuations cause instantaneous deviations away from a spherically symmetric distribution. For instance, a polymer whose initial configuration has both ends on the same side of the center of mass and is subject to a τ > 1/τs,exp would most likely become folded, because there is not enough time (τs,exp) for an end to move to the other side of the molecule. Variations similar to those in our experimental data have been observed in the simulations of Larson (31), Hinch (16), and Keunings (33).

Although we observe a sudden increase in the steady-state extension of polymers at a critical strain rate, our data indicates that the concept of a discrete and abrupt collapse transition is limited to the steady state. Polymers do not undergo a simple, collective and simultaneous unwinding as soon as τ > τs,exp. The mismatch between x(t,x,exp) and xsteady implies that the non-Newtonian properties of dilute polymer solutions in most practical elongational flows (where \( \varepsilon_{\text{dyn}} < 5.5 \)) are dominated by the dynamic and not the steady-state properties. Our data should serve as a guide in developing improved microscopic theories for polymer dynamics and the bulk rheological properties of such solutions.

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Fig. 4. A comparison between the rates of deformation of the fluid element (Rbead = 6) and the polymer as characterized by the master curve. We subtract \( k_{\text{bead}} \), the experimentally determined critical strain rate, from the \( k_{\text{total}} \) to account for the necessary difference between it and \( k_{\text{total}} \). This difference leads to the hydrodynamic force that stretches the polymer against its native elasticity. The master curve is used instead of \( x(t_{\text{fit}}) \), as it better represents the unwinding dynamics by suppressing the variation in it. To define a molecule strain rate from the master curve, \( k_{\text{total}} \) was calculated from fitting \( x(t) = k_{\text{total}} x + b \) over the region where \( x(t) \) is a linear function of \( t \). As shown in the inset, this approximately linear behavior is seen for \( x < 11.8 \mu \text{m} \) at \( \varepsilon = 0.86 \text{ s}^{-1} \). For the lower \( k_{\text{bead}} \), \( x(t) \) is linear in \( x \) up to 5.2, 7.8, 8.5, and 10.2 \( \mu \text{m} \) in increasing order of \( \varepsilon \). If affine deformation becomes an increasingly valid approximation at higher \( \varepsilon \), then the data would line along the curve described by \( k_{\text{total}} = k_{\text{bead}} - k_{\text{aff}} \) (dotted line). \( k_{\text{aff}} \) determined for those molecules that best typify a dumbbell configuration is plotted as an open symbol (see Fig. 2C, inset). The inset \( x(t) \) versus \( x \) for all molecules at \( \varepsilon = 0.86 \text{ s}^{-1} \) (filled circles) and molecules in the dumbbell configuration (open circles). The solid line represent the dynamics predicted by the dumbbell model using the values \( k_{\text{total}} \), \( L \) determined by the fit of \( x_{\text{steady}} \) to \( k \) shown in Fig. 3B. More explicitly, we sum the hydrodynamic and entropic forces and set them to zero \( F_{\text{total}} = F_{\text{hydro}} - F_{\text{spring}} = 0 \), where \( F_{\text{hydro}} = 6\eta v\rho_{\text{bead}} L \) and \( F_{\text{spring}} \) as given in (20). Because \( x_{\text{predicted}} \) is calculated at a given \( x \), rather than \( F_{\text{total}} \), \( L \) is a function of only \( x \) and \( \varepsilon \) given \( F_{\text{hydro}} \) and \( L \) from the fit to the steady-state data. We can then calculate the predicted rate of stretching of the total chain at each \( x \) to be \( x_{\text{predicted}} = 2(6\eta v\rho_{\text{bead}} L) F_{\text{hydro}} / (6\eta v\rho_{\text{bead}} L + F_{\text{spring}}) \). The fluid element \( F_{\text{hydro}} \) (dashed line). Thus, this simple model overestimates the average dynamics although it agrees closely with the dynamics of molecules in the dumbbell configuration. The introduction of a term \( (1 + \alpha) \) multiplying \( x_{\text{predicted}} \) in \( F_{\text{hydro}} \), where \( \alpha = 0.55 \) approximately compensates for this overestimation of the dynamics at \( \varepsilon = 0.86 \text{ s}^{-1} \). Note that the variation in Fig. 4 away from \( k_{\text{total}} = k_{\text{bead}} - k_{\text{aff}} \) shows that this coefficient \( \alpha \) is dependent on \( k \).
ging, S. Smith, Science 265, 1599 (1994). Their ana-
lytic approximation for the elasticity is given by $\frac{k_B T}{k_F} = 0.25(1 - v/L) + 0.25 \times v/L$, where $L$ is the feature length across the ends, $v$ is the persistence length, and $k_B T$ is the thermal energy.


23. Following the example of W. D. Volkmann and R. H. Austin [Nature 358, 600 (1992)], we made our flow cell generator by sandwiching water with KOH and anion-
cally bonding Pyrex coverslips to the silicon to seal the top surface of the channels. Vertical side walls along the inlet and outlet channels were achieved by rotating the silicon by 45° to the crystal axis [C. Hu and S. Kim, Appl. Phys. Lett. 29, 582 (1978)]. To generate the flow, we used a syringe pump (Isco model 100 D) which was temperature-stabilized at 22.7°C. A 100-1 fluid shunt bypassing the flow cell was used to operate the pump at flow rates of 0.25 mm/min. The velocity field was calibrated by tracking fluorescent beads near the stagnation point and showed that $v_{z_{center}} = v_z$, confirming that a uni-
form elongational flow had been achieved. Our imaging and digitization system was the same as previously described [17], except we used a water immersion (40x C-APo NA 1.2) and stroboscopically illuminated the DNA to elim-
inate the blurring of the image due to camera lag. Our measurements of the equilibrium coil size gener-
ally did not yield $\langle r^2 \rangle = \tau_\alpha^2$, because $\tau_\alpha$ is slightly larger than $\tau_{relax}$ due to blooming in the camera. Once the chain is extended, a large number of times, the polymer is not in a linear elongational manner. It is not only the polymer, but something slightly larger than $\tau_{relax}$ is also blooming in the camera. Once the chain is bloomed, our characterization of the dynamics of long chains, because we did not have an adequately mono-dis-
perse sample and could not independently measure the length of individual molecules.

24. To prevent any predeformation of the polymer before entering the elongational flow, we used a cross-slot 0.25 mm wide and 220 mm deep and imaged the polymers at the center of depth of the channel ($z_{center} = 110$ mm) where the applied shear ($\dot{\gamma} = 0.02/s$) was negligible. Because of the requirement of a low refraction of the immersion fluid (water) and the high viscosity buffer, we deter-
duced $z_{center}$ by measuring the velocity of fluores-
cent beads in the inlet as a function of depth and adjusting as needed. The flow cell was mounted on a copper block and stabilized to 22.7°C ± 0.2°C.

25. At the lower strain rates, there were not distinct, well-
templated areas, but rather a general aspect of a dumbbell configuration subject to large fluctuations in extension and shape.

26. We determined $\tau_{relax}$ = 3.89 s by averaging the re-
laxation of 10 individual $R_F$ distributions. The relaxation time we report is from a fit over the region where $x/L < 0.3$ to $(x/L)^2 = c \exp(-x/L)$ with $c = 2.6$, where $x_{center}$, $c$, and $R_F$ were free parame-
ters. This is directly related to the relaxation of the stress in the fluid via $\tau_{relax} = \frac{n \langle x^2(t) \rangle}{(x^2(t))}$, because in the

limit of small deformation, $F(x)$ is linear in $x$ and this yields $\tau_{relax} = \tau_\alpha$. Thus, our characterization $\tau_{relax}$ via stress relaxation is made in the region ($x/L < 0.3$) where the distribution of higher order relaxation modes should be negligible (2). When scaled to $\tau_\alpha = 1$ centipoise and for the change in length caused by staining, our $\tau_{relax}$ for unstained, A-DNA in water is $R_F$, which yields $\tau_\alpha = 6.2$, and is the relative time we reported in an earlier experiment where the DNA was teth-
ered to a bead (35).

27. The longer DNA molecules were concentrators of $\lambda$-DNA up to 250 μm. We were unable to system-
atically investigate the dynamics of longer molecules, because we did not have an adequately mono-dis-
perse sample and could not independently measure the length of individual molecules.

28. The results presented here should not be general-
ized to polymers in a mixed elongational and shear-
ing flow or to polymers in a gravitational flow that were prehydrated. Our data indicates that the pro-
cesses involved in the diverse dynamics arise from the variation in $z_{center}$ and from internal configurations which is, folds. Mixed flows, a large fraction of the molecules are partially extended because of shear-
ning, and this effect may eliminate some of the internal constraints that led to the observed dynamics.


36. We thank G. Fuller, D. Hoagland, R. Larson, R. Pecora, and B. Zimm for helpful discussions and J. Shott for generous aid in topographic design. Sup-
ported in part by grants from the U.S. Air Force Office of Scientific Research, NSF, and the Human Frontiers Foundation and by an endowment established by Theodore and Frances Geballe. We acknowledge the generous assistance of J. Spudich, including support from NIH grant GM33289 to J. Spudich. D.E.S. was supported by a fellowship from the Program in Mathematics and Molecular Biology at the University of California, Berkeley, through NSF grant DMS 9406584. S.C. was supported in part by a Guggen-
heim fellowship during the course of this work.

18 February 1997; accepted 8 May 1997

Surface Stress in the Self-Assembly of Alkanethiols on Gold

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Surface stress changes and kinetics were measured in situ during the self-assembly of alkanethiols on gold by means of a microcantilever sensor. Self-assembly caused compressive surface stress that closely followed Langmuir-type adsorption kinetics up to monolayer coverage. The surface stress at monolayer coverage increased linearly with the length of the alkyl chain of the molecule. These observations were interpreted in terms of differences in surface potential. This highly sensitive sensor technique has a broad range of applicability to specific chemical and biological interactions.

Molecular and biomolecular layers are scientifically appealing for a wide range of potential applications (1, 2). Alkanethiols, which are known to self-organize into well-ordered, densely packed films, represent a model molecular system for controlling surface properties (3, 4). These self-assembled monolayers (SAMs) are used in applications such as mi-
crocontact printing (5) and voltammetric micro-
sensors (6), and they have recently been applied to molecular host-guest recognition (7).

There is little information available on the mechanical properties of SAMs, particularly concerning the nature of surface stress in films during the formation process, because it is difficult to follow the structural evolution of monolayer self-assembly. One recent approach (8) used scanning tunneling microscopy to infer the growth kinetics of alkanethiol SAMs indirectly from snapshot images ob-
tained at various coverages. Here, we used micromechanical sensors to gather quantitative data on surface stress changes that develop during the self-assembly process of HS-