Nonmonotonic dependence of polymer glass mechanical response on chain bending stiffness

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We investigate the mechanical properties of amorphous polymers by means of coarse-grained simulations and nonaffine lattice dynamics theory. A small increase of polymer chain bending stiffness leads first to softening of the material, while hardening happens only upon further strengthening of the backbones. This nonmonotonic variation of the storage modulus G' with bending stiffness is caused by a competition between additional resistance to deformation offered by stiffer backbones and decreased density of the material due to a necessary decrease in monomer-monomer coordination. This counter-intuitive finding suggests that the strength of polymer glasses may in some circumstances be enhanced by softening the bending of constituent chains.

Introduction The study of polymer dynamics has 56 been at the heart of soft matter research for decades, 57 yet a comprehensive theoretical basis that links monomer 58 chemistry to mechanical properties remains under devel-59 opment [1, 2]. Polymers below their glass transition tem-60 perature, which find application in everyday consumer 61 goods and high-technology material applications, pose 62 a particular challenge as understanding their properties 63 further requires an assimilation of glassy dynamics, itself 64 a topic of ongoing debate [3].

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Throughout the historical development of polymer ⁶⁶ physics, it has proven constructive to consider two ide-⁶⁷ alised linear polymer models: freely-jointed, in which ⁶⁸ chains are assumed to comprise random walks of fixed ⁶⁹ step length with no monomer interactions; and freely-⁷⁰ rotating, in which the angle formed by three consecutive ⁷¹ monomers is strictly fixed but the monomers are oth-⁷² erwise unconstrained. Here we explore the mechanical ⁷³ properties of polymer glasses between these limits as the ⁷⁴ monomer motions become increasingly constrained by a bending penalty. We further enforce excluded volumes ⁷⁵ around individual monomers.

It is already established that increasing the number 78 of constraints on particles in a many-body system reduces the critical coordination, and hence the critical 80 density, at which the system achieves marginal stability [4]. This has been apparent in granular systems 82 for some time, when comparing frictionless to frictional 83 packings [5]. Indeed, constraint-counting arguments underpin recent theories for shear thickening in athermal 85 suspensions [6]. An analogy between friction in granular systems and bending in polymers has been proposed 87 theoretically [7] and in experiments on 'granular polymers' [8] and is a promising lead towards unifying the 89 understanding of marginal stability across a surprisingly 90 broad class of soft matter systems [9, 10].

The introduction of bending constraints in bead-spring 92 polymer chains is expected to reduce the critical coordi-93 nation Z_q , *i.e.* the sum of inter- and intra-chain interac-94

tions at the glass transition, and, therefore, the critical density [11]. Such a density reduction is reminiscent of the role of plasticising additives [12], designed to reduce the mechanical strength of the material by increasing the free volume. By contrast, one might expect enhanced bending stiffness to increase the strength of the bulk material. The question remains, therefore, what overall effect the introduction of such constraints has on the mechanical properties of glassy polymers.

In this Rapid Communication we show using simulations and theory that the competing effects of increasing backbone strength and increasing free volume lead to nonmonotonic behaviour of the shear modulus of glassy polymers as a function of bending stiffness. This finding offers a connection between monomer chemistry and polymer glass rheology, demonstrating that manipulating bending constraints at the monomer level can have nontrivial influence on the bulk mechanical properties of the material.

Simulation details A non-overlapping random-walk algorithm is used to generate initial loose configurations of $N_p = 10^4$ monomers, in chains of length 10^2 . For each monomer in our system we use LAMMPS [14] to solve the Langevin equation with coefficient of friction $1/\xi$ and random forces $f_B(t)$ satisfying $\langle f_B(t)f_B(t')\rangle =$ $2mk_BT\delta(t-t')/\xi$ at time t. Monomers of uniform mass m interact through potentials U given by the Kremer-Grest model [15], comprising a Lennard-Jones potential $U^{\rm LJ}$ of depth $\varepsilon_{\rm LJ}$ and rest length $2^{1/6}\sigma$ acting between monomer pairs within a cut-off range $r_c = 2.5\sigma$ and a finitely extensible nonlinear elastic potential U^{FENE} with maximal length R_0 and emerging rest length $\approx 0.96\sigma$ acting between sequential monomer pairs along each chain [16]. $\varepsilon_{\rm LJ}$ sets the LJ energy scale and $\varepsilon_{\rm FENE}$ is the bond energy scale where $\varepsilon_{\text{FENE}}/\varepsilon_{\text{LJ}} = 30$. With reference to fundamental units of mass ν , length d, and energy ϵ , we set $\sigma = 1$ and m = 1, giving a time unit of $\tau = \sqrt{m\sigma^2/\varepsilon_{\rm LJ}}$, and we set $\xi = 100\tau$. We define a third energy associated with chain bending given

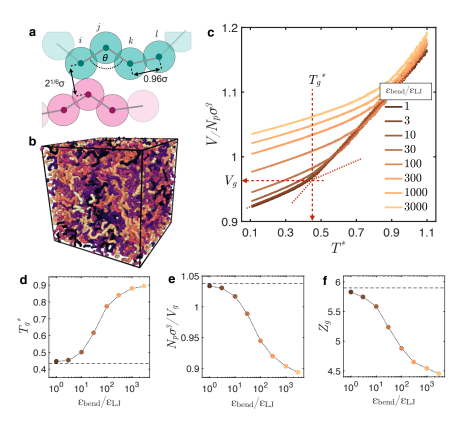


FIG. 1. Entry into the glassy state and its structural properties for $\theta_0=109.5^\circ$. (a) Sketch of polymer chain illustrating the angle θ and rest positions for LJ ($2^{1/6}\sigma$) and FENE (0.96σ) interactions. (b) Snapshot of glassy polymer in periodic box [13]. (c) The decrease of volume associated with decreasing T^* at fixed pressure, for several values of $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. We approximate the low- and high-temperature dependences as linear, and take their intersection to occur at $T^*=T_g^*$. (d) Variation of glass transition temperature T_g^* with $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. (e) Variation of $N_p\sigma^3/V_g$, the density at T_g^* , with $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. (f) Variation of Z_g , the coordination number at T_g^* , with $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. Dashed line in (d)-(f) indicates $\varepsilon_{\rm bend}=0$.

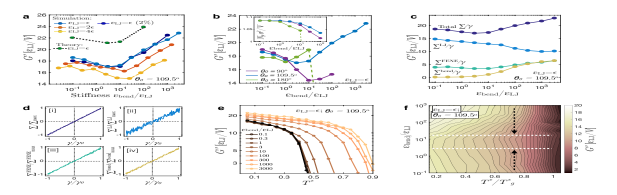


FIG. 2. Nonmonotonic mechanical response of a polymer glass as a function of chain bending stiffness. (a) Elastic modulus G' as a function of $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ for three values of $\varepsilon_{\rm LJ}$. We used five realisations and found the variation between realisations to be smaller than the marker size. (b) Elastic modulus G' as a function of $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ for three values of θ_0 . Inset: decreasing density with increasing $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. (c) Elastic modulus contributions from LJ, FENE and angular potentials for $\varepsilon_{\rm LJ} = \epsilon$ and $\theta_0 = 109.5^{\circ}$. (d) Lissajous-Bowditch plots showing linear elastic stress contributions [i] the total Σ ; [ii] $\Sigma^{\rm LJ}$; [iii] $\Sigma^{\rm FENE}$; and [iv] $\Sigma^{\rm bend}$, each rescaled by their maximal values. Strains are rescaled by the amplitude γ_0 . (e) Temperature dependence of G' across a range of bending stiffnesses $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. (f) Contour plot showing G' as a function of rescaled temperature T^*/T_g^* and $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$. Dashed black arrows indicate decreasing G'; dotted white lines show region of minimal G'.

by $U^{\mathrm{bend}}(\theta) = \varepsilon_{\mathrm{bend}}[1 - \cos(\theta - \theta_0)]$ for energy scale₁₅₂ $\varepsilon_{\mathrm{bend}}$ and rest angle θ_0 . The angle θ is formed between₁₅₃ consecutive monomer triplets along each linear chain,₁₅₄ Fig 1a. A dissipative timescale emerges as $m\sigma^2/\xi\varepsilon_{\mathrm{LJ},155}$ and a thermal timescale emerges as $m\sigma^2/\xi k_B T$. The₁₅₆ state of our system, i.e. whether it is in the melt or glassy₁₅₇ state, is given simply by the ratio of these timescales, as₁₅₈ $T^* = k_B T/\varepsilon_{\mathrm{LJ}}$ [17]. A snapshot of the polymer glass is₁₅₉ given in Fig. 1b.

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Decreasing density with $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ Using periodic¹⁶¹ boundaries we equilibrate the system in a melted state at $T^* = 1.2$, maintaining zero external pressure using a^{162} Nose-Hoover barostat. We then cool the system by de-163 creasing T^* at rate $1/\tau_c$, with $\tau_c \sim \mathcal{O}(10^5)\tau$. Results¹⁶⁴ are presented in Fig. 1 for $\theta_0 = 109.5^{\circ}$. The system un-165 dergoes a decrease in volume V as it is cooled, with a 166 change of slope at $T^* = T_g^*$, Fig. 1c [18]. The coordination of the system is quantified by counting all neighbour-167 ing monomers that are within the repulsive part of the Lennard-Jones, i.e.: $Z=\frac{N_p\sigma^3}{V}\int_0^{2^{1/6}\sigma}g(r)4\pi r^2dr$ where g(r) is the monomer-monomer radial distribution func-168 tion. The glass transition occurs at $T^* = T_g^*$, where $V = V_g$ and $Z = Z_g$. As expected [19], T_g^* increases vith $\varepsilon_{\rm bend}$, with apparent limiting values occurring for for T_g $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ} \to 0$ and $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ} > 10^2$ (Fig. 1d), while 172 the associated density $N_p \sigma^3/V_g$ (Fig. 1e) and coordina-173 tion Z_g (Fig. 1f) decrease. Z_g varies from 5.9 to 4.4,174 close to the expected values when transitioning from a^{175} purely central force network to one bound by bending constraints [4, 20]. A value closer to $Z_g = 4$ is expected₁₇₆ for chain lengths $\gg 10^2$ and for $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ} \to \infty$, while further constraints such as torsional rigidity are expected to lead to further reduction [4]. Thus, adding constraints¹⁷⁷ to the monomers reduces the critical coordination and 178 density of the system. The trends in Figs. 1e,f remain¹⁷⁹ the same at any fixed $T^* < T_g^*$; the shearing simulations described below are run at $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at any fixed $T^*=10^{-3}$ for comparison of the same at $T^*=10^{-3}$ for comparison of the same at $T^*=10^{-3}$ for comparison of $T^*=10^{-3}$ for $T^*=10^{-3}$ for with athermal theory and at higher temperatures to test¹⁸² the robustness of the nonmonotonic response near $T_g^{*,\,\mathrm{183}}$ Further structural description is given in Fig S1.

Nonmonotonic dependence of G' on $\varepsilon_{\mathrm{bend}}/\varepsilon_{\mathrm{LJ}}$ The storage modulus G' is obtained for bending stiffnesses₁₈₇ in the range $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}=0.01\to3000$ and rest angles₁₈₈ $\theta_0 = 90^{\circ}$, 109.5° and 180° by two means. In the first, we use dynamic simulation to apply an oscillatory shear₁₉₀ deformation to the system at $T^* = 10^{-3}$ and zero external pressure, with strain amplitude $\gamma_0 = 1\%$ and period₁₉₂ 200τ . For these parameters the system remains in the linear elastic regime. From the potentials $U^{\rm LJ}$, $U^{\rm FENE}$ and $_{_{104}}^{_{104}}$ U^{bend} described above, we obtain per-monomer forces₁₉₅ as, e.g., $f^{LJ} = -dU^{LJ}/dr$ and compute shear stresses in $_{196}^{12}$ xy (with velocity x, gradient y, vorticity z) according to $\frac{1}{197}$ $\Sigma^{\mathrm{LJ}} = \frac{1}{V} \sum_{n=1}^{N^{\mathrm{LJ}}} r_{x,n} f_{y,n}^{\mathrm{LJ}}$, where N^{LJ} represents the total₁₉₈ number of LJ interactions and r is the vector between₁₉₉ interacting monomers, and similarly for FENE $(\Sigma^{\text{FENE}})_{200}$ and bending (Σ^{bend}) interactions. We take the total₂₀₁ Σ and compute G' from the linear oscillatory stress re-202

sponse in the usual way after $\mathcal{O}(10^2)$ cycles.

In the second, we use the nonaffine lattice dynamics formalism [21–23] to theoretically predict the zero-temperature elastic response from the amorphous structure. The modulus comprises an affine term G_A [24] and a nonaffine correction that originates in the lack of local inversion symmetry of the polymer glass. From the interaction potentials and particle coordinates, we obtain the affine contribution to the elastic modulus as $G_A = \frac{1}{V} \frac{\partial^2 U}{\partial \gamma^2} \Big|_{\gamma \to 0}$, where U is the overall interaction potential energy and γ is the strain amplitude. To obtain the nonaffine contribution, we first construct the Hessian matrix H_{ij} for the system at a given configuration as the second derivative of the energy following Ref [21], where the entries can in general be written as

$$\frac{\partial^2 U(z)}{\partial r_n^a \partial r_m^b} = \underbrace{\frac{d^2 U(z)}{dz^2}}_{\text{stiffness}} \frac{\partial z}{\partial r_n^a} \frac{\partial z}{\partial r_m^b} + \underbrace{\frac{d U(z)}{dz}}_{\text{tension}} \frac{\partial^2 z}{\partial r_n^a \partial r_m^b}. \tag{1}$$

Here z represents a generic argument that in practice is represented by either the monomer-monomer separation r or the angle θ ; we give a detailed form of the corresponding matrix entries in the SI. H_{ij} thus includes stiffness and tension contributions from Lennard-Jones, FENE and angular potentials [25]. The eigenvalue problem $\omega_k^2 m e_i^k = \sum_j H_{ij} e_j^k$ is then solved directly, after which we compute the storage modulus as

$$G'(\Omega) = G_A - \operatorname{Re}\left(\frac{1}{V}\sum_k \frac{\Gamma(\omega_k)}{m\omega_k^2 - m\Omega^2 - i\Omega\nu}\right),$$
 (2)

where $\Gamma(\omega_k)$ is the affine force field correlator, e_i^k , e_j^k are eigenvectors and the sum is over the k eigenvalues of the system.

In Fig. 2a we present G' as a function of bending stiffness from both simulation and theory, for $\theta_0 = 109.5^{\circ}$. Shown are results for three values of $\varepsilon_{\rm LJ}$. We verified that our results are valid throughout the linear elastic regime by repeating the $\varepsilon_{\rm LJ} = \epsilon$ calculations at $\gamma_0 = 2\%$. There is clear nonmonotonic dependence of G' on $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ in all cases, with a minimum in G' occurring at $2 < \varepsilon_{\rm bend}/\varepsilon_{LJ} < 20$. The theoretical prediction provides a strong qualitative match to the simulation result at $\varepsilon_{\rm LI} = \epsilon$, also showing nonomontonic behaviour. In the present article, we limit our discussion of the theoretical approach to its corroboration of the simulation result. Future works will focus on the detailed interpretation of the arising features of the density of vibrational states. When expressed in units of ϵ/V (Fig S2), there is a strong increase of G' with $\varepsilon_{\rm LJ}$ as expected. In units of $\varepsilon_{\rm LJ}/V$, though, G' collapses with a small offset for all $\varepsilon_{\rm LI}$, as expected due to the decreasing relative contribution from FENE bonds in each case. In Fig. 2b we present G' at three rest angles θ_0 . Nonmonotonic behaviour is recovered in each case. For $\theta_0 = 90^{\circ}$, we observe an enhanced minimum, with a substantial reduction in G' of approximately 25%, correlated with a consequent decrease in

density relative to $\theta_0 = 109.5^{\circ}$, Fig. 2b Inset. A shallow₂₆₁ minimum is also observed for $\theta_0 = 180^{\circ}$, though at larger₂₆₂ values of $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ individual chains become rod-like, at₂₆₃ which point both the density and G' of the material have₂₆₄ anomalous behaviour (Fig S4).

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To eludicate the origin of the minimum in G', we de-266 compose the contributions Σ^{LJ} , Σ^{FENE} , Σ^{bend} from the²⁶⁷ simulation result for $\theta_0 = 109.5^{\circ}$ and $\varepsilon_{\rm LJ} = \epsilon$, Fig. 2c,268 verifying that each remains linearly elastic, Fig. 2d. Con-269 sistent with the decrease of $N_p\sigma^3/V_g$ and Z_g with increas-270 ing stiffness (Fig 1e,f), we find a steady decrease in $\Sigma^{\mathrm{LJ}_{271}}$ as stiffness is increased. The increasingly rigid built-in272 three body correlations arising from increasing $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ^{273}}$ necessitate a smaller number of pairwise monomer inter-274 actions for marginal stability, which can be achieved at 275 a lower density, or equivalently at a higher free volume.276 As such the stress contribution from Lennard-Jones in-277 teractions (which is proportional to the packing density) 278 decreases monotonically. Since Σ^{LJ} is the dominant con-279 tribution, this corresponds initially to an overall drop in₂₈₀ G'. As expected, though, we find a monotonic increase₂₈₁ in Σ^{bend} as stiffness is increased, as deformation requires₂₈₂ an increasing energy input to move three-body config-283 urations away from their resting positions. There is a_{284} minor nonmonotonicity observed in the FENE contribu-285 tion to G', with the minimum being attributable to the₂₈₆ removal of LJ interactions allowing minor relaxations of₂₈₇ FENE bonds to their resting positions. This magnitude₂₈₈ of this effect is, though, largely outweighed by the other,

We next test the robustness of the nonmonotonic be-291 haviour away from the low temperature limit, as the292 glass transition temperature T_g^{*} is approached from be-293 low. A plot of G' as a function of temperature is given₂₉₄ in Fig. 2e. For low temperatures, the shear modulus de-295 creases slowly with increasing temperature, until a criti-296 cal value is reached at which point the mechanical rigidity₂₉₇ is lost [20]. Rescaling the temperature axis with the ap-298 propriate values of T_g^* (obtained from Fig 1c), Fig 2f, we₂₉₉ find a good collapse of the loss of rigidity G' as $T^* \to T_q^*$.300 Similarly, we find the minimal G' occurring in the same₃₀₁ range of $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$ as in Fig 2a across temperatures, high-302 lighted by white dotted lines in Fig 2f. The nonmono-303 tonic behaviour of G' thus remains even very close to the 304 glass transition. This raises the question of the mech-305 anism by which marginal stability is achieved in semi-306 flexible polymers at T_g^* , which might extend recent work₃₀₇ in the T=0 limit by Ref [11]. The values of T_g^* and₃₀₈ Z_q vary monotonically with chain bending stiffness be-309 tween asymptotic limits (Fig. 1d,f), yet the mechanical₃₁₀ strength at the glass transition retains a minimum for₃₁₁ intermediate $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$.

In general, therefore, one might expect that any chemi- $_{313}$ cal change that decreases the monomer-monomer coordi- $_{314}$ nation of the system, i.e. adding bending constraints or $_{315}$ frustrating packing by inclusion of plasticisers, will result $_{316}$ in a decrease in the contribution to G' from non-bonding $_{317}$ interactions (represented here as Lennard-Jones). We $_{318}$

have demonstrated here with the $\theta_0 = 90^{\circ}$ case that this might be 'designed for' in practice by adjusting the rest angles of linear chains to enhance this decrease. Conversely, it follows trivially that increasing bending stiffness of polymer chains will generally increase the G' contribution from angular potentials.

The two contributions to the storage modulus G' from non-bonding and bending interactions thus have opposite responses to increases in chain bending stiffness. As a result, there is a competition between these contributions that leads to an overall nonmonotonic dependence of G' on bending stiffness, with there being a minimum in G' at $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}=2\to 20$. Parameter exploration in $\varepsilon_{\rm LJ}$ and θ_0 demonstrate that both the depth and location of the minimum in G' can be tuned by manipulating monomer chemistry, suggesting ways in which one might exploit or suppress the nonmonotonicity. Together, these findings predict that nonmonotonicity in G' is a generic feature across glassy polymeric materials.

Given the monomer chemistry of some novel polymeric system, one might use ab-initio computations to derive coarse-grained forms of the non-bonding and bending interactions, with energy scales that serve as proxies for $\varepsilon_{\rm LJ}$ and $\varepsilon_{\rm bend}$, respectively [26, 27]. Our results here can then serve to guide the synthesis of materials by predicting whether the mechanical response will be in the nonmonotonic region, based on the value of the control parameter $\varepsilon_{\rm bend}/\varepsilon_{\rm LJ}$.

Outlook Nonmonotonic dependence of polymer glass mechanical properties results from two contrasting effects as polymer chain bending stiffness is increased: decreased density (and coordination) as monomer-monomer bending constraints are added; and increased mechanical rigidity of the chains. Our results strongly support this being a general phenomenon, as it is robust all the way up to the glass transition temperature and persists for various sets of model parameters. Since bead-spring models form the basis of much contemporary theory for polymer glasses and their material properties, this finding has broad consequences across polymer physics. Indeed, nonmonotonicity of dynamic quantities with respect to chain length and stiffness is emerging as a widespread feature of polymeric systems in various contexts [28, 29]. It is, so far, difficult to isolate bending stiffness experimentally, since many other factors can influence the mechanical properties. Model systems such as colloidal and granular polymers (CGPs) [30] might be good candidates for verifying our predicted nonmonotonicity, though, as they offer a very high level of control over coarse-grained properties.

The density of vibrational states from which we constructed the theoretical calculation of G' using nonaffine lattice dynamics promises to offer additional insights into the structural and dynamic properties of polymer glasses in future works, both under shear induced yielding [31, 32] and approaching T_g^* , and under imposed pressure [33]. Future work might extend the present finding to coarse-grained potentials that represent more specific

materials [7, 27, 34]. Moreover, the present result repre-332 sents the limit of long chains, while future work might ex-333 plore the minimum chain length required to observe non-334 monotonicity. This is further relevant to colloidal gels,335 where specific adhesive forces have been shown to lead336 to bending moments among small aggregates [35] that337 could influence the rheological properties [36] in an analo-338 gous way to that discussed here. Indeed, returning to the analogy with granular materials, it is not clear whether339 similar nonmonotonicity in G' might be observed exper-340 imentally for increasing particle-particle friction. Recent341 theory [37] suggests otherwise, as endogenous noise gen-342 erated in such packings is responsible for rapidly opening343

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and closing contacts meaning both the friction coefficient and Z are rather poorly defined. Understanding the role of rigidity in the mechanical properties of polymers will be useful in applications as diverse as packing genetic material in cells [38], the structure of polyelectrolyte aggregates [39] and high-rate deformation of advanced materials [40].

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