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Monte Carlo study of multiply crosslinked semiflexible polymer networks

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We present a method to generate realistic, three-dimensional networks of crosslinked semiflexible polymers. The free energy of these networks is obtained from the force-extension characteristics of the individual polymers and their persistent directionality through the crosslinks. A Monte Carlo scheme is employed to obtain isotropic, homogeneous networks that minimize the free energy and for which all of the relevant parameters can be varied: the persistence length and the contour length as well as the crosslinking length may be chosen at will. We also provide an initial survey of the mechanical properties of our networks subjected to shear strains, showing them to display the expected nonlinear stiffening behavior. Also, a key role for non-affinity and its relation to order in the network is uncovered.

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I. INTRODUCTION

Networks of semiflexible polymers have become one of the focal points in current soft-matter research. The reason for this interest is twofold: on the one hand, most relevant structural biological materials, both intra- and extracellular, share the common architecture of crosslinked semiflexible polymer networks. Two archetypical examples are the cytoskeleton and the extracellular matrix. At the same time, there is a widespread realization that semiflexible networks represent an interesting soft-matter system in their own right, outside of any biological context, resulting in a much more fundamental interest in the microscopic and geometrical origins of their mechanical behavior.

The mechanoelastic characteristics of networks of semiflexible polymers have been studied to analyze and characterize different types of these networks, both in vivo [1] and in vitro [2]. The contributions of theory have been many and insightful, but analytical progress has typically only been possible in certain limiting cases where simplifying assumptions may be believed to hold, most notably the assumption of affine deformations [3,4]. At the same time, computer simulations have been used to study these networks, but they too have had to rely on simplifications—either reducing the system to two dimensions and limiting to the small-strain regime [5,6] or ignoring the nonlinear nature of the constituent filaments [7,8].

We believe that the time is right for more realistic numerical modeling of these networks that allows for a detailed microscopic look at the relations between structure, geometry, and mechanical properties. To this end, we present a computer model to simulate these semiflexible polymer networks in three dimensions. Networks are considered to consist of filaments, described as semiflexible polymers. These filaments are crosslinked in various locations, which might induce extra bending of filaments, thus increasing the free energy of the system. We start with a homogeneous, isotropic initial random network with a high free energy and employ a Monte Carlo scheme to relax this network. This approach allows us to generate realistic three-dimensional networks containing hundreds of crosslinks, which are nonetheless well equilibrated and thus represent realistic initial conditions for further mechanical loading in three dimensions. The methodology to generate such networks is the first main result presented in this paper and is described in the first part of the paper (Sec. II).

In the second part of this paper (Sec. III), we subject these networks to shear and analyze their behavior as a function of the network parameters: e.g., the stiffness and the length of the filaments. The results of these computer experiments are compared with experiments to validate our model and yield novel predictions for the mechanical behavior of semiflexible networks.

II. NETWORK GENERATION AND EQUILIBRICATION

We begin our discussion with a detailed look at the generation of our semiflexible networks based on single polymer energies and how the Metropolis Monte Carlo scheme is implemented and adapted for our specific purposes.

A. Network free energy

The networks considered in this paper consist of filaments, which are linked by crosslinks $i=1,\ldots,N_c$. Figure 1 shows a schematic representation of a part of the network, indicating important parameters of the network and the notation used. Each filament is an inextensible semiflexible chain, whose energy in the presence of an external force is given by

$$E_{fil} = \int_0^{l_i} \left( \frac{\kappa}{2} \left( \frac{d\hat{r}(s)}{ds} \right)^2 + \frac{\gamma}{2} \left| \hat{r}(s) \right|^2 \right) ds,$$

where $s$ is the arclength coordinate running along the filament, $\kappa$ is the bending stiffness which is related to the per-

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to-end vectors of neighboring segments along the same filament.

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with the possibility of dangling ends which are discarded
our networks are therefore strictly tetrafunctional—albeit
the segment level but also through crosslinks.

A filament can thus consist of many segments, but is always
a single mechanical entity, satisfying persistence not just at
the segment level but also through crosslinks.

Each crosslink connects segments of two filaments, and
our networks are therefore strictly tetrafunctional—albeit
with the possibility of dangling ends which are discarded (we
do not take steric avoidance into account). Compared to the
other scales in the network, crosslinks are assumed to be
exceedingly small so that their only action, effectively, is to
force a binary bond between two distinct filaments or remote
regions of the same filaments.

In our computer simulations, we store a complete list of
all positions \(r_{ij}\) of the crosslinks, a complete list of the con-
tour lengths \(l_{c,ij}\) of the segments between crosslinks \(i\) and \(j\),
and a connectivity table which lists which segments are
linked by each of the crosslinks. We do not keep track of the
spatial configuration of a segment between two crosslinks.
Instead, we use the exact radial distribution function as com-
puted from Eq. (2) [9] to assign to each segment a contour
length drawn from the radial distribution function computed
at the segment’s end-to-end length and persistence length. In
this manner, we can perform an important part of the full
ensemble sampling in a straightforward manner: different
assignments of the contour lengths correspond to different
realizations of semiflexible networks with a prescribed
spatial distribution of crosslinks. The relative likelihood of a
given distribution of lengths is computed from the free en-
ergy of the resultant network, which we compute as follows.

For a given network realization we partition the free en-
ergy in an internal segment part \(F_2\) and an intersegment part
\(E_S\). As stated above, the internal degrees of freedom of the
segments are integrated out. Thus, we express the free en-
ergy of a segment as a function of the distance between the
crosslinks \(r_{ij}\) and the length of the segment \(l_{c,ij}\). If the
applied force \(f\) in Eq. (1) is positive (i.e., stretching the fila-
ment), \(F_2\) can be computed from Eq. (1) by employing a
 semiflexible analog of the Marko-Siggia interpolation for-
mula [11]; an expression for this is given in the next section.
The semiflexible wormlike chain (WLC) force-extension for-
mula is not particularly accurate for negative forces, as the
filaments quickly assume configurations with considerable
transverse displacements under compressive loading. The
crucial feature of compressive loading, however, is that the
forces involved are always considerably smaller than those
encountered for extensional loads—indeed, this asymmetry
in the force extension curve is responsible for many me-
chanical features of semiflexible networks. For negative
forces, we find that the force extension is adequately de-
scribed by an exponential approach to the asymptote set by
the classical Euler buckling force. Integrating the force-
extension curve yields the following expression for the en-
ergy:

\[
\beta F_2 = \left\{ \begin{array}{ll}
\frac{-9g(r_{ij})^2[5 + 6g(r_{ij})]}{1 + 6g(r_{ij})} & \text{if } f > 0, \\
\frac{-\left(1 + \exp[90g(r_{ij})/\pi^2]\right)[\pi^4 + \pi^2]}{90} & \text{if } f < 0,
\end{array} \right.
\]

(2)

where \(g(r_{ij})\) is the scaled extension given by

\[
g(r_{ij}) = -1/6/l_{c,ij} + 1 + l_{p}r_{ij}/l_{c,ij}^2.
\]

(3)

These equations are not only computationally convenient;
they also provide an excellent fit to the full, analytical force-
extension curves as shown in Fig. 2(a), where we plot the
force versus the scaled extension \(g(r_{ij})\). In addition to the
single-segment force extension, we also need to keep track of
their persistence through crosslinks. There is no analytical
formula for this contribution, and we have therefore simu-
lated many individual filaments to obtain a reliable numeri-
ical expression for this contribution. If the applied force \(f\)
in Eq. (1) is positive (i.e., stretching the filament), it turns out
that we can capture the essential behavior by

\[
\beta E_S = \frac{\theta_{ij}^2}{l_{c,ij} + l_{c,jk}^2},
\]

(4)

where \(l_{c,ij}\) and \(l_{c,jk}\) are the contour lengths of the segments
and \(\theta_{ij}\) is the angle between the two end-to-end vectors of the
segments. Note that this contribution to the total energy is not
accompanied by an entropic contribution, since it is
defined by explicit variables in our network.

To assess the quality of the segment-segment energy
function, we compare the distribution function of this energy
with simulations. We simulate a single wormlike chain of length \(L_w\)
at a fixed temperature and a persistence length \(l_p\), and count the probability \(P_{\text{shk}}(\theta)\) of an angle \(\theta\)
between the vectors \(\vec{r}_{L_w}\) and \(\vec{r}_N\). Here, \(N\) is anywhere
on the chain. Our approximate expression for this probability is $P_{\text{app}}(\theta) \sim N[l_p/(l_{i_1}+l_{i_2})] \theta \exp(-\beta E_\theta(\theta))$, in which the energy $E_\theta$ is given by Eq. (4), and $N[l_p/(l_{i_1}+l_{i_2})]$ is a normalization factor. This histogram is plotted in Fig. 2(b). Although the correspondence is not perfect, this formula does reflect the essentials of the angle distribution, capturing the broadening and shift of its peaks.

In summary, we attribute to a specific network configuration an energy which is the sum of single-segment energies given by Eqs. (2), plus a sum over all segment-pair energies given by Eq. (4), which runs over all pairs of segments belonging to the same filament and meeting in the same crosslinks.

B. Interpolation formula for the segment free energy

Equation (1) enables us to derive an analytic approximation for the semiflexible force-extension relation. To simplify notation, we will pass to dimensionless quantities, rescaling all forces by a factor of $l_p^2/k$ and all lengths by $l_p/l_c$. Based upon Eq. (1), we can express the scaled difference between the total rescaled length of the polymer ($\tilde{l}_c$) and the end-to-end length at rescaled force $\phi$ ($\tilde{l}_\phi$) as

$$\tilde{l}_c - \tilde{l}_\phi = \frac{1}{\pi^2} \sum_{n=1}^{\infty} \frac{1}{n^2 + \phi^2},$$

which gives

$$\tilde{l}_c - \tilde{l}_\phi = -\frac{1 + \sqrt{\phi} \coth \sqrt{\phi}}{2\phi}.$$  

At zero force this gives $\tilde{l}_c - \tilde{l}_\phi = 1/6$. With this, we can define the differential extension at force $\phi$ (i.e., the incremental extension compared to that at zero force) as $\delta \tilde{l} = \tilde{l}_\phi - \tilde{l}_0$. We use these equations to construct an interpolation formula for $\phi(r, l_c, l_p)$, which is the direct analog of the Marko-Siggia interpolation for the WLC formula [11]. Around $\tilde{l}_0$, Eq. (6) gives as a first-order approximation:

$$\phi = 90 \delta \tilde{l}.$$  

In the large force regime we can expand Eq. (5) to yield

$$\phi = \frac{1}{4(1/6 - \delta \tilde{l})^2}.$$  

Tying the two asymptotes together yields

$$\phi = -18 \delta \tilde{l} + \frac{1}{4(1/6 - \delta \tilde{l})^2} - 9,$$  

which can be integrated once to yield Eq. (2). Figure 2(a) shows the comparison between this formula and the exact solution—the difference between the two does not exceed 6%.

C. Network generation

The task at hand is obviously to determine network configurations that minimize the free energy thus defined. To this end, we use a Monte Carlo minimization scheme: starting from an isotropic, random network, we propose random changes in topology, each of which is either accepted or rejected according to the Metropolis criterion.

The initial network is constructed by placing $m$ nodes with random coordinates in a cubic periodic cell. To connect these nodes into a fourfold-coordinated network, we proceed iteratively: we begin by identifying three nodes which are close to each other and connect these with a loop of three bonds. This loop is then extended one bond at a time: we identify a node $A$ which is not fully connected and which is closest to an existing bond $BC$ and then replace this existing bond by two bonds $AB$ and $AC$. This process is repeated until all nodes are fourfold connected.

In the resulting fully fourfold-coordinated network, each bond is considered to be a segment of a single, long filament. This network as a whole can therefore be considered a
FIG. 3. Schematic representation of the three Monte Carlo moves. (a) Four crosslinks that are connected as shown in the above figure are randomly selected in the network. Bonds $AB$ and $CD$ are broken, and bonds $AC$ and $BD$ are created, such that the configuration of the lower figure is formed after energy relaxation. (b) A crosslink is randomly chosen at which crosslinks $A$ and $B$ are part of the same filament, as are crosslinks $C$ and $D$ (above figure). Now $A$ and $C$ become part of the same filament as do $B$ and $D$. This alters the three-crosslink free energy $E_3$. (c) A randomly chosen length $(dl)$ is removed from the length of one segment of a filament and transferred to a neighboring segment of the same filament, such that the configuration of the lower figure is formed after relaxation.

single, circular filament which is crosslinked to itself at various places. We then proceed to minimize the free energy—computed as detailed before—of this initial network, using the standard local minimization method of damped molecular dynamics.

The initial network will be highly stressed and, in general, far removed from a realistic equilibrium configuration. Chiefly, this is due to considerable filament bending, with intrafilament bends at crosslinks often exceeding $90^\circ$. As initial large strides towards an optimal configuration will proceed along downhill directions related to the release of precisely these dominant bending stresses, we first focus on rearranging the topology of the network, analogous to the continuous random network approach, pioneered by Wooten, Winer, and Weaire [12] and further extended and optimized as detailed in [13]. This is realized by a series of Monte Carlo moves that alter the topology; these are moves (a) and (b) in Fig. 3. To the initial configuration with a topology $L'$ with minimized coordinates $\vec{x}'$, we assign a free energy $F'$ as obtained from Eq. (4) plus a quadratic function around the average bond distance to prevent crosslinks from clustering and to tune the final network topology. The average bond distance determines whether the final network will be densely or loosely crosslinked. We then change the topology to $L''$ by one of the moves and relax the network with this new topology, resulting in the coordinates $\vec{x}''$ and a free energy $F''$. Depending on the change in free energy $\Delta F=F''-F'$, the topological change is accepted or rejected, using the Metropolis algorithm. Note that in this stage, we assume that the free energy of a network with minimized crosslink coordinates is representative for the free energy of all networks with the same topology up to some additive constant that is topology independent.

Once such topology-altering moves no longer significantly affect the overall energy—this typically happens in configurations where the bending angle of the filament in each node is on average around $20^\circ$—contour lengths are attributed to the segments. As explained, for a segment $AB$ with end-to-end distance $r_{AB}$, the length $l_{AB}$ is drawn from the corresponding distribution for the WLC formula with the desired persistence length $\ell_p$. Next, we chop up the single continuous filament into many smaller ones by random deletion of segments under the constraint that all crosslinks stay connected up to the point where the desired number of filaments (or, alternatively, mean filament length) is reached. This network is then further equilibrated with the Monte Carlo moves (b) and (c) shown in Fig. 3, each of which is now accepted to a comparable degree. To avoid computational instabilities for floppy filaments we add a short-range repulsive force between crosslinks. A typical network generated with this approach is shown in Fig. 4.

III. MECHANICAL RESPONSE OF THE NETWORK

The ultimate goal is to understand the relationship between the structure of a network and its mechanical properties. In the following sections, we explore some of the basic mechanical properties of our system in an attempt to check whether well-known behavior is correctly reproduced and simultaneously to offer a glimpse of the relevant microscopic processes that we are now able to study in detail and their role in the overall mechanics.

The behavior of biopolymer networks under strain depends on many experimental parameters, such as the concentration of biopolymer, the amount of capping proteins, and the concentration and characteristics of binding proteins. In our simulations, we can reproduce such changes by varying the crosslinking length, the persistence length, and the average number of crosslinks per filament. In this paper we consider networks that consist of $10^3$ crosslinks connecting 2 $\times$ $10^3$ segments. Periodic boundary conditions are applied in all three directions. We do not take into account the contributions of the dangling ends of filaments; nor do we consider
the excluded volume. Our networks are typically very densely crosslinked, which implies that filaments which are close to each other have a high probability to be crosslinked, and we therefore feel it is justified to assume the crosslinking constraints to dominate over the effect of entanglements. Additionally, most naturally occurring biopolymer networks, as well as most in vitro biomimetic experiments considered occur at fairly low polymeric volume fractions, further reducing the importance of excluded volume. The persistence length in the networks we use ranges from $l_p/l_c=1$ to $l_p/l_c=16$ and the average length of filaments from $L/l_c=5$ to $L/l_c=20$. The data graphed in this paper are obtained from single, representative networks as large as practically possible to minimize the effects of finite system size on the observables that we measure.

The experimental techniques used to probe the mechanical response are essentially twofold: on the one hand, in vitro networks are often subjected to global shears in commercial rheometric setups to characterize their macroscopic viscoelastic properties [14,15]. On the other hand, many experiments focus rather on the microscopic processes involved by injecting small particles ($\sim 1 \text{ m}$m) in the network to monitor the behavior at the filament scale of the network [14,17]. Our computational method allows us to work at both levels by direct and simultaneous measurement of the overall stiffness as well as all individual displacements and forces in the system, to high accuracy.

We model shearing by virtually displacing all crosslink positions affinely by small shear-increments of 0.2%. After each shear increment we allow for nonaffine relaxation of all individual crosslinks in order to minimize the free energy of the network. During this procedure, the forces and displacements are recorded and can be used for further analysis of the network response. It is important to note that we allow for full relaxation after each strain increment—this would be appropriate for adiabatically slow shears and should therefore be compared to the zero-frequency limit in oscillatory rheology, as indeed we shall do.

A. Strain stiffening

To characterize our networks, we first consider the differential stiffness $K=\partial \sigma/\partial \gamma$ during shear. An important and characteristic feature of these networks is their highly nonlinear stiffening behavior under relatively small shear stresses [14]. As argued in Ref. [4], all experimental curves of the modulus of networks of semiflexible polymers collapse for small shears on a master curve by scaling the stiffness by the initial stiffness ($K/K_0$) and scaling the shear by its value at which the stiffness is 4 times the initial stiffness ($\gamma/\gamma_4$). Figure 5(a) shows the scaled strain-stiffness curves of our networks under shear, where we plotted the differential modulus $K$, as a function of shear for different ratio’s between $l_p$ and $l_c$ (the average contour length of the segments). We observe the same universal scaled stiffening behavior as observed in experiments. For comparison, we plotted the theoretical curve that incorporates the typical force-extension curve of single filaments combined with the assumption that the filaments deform affinely, which has shown to represent this same master curve. Note that we do not account for rupture and backbone stretching of the filaments, which becomes relevant at larger shears. The inset shows the original curves, where one clearly sees an increase in the initial stiffness as well as a small decrease in the strain at which the networks start to stiffen by increasing the stiffness of individual filaments. In our simulations, $l_p/l_c=16$ is more or less comparable with an actin network with an average persistence length of 15.7, 3.81, and 0.77, of which only the latter is distinguishable at large strains (dotted blue line). Besides, we plotted the scaling from affine theory, which overlaps with the other curves. The inset shows the original strain-stiffness curves, from top to bottom with scaled persistence lengths of 15.7, 3.81, and 0.77. Note that we plot all data points and draw a curve through them. However, a couple of the data points lie well outside the curve (see Sec. III C). (b) Differential modulus of the networks as a function of the scaled shear stress $\sigma$ for 18 networks with varying $l_p/l_c$ and $L/l_c$. Another way to compare our results with experiments is to look at the scaling in the large strain limit. By superposition of a small oscillatory stress on a prestress, the differential modulus can be experimentally measured. From these measurements it is known that $K \sim \sigma^{1.5}$ for large stresses $\sigma$. 

FIG. 5. (Color online) (a) Master curve of scaled differential modulus as a function of scaled shear strain $\gamma/\gamma_4$, where $\gamma_4$ is the strain at which the modulus is 4 times the initial modulus $K_0$. For the curves shown, the average number of crosslinks per filament is $L/l_c=6$. The values for the scaled persistence length $l_p/l_c$ used are 15.7, 3.81, and 0.77, of which only the latter is distinguishable at large strains (dotted blue line). Besides, we plotted the scaling from affine theory, which overlaps with the other curves. The inset shows the original strain-stiffness curves, from top to bottom with scaled persistence lengths of 15.7, 3.81, and 0.77. Note that we plot all data points and draw a curve through them. However, a couple of the data points lie well outside the curve (see Sec. III C). (b) Differential modulus of the networks as a function of the scaled shear stress $\sigma$ for 18 networks with varying $l_p/l_c$ and $L/l_c$. Another way to compare our results with experiments is to look at the scaling in the large strain limit. By superposition of a small oscillatory stress on a prestress, the differential modulus can be experimentally measured. From these measurements it is known that $K \sim \sigma^{1.5}$ for large stresses $\sigma$. 

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influence each others displacements, thus restricting the freedom to adapt to stresses. Besides, when crosslinks connect two or three segments instead of four, these crosslinks are more flexible to reorient when sheared. Therefore, networks with short filaments are softer during shearing.

Figure 6(b) also shows that the stiffness becomes nearly zero for short filaments, a behavior independent of \( l_p \). This decrease is related to the percolation of the network. When the filaments become too short, no real network will be formed. In that case, shearing will shear the liquid in which the filaments are immersed, but the filaments will not be constrained in their movement and thus the stiffness will vanish. Note that we employ a specific procedure to remove material from the network to generate increasingly sparse networks, which implies that the filament length at which the modulus vanishes cannot be directly related to experiments. The overall trend, however, is representative of real networks.

B. Nonaffine behavior and ordering

While the system deforms in our simulations, we allow for nonaffine reorientations of the segments. It has been suggested that such nonaffine deformations greatly alter the mechanical response [7], and indeed we find that this is true. First, a few words about the definition and measures of nonaffinity. In general, an applied macroscopic strain maps any material point \( x \) in the reference space in the network onto a new point \( x' \) in the target space. The location of the point in the target space may be thought of as arising from a combination of an affine deformation and a nonaffine contribution:

\[
    x' = \Lambda(\gamma) x + \Delta(x, \gamma),
\]

where \( \Lambda(\gamma) \) is the deformation gradient tensor, which for the case we consider—three-dimensional simple shear in the \( \hat{e}_1 \) direction—is given by

\[
    \Lambda(\gamma) = \begin{bmatrix}
        1 & \gamma & 0 \\
        0 & 1 & 0 \\
        0 & 0 & 1 
    \end{bmatrix}.
\]

The observation that the nonaffine contribution \( \Delta(x, \gamma) \) depends both on the applied strain and the (original) location of the point under consideration immediately raises the question of what, precisely, it means for a system to be affine. In the strictest sense, an affine system may be defined as one obeying \( \Delta(x, \gamma) = 0 \) for all \( x, \gamma \). This definition, however, is highly restrictive as it does not allow for any nonaffine motion at any point. For systems that do behave nonaffinely to some extent, the most general measure of the extent of this nonaffinity was shown in [20] to be the nonaffinity correlation function

\[
    A_{ij}(x,x'; \gamma, \gamma') = \langle \Delta_i(x, \gamma) \Delta_j(x', \gamma') \rangle,
\]

where the average \( \langle \cdot \cdot \cdot \rangle \) is over all crosslinking points in the network. Both its spatial dependence and the strain dependence are of interest—in a moment we will investigate the strain-dependent aspects by focusing on the trajectory that single points trace out during a deformation. Following [20],

\[\text{(10)}\]

\[\text{(11)}\]

\[\text{(12)}\]
we shall measure to this end the equal argument limit of the trace of $A_{ij}(x,x'; \gamma, \gamma')$, which we shall call simply $\mathcal{A}(\gamma)$:

$$\mathcal{A}(\gamma) = \frac{1}{\gamma^2} \langle |\Delta(x, \gamma)|^2 \rangle. \quad (13)$$

Note that this measure need not approach zero at large strains, even though one may expect all segments to become aligned with the direction of maximal strain in this limit and experience, in effect, a purely extensional strain. The reason $\mathcal{A}(\gamma)$ does not tend to zero lies in the fact that even though the deformation becomes differentially affine, it does not become affine in the absolute sense. To focus on this differential affinity, which we feel is a more appropriate measure of (asymptotic) affinity, we introduce a second measure by considering the differential displacement from the initial point $x_i$ to the final point $x_f$ before and after a small strain increment $d\gamma$:

$$x_f = \Lambda(d\gamma)x_i + \Delta(x_i, \gamma + d\gamma). \quad (14)$$

We use this to define the differential nonaffinity measure

$$\delta\mathcal{A}(\gamma) = \frac{1}{(d\gamma)^2} \langle |\Delta(x, \gamma)|^2 \rangle. \quad (15)$$

This measure does go to zero as $\gamma$ becomes very large. Later, our simulations will show that we do not expect this limit to be attained in experiments as, for realistic parameter values, the system will have failed long before. $\mathcal{A}$ and $\delta\mathcal{A}$ may be expressed in terms of each other, and the latter tending to zero implies that, asymptotically, $\mathcal{A}$ should become constant, with the magnitude of this constant reflecting the overall strength of the past nonaffinity.

Ultimately, we are interested to see to what extent non-affinity affects the mechanical response. To monitor this influence, we perform a shear without relaxation after each strain increment, thus obtaining $K_{\text{affine}}$. Figure 7(a) shows both $K_{\text{affine}}$, which is independent of $L/l_c$ and $K$ for networks with different filament length, all having $l_p/l_c = 1$. As can be seen, even for long filaments, the difference between affine deformation and nonaffine deformation is striking, both for the initial modulus $K_0$ and for the onset of stiffening. This puts the so-called linear (i.e., small-strain) regime of network elasticity in another perspective: even though the strains are small, there is always a finite amount of nonaffinity which greatly affects the overall small strain response. It is thus crucial to understand the role of nonaffinity, even at small strains, to predict the network modulus.

Even though the Hamiltonian of the network remains the same, the difference between the strain-stiffness curves is striking. These differences can only be due to nonaffine behavior of the network, as all other determinants—topology, filament length, density, and persistence length—are identical. There has been some debate whether the origin of stiffening is ultimately entropic or mechanical, but our results suggest that, rather, we should focus our attention on the degree of nonaffinity which acts to delay and attenuate the stiffening.

To see whether our systems tend to affinity at the largest strains, we measure the differential nonaffinity $\delta\mathcal{A}$ as a function of the applied macroscopic strain. To relate $\delta\mathcal{A}$ to other length scales in the system, we plot $\delta\mathcal{A}/r_c^2$, where $r_c$ is the average distance between crosslinks. Figure 7(b) shows a strong increase in $\delta\mathcal{A}$ with increasing strain. From the inset of Fig. 7(b) a strong correlation between the stiffening of the networks and the amount of nonaffine displacements is revealed. Apparently, to prevent the extreme extension that filaments would experience at high strains in an affine setting, the network shows a strong nonaffine reorientation. As indicated before, we expect that for high shear all filaments will be aligned in the direction of shear and deform purely by stretching. Since stretching is an affine deformation, we expect $\delta\mathcal{A}$ to ultimately tend to zero at large strain. This figure and the inset make it clear that this asymptotically (differentially) affine regime is never actually attained, and nonaffinity will continue to feature prominently all the way up to the point of failure.

Shorter filaments, or filaments that are less densely crosslinked, are less constrained in their motions, which should, in principle, allow for greater nonaffine motions. To verify whether indeed this is the case, we evaluate the nonaffinity as a function of filament length. To better compare to existing experiments, we shall use $\bar{\Lambda}$, the overall nonaffinity parameter, instead of the differential measure $\delta\mathcal{A}$. Figure 8(a) indeed shows a pronounced increase in nonaffinity as the...
indicate the ordering during shearing in networks with line indicates the ordering due to affine shearing. The dotted lines initially isotropic material at shear \( \gamma = 0 \) suggest that, asymptotically, we recover the classical picture of rubber elasticity which does very well for long, flexible polymers, but rather poorly for semiflexible systems); as the persistence length decreases, the polymer configurations become increasingly random (i.e., Gaussian), which is accompanied by a decrease in the nonaffinity. This is precisely the rubber limit: Gaussian polymers deforming affinely. Note that these nonaffine deformations are the sole possible origin of the large difference in stiffness between affine and nonaffine deformations shown above. Thus, even though the magnitude of the nonaffine deformations is small, they do have an important effect on the network response. This is a striking example of the value of simulations in this field: microscopic structure and motion are of crucial importance to properly understand the macroscopic behavior.

Apparently, the macroscopic result of the microscopic nonaffinity is to lower the overall stiffness of the system. This suggests an interesting question: if the filaments do not go to their affine positions, where do they go? To begin to answer this, we consider the orientational order of our networks and compute the nematic order parameter \( \omega \), defined as \( \omega = (3 \cos^2 \theta - 1)/2 \). Here the average is taken over all vectors connecting crosslinks that are connected by segments of filaments and \( \theta \) is the angle between such a vector and the average orientation. An isotropic network has \( \omega = 0 \), while a fully ordered network has \( \omega = 1 \). Even when we shear a network affinely, the order will increase from 0 to 1. To appreciate the effect nonaffinity has, we should therefore compare to the affine ordering. This affine ordering is represented by the solid line in Fig. 8(b). The dotted lines show the effect of nonaffine reorientations on the ordering of the network for different filament lengths. Interestingly, nonaffine reorientations tend to increase the order in the network, a behavior independent of \( l_p \).

To get insight into the direction of the ordering, we plot the distribution of the angle \( \phi \) of the end-to-end vectors of segments with respect to the \( x \) axis at \( \gamma = 0.7 \), as shown in Fig. 8(c). By comparing the distribution in a nonaffine network deformation (straight bars) with the distribution of an affine network deformation (dotted bars) we see that the nonaffinity increases the number of segments oriented at a small angle. To appreciate the differences in the two distributions, we plot the analytic expression for the distribution of an initial isotropic medium that is sheared affinely. Interestingly, the maximum of \( P(\phi) \) coincides with the maximal extensional strain experienced as a function of angle. As the figure clearly shows, the additional ordering is in the direction of maximal extensional strain. This might seem counterintuitive—the order appears to be increasing in the direction of increasing filament extensional strain, which would be highly unfavorable from an energetic point of view. However, one should keep in mind that nonaffine motions are not purely rotational: they may encompass additional and simultaneous overall shifts and extensional and compressional components. It would be most interesting to see if this increased order is also observed in experiments. Our simulations suggest that systems containing long filaments are the best place to look for this effect, even though these tend to display lower overall nonaffinity.

So far, we have considered only the nonaffine motion of single points. The nonaffinity correlation function is not only a function of strain; it may also be evaluated for spatially
separated points $x$ and $x'$, To this end, we consider $\delta N(r) = \langle (\bar{r} - \bar{r}_{\text{aff}})^2 \rangle / \delta \gamma^2$, where $\bar{r}$ is the actual vector between two crosslinks and $\bar{r}_{\text{aff}}$ is the vector between the crosslinks if they would have moved affine during $\gamma$. The average, now, runs over all pairs of crosslinks whose separation is $r$.

As explained in Ref. [17], there are two limiting cases in the behavior of $\delta N(r)$. If filaments were stiff rods, the only way to adapt to strain would be by rotating the whole filament. In that case, doubling $r$ would double $\bar{r} - \bar{r}_{\text{aff}}$ and thus $\delta N(r) \sim r^2$. In the other limiting case, segments along a filament behave totally uncorrelated, leading to $\delta N(r) \sim r^0$. The latter is also the limit for $r \to \infty$. However, the net effect of correlated motion of segments along a filament will be highly sensitive to the actual network configuration.

Figure 9(a) shows $\delta N(r)$ of a network at different strains. Note that the larger scatter for small values of $r$ is due to the

Figure 9. (Color online) Correlation $\delta N(r_{rc})$ of nonaffine behavior, as a function of distance $r/r_{rc}$, (a) for strains $\gamma=0.0$ ($K=K_0$, bottom curve), $\gamma=0.2$ ($K=4.0K_0$), $\gamma=0.25$ ($K=7.6K_0$), $\gamma=0.29$ ($K=25K_0$), and $\gamma=0.33$ ($K=260K_0$, top curve); (b) for networks with different $l_p/l_c$ (solid line, $l_p/l_c=0.77$; dash-dotted line, $l_p/l_c=3.81$; and dashed line, $l_p/l_c=15.7$); for all networks, $L/l_c=6$; (c) for networks with different $L/l_c$, ranging from 4 to 20. All curves collapse, except the dash-dotted (brown) curve with $L/l_c=4$. For all networks, $l_p/l_c=15.7$. 

FIG. 10. (Color online) Illustration of a collective reorientation in one of our networks with $l_p/l_c=15.7$ and $L/l_c=6$ during deformation at, respectively, $\gamma=0.268$, $\gamma=0.270$, and $\gamma=0.272$. The thickness of the segment indicates the size of its displacement. The modulus $K$ belonging to the deformation of this network is shown as the upper curve in the inset in Fig. 5(a).
smaller number of pairs of crosslinks. As can be seen, $\delta N(r)$ varies with strain: low strains give a low initial value of $\delta N(r)$ and a steep increase, while high strains show just the opposite. This behavior is strongly correlated to the stiffening behavior shown in Fig. 9(a) (upper line). The observed strain dependence of $\delta N(r)$ is indiscernible when normalizing with respect to $\gamma$ rather than of $\delta y$, which might explain why experiments report no strain dependence [17].

We observe a small but systematic dependence on $l_p$, as shown in Fig. 9(b). Interestingly, thus far we hardly observe any length dependence of the correlation in nonaffine behavior, which is shown in Fig. 9(c) for networks with $l_p/l_c = 15.7$. For filaments with $l_p > l_c$, one would naively expect that the behavior of segments along a filament will be much more correlated than the behavior of segments belonging to different filaments. Thus, one might expect to find increasing spatial correlations for systems composed of larger filaments. That we do not see this behavior suggests that it is approximately balanced by another effect: larger filaments have more links to the rest of the system and are therefore more constrained. While the individual segments along a single filament would like to line up, they become increasingly unable to do so. Interestingly, the first experiments to measure $\delta N(r)$ do show a length dependence [17]. We cannot rule out that we will see this behavior at larger system sizes, but for now are unable to reproduce it.

### C. Collective rearrangements

A closer look at Fig. 5(a) reveals some outliers in the $K$ versus $\gamma$ curve. These discontinuities in $K$ are accompanied by an increase in $A$. This is not a glitch, but rather reflects an interesting microscopic aspect of our networks. To understand this behavior we look at the displacements of individual segments during shearing. Figure 10 shows a network in which the thickness of the segments indicates their displacement during a strain increment of 0.2%. Here we see what happens: during such a strain increment, a significant fraction of the segments has a relatively large incremental displacement in comparison with the average displacement of segments during shear increments.

The noteworthy feature is not so much that there are large displacements, but rather that these displacements are localized and occur in correlated fashion. This is reminiscent of the behavior of so-called collectively rearranging regions, observed in simulation and experiment in glassy systems and colloidal suspensions. These events are rare over the time courses that we have simulated, but may turn out to play an important role in the long-time behavior of these materials. It would be most interesting to check whether these events are also seen in experiments—while these may not be able to resolve the blip in $K$, they might be able to register the accompanying peaks in $A$. The weight in determining $A$ of a reorientation of a certain size decreases with increasing $\gamma$, since $A$ measures the total nonaffinity relative to the total shear. This implies that for small shears, reorientations might induce huge peaks in $A$, while these peaks are absent for larger shears even though the reorientations are still present.

### IV. Conclusions

We have presented a method to generate and deform three-dimensional networks of biopolymer filaments. By an adequate choice of energies both the entropic stiffness of individual segments and the persistence of filaments through crosslinks can be taken into account. By a Monte Carlo thermalization the networks find a local minimum, without further interference from our side.

This method enables us to relate the macroscopic network response to microscopic behavior of individual segments and crosslinks, at both small and large strains. Although a quantitative comparison between experiments and our simulations is hard to obtain, the first results from these simulations agree well with experiments. Both the stiffening during shearing and the length dependence of the nonaffinity are as expected and fit well into the general framework of the behavior of semiflexible polymers. Besides, the stress dependence of the stiffness for large shears is the same as experiments have shown. This confirms that our model captures the right features that decide the network behavior.

Our model proves an excellent tool to compare affine deformations with deformations that allow for nonaffine displacements. We have shown that nonaffine displacements have a large influence on the stiffness of a network and the onset of stiffening. This accounts for the important role of filament length. Besides, the accuracy of analysis of the behavior of the filaments during deformation reveals some surprising results that are hard to obtain by experimental analysis. Thus far unobserved, the nonaffinity increases the order in the network.

Thus far, we have only considered networks of a single type of filaments. Both in cells and in the extracellular matrix, the important load-bearing biopolymer networks are made up of different kinds of filaments: vessel walls are composed of collagen and elastin, and the cytoskeleton too is a composite system containing f-actin, intermediate filaments, and microtubules. This method is a promising tool to explore the behavior of such composite networks under strain, and we are currently exploring their properties.

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