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Effective-medium approach for stiff polymer networks with flexible cross-links

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Recent experiments have demonstrated that the nonlinear elasticity of in vitro networks of the biopolymer actin is dramatically altered in the presence of a flexible cross-linker such as the abundant cytoskeletal protein filamin. The basic principles of such networks remain poorly understood. Here we describe an effective-medium theory of flexibly cross-linked stiff polymer networks. We argue that the response of the cross-links can be fully attributed to entropic stiffening, while softening due to domain unfolding can be ignored. The network is modeled as a collection of randomly oriented rods connected by flexible cross-links to an elastic continuum. This effective medium is treated in a linear elastic limit as well as in a more general framework, in which the medium self-consistently represents the nonlinear network behavior. This model predicts that the nonlinear elastic response sets in at strains proportional to cross-linker length and inversely proportional to filament length. Furthermore, we find that the differential modulus scales linearly with the stress in the stiffening regime. These results are in excellent agreement with bulk rheology data.

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I. INTRODUCTION

The mechanical response and locomotion of living cells is mainly controlled by the cellular cytoskeleton. The cytoskeleton is a highly composite network of various stiff biopolymers, along with various binding proteins for force generation, cross-linking and polymer growth regulation. Understanding the basic physics that governs the mechanical properties of a composite biopolymer network represents an important biophysical challenge that will help elucidate the mechanics of a living cell. In addition to their importance for cell mechanics, cytoskeletal networks have also demonstrated novel rheological properties, especially in numerous in vitro studies [1–10]. However, there have been few theoretical or experimental studies that address the composite nature of the cytoskeleton [11–16]. Recent experiments on F-actin networks with the highly compliant cross-linker filamin, in particular, have demonstrated several striking features: These networks can have a linear modulus as low as 1 Pa, which is significantly lower than for actin gels with incompatible cross-links, and yet they can withstand stresses of 100 Pa or more and can stiffen dramatically by up to a factor of 1000 under applied shear [10,11,16]. Both the linear and nonlinear elastic properties of actin-filamin gels appear to be dramatically affected by the flexible nature of the cross-links, resulting in novel behavior as compared to actin networks with incompatible cross-links, and to synthetic polymer gels. This suggests new network design principles that may be extended to novel synthetic materials with engineered cross-links [12]. However, the basic physics of networks with flexible cross-links remain unclear.

In this article we provide a detailed description of an effective-medium approach to describe the nonlinear elastic properties of composite networks consisting of stiff filaments linked by highly flexible cross-links [15]. A schematic image of the network we aim to model is shown in Fig. 1. The network is composed of randomly oriented filaments or rods of length L, which are linked together by highly flexible cross-linkers. The cross-links consist of two binding domains interconnected by a thermally fluctuating flexible polymer chain of length ℓp. The compliance of such a cross-linker is entropic in nature.

Adopting the wormlike chain (WLC) model, we can fully characterize the cross-linkers with a contour length ℓ0 and a persistence length ℓp [17,18]. The WLC force-extension curve, which is shown in Fig. 2(c), demonstrates the dramatic stiffening of the cross-linker as it reaches its full extension. Indeed, atomic force microscope (AFM) measurements show that an actin cross-linker such as filamin can be accurately described as a WLC [19,20]. At large mechanical loads, however, the experimental force-extension curve deviates significantly from WLC behavior. The polymer chain in cross-linkers such as filamin consists of repeated folded pro-

FIG. 1. (Color online) Schematic figure of an isotropic stiff polymer network with highly compliant cross-linkers.
tein domains, which unfold reversibly at sufficiently large mechanical loads. The experiments by Furuike et al. [20] show that after an initial stiffening regime at a force threshold of ≈100 pN one of the protein domains unfolds reversibly. The accompanied increase in contour length results in a strong decrease in the cross-linkers stiffness. This softening is immediately followed by WLC stiffening as the thermal undulations of the lengthened cross-linker are stretched out. This leads to an elastic response that alternates between entropic stiffening and softening caused by domain unfolding, resulting in a sawtooth force-extension curve.

It has been suggested that the unfolding behavior of filamin is crucial for the mechanical properties of networks with such cross-linkers [11,13,20]. Simulations of stiff polymer networks, assuming a sawtooth force-extension curve for the unfoldable cross-links, reveal that such networks exhibit a fragile state in which a significant fraction of cross-linkers is at the threshold of domain unfolding [13]. This results in strain softening of the network under shear, inconsistent with the pronounced stiffening response observed experimentally in actin-filamin gels [10,11]. We estimate, however, that under typical in vivo experimental conditions, domain unfolding in the cross-links is highly unlikely. For domain unfolding to occur with multiple filamin cross-links experiencing forces of order 100 pN, the resulting tension in the actin filaments is likely to exceed rupture forces of order 300 pN of F actin [21]. Also, a simple estimate of the macroscopic stress corresponding to even a small fraction of filamin under 100 pN tensions is larger than the typical limit of shear stress before network failure is observed. Therefore, we do not expect domain unfolding to occur. Rather, it seems likely that cross-link unbinding occurs before sufficiently large sufficiently large forces are attained for a significant amount of domain unfolding. Detailed estimates based on experiments suggest filamin tensions only of order 1–5 pN at network failure [16]. It has also been shown in single molecule experiments [22] that filamin unbinds from F actin at forces well below the forces required for unbinding, which indicates that cross-linker unfolding is highly unlikely to occur in typical network conditions. Therefore, we consider only the initial stiffening of the cross-links, which we show can account well for the observed nonlinear elasticity of actin-filamin gels.

Our model consists of a network of stiff filaments connected by flexible cross-linkers. The compliance of such a network is expected to be governed by the cross-linkers. The stiff filaments provide connectivity to the network and constrain the deformation of the cross-linkers, thereby setting the length scale of the effective unit cell of the network. Consequently, we expect that the elasticity of the network will be controlled by the filament length L and network connectivity, which is expressed in terms of the number of cross-link per filament n. Therefore, we describe the network with a model in which the basic elastic element consists of a single stiff rod and many compliant cross-linkers that are connected to a surrounding linear elastic medium.

II. EFFECTIVE-MEDIUM APPROACH

Networks of semiflexible polymers with pointlike incompliant cross-links have been studied extensively [4,5,23–26]. These systems exhibit two distinct elastic regimes: One in which the deformation is affine and a regime that is characterized by highly non-affine deformations. The network is said to deform affinely if the strain field is uniform down to the smallest length scale of the network. Simulations [27,28] have shown that the deformation of these networks becomes more affine with increasing cross-link concentration and polymer length, which has been borne out by experiments [4,29]. The elastic response of the network can fully be accounted for by the stretching modes of the polymers in the affine regime. In addition to stretching modes, stiff polymers can also store energy in a non-affine bending mode. Indeed, it has been shown that in sparser networks, in which there are fewer constraints on the constituting polymers, non-affine bending modes dominate the elastic response [24,27,28]. We will, however, not consider the sparse network limit here.

We expect the soft stretching modes of the cross-linkers to govern the elasticity of a dense network of stiff polymers with highly flexible cross-links. However, the large separation in size and stiffness between cross-links and filaments does imply a nonuniform deformation field for the cross-links at the subfilament level. On a coarse-grained level the network deforms affinely and stretches the cross-links as depicted in Fig. 2(b). The network surrounding this particular rod is shown here as a gray background. The deformation of
the cross-links increases linearly from 0 in the center toward a maximum value at the boundaries of the rod. At small strains the cross-links are very soft and follow the deformation of the stiffer surrounding medium. However, at a strain $\gamma \sim \ell_0/L$ the outermost cross-links reach their full extension and, consequently, stiffen dramatically. This suggest the existence of a characteristic strain $\gamma_c$, for the onset of the nonlinear response of the network.

The macroscopic elasticity of the network results from the tensions in all the constituting filaments. The tension in a particular filament can be determined by summing up the forces exerted by the cross-links on one side of the midpoint of the filament. We will employ an effective-medium approach to calculate these forces as a function of filament orientation and the macroscopic strain. Thus, we model the network surrounding one particular rod, as an affinely or uniformly deforming continuum, which effectively represents the elasticity of the network, as depicted in Figs. 2(a) and 2(b). We then proceed by considering contributions from the rods over all orientations to calculate the macroscopic response of the network.

The remainder of this article is organized as follows. First we study a model in which the effective medium is treated as a linear elastic continuum. In this model we will describe the cross-links both as linear springs with finite extensibility, and also as WLC cross-links. We analyze our model in a fully three-dimensional (3D) network, as well as a simplified one-dimensional (1D) representation, which already captures the essential physics of the nonlinear behavior. At large strains, when many of the cross-links are extended well into their nonlinear regimes, it is no longer realistic to model the surrounding network as a linear medium. Therefore, we extend our linear medium model in a self-consistent manner, replacing the embedding medium by a nonlinear effective medium whose elastic properties are determined by those of the constituent rods and linkers. This self-consistent model can quantitatively account for the nonlinear response found in prior experiments on actin-filamin networks [11,16]. Finally, we show how we can compute the tension profiles along the filaments, and we demonstrate how to use these to express the macroscopic stress in terms of the maximum force experienced by a single cross-link.

### III. LINEAR MEDIUM MODEL

We first develop a one dimensional representation of our model, which will be used in Sec. V to construct a more realistic three dimensional model. Also we will restrict the treatment here to a linear description of the effective medium, a constraint that we lift in Sec. IV.

Consider a rigid rod of length $L$ connected by $n$ flexible cross-links to an elastic medium. We shall refer to such an elastic unit as a Hairy Rod (HR). The medium is subject to an externally imposed extensional strain $\epsilon$ parallel to the orientation of the rod. Throughout this paper we denote a 1D extensional strain with $\epsilon$ and a 3D strain with $\gamma$. The presence of the HR in the medium reduces the deformation of the medium at a position $x$ in the rest frame of the rod by an amount $u_{EM}(x, \epsilon) = \epsilon x - u_{cl}(x, \epsilon)$, where $u_{cl}(x, \epsilon)$ is the extension of a cross-linker at a distance $x$ from the center of the rod. The magnitude of $u_{cl}(x, \epsilon)$ and $u_{EM}(x, \epsilon)$ are set by requiring force balance between the cross-links and the medium,

$$f_{cl}[u_{cl}(x, \epsilon)] = K_{EM} u_{EM}(x, \epsilon),$$

where $f_{cl}(u)$ is the force-extension curve of a single cross-linker. The tension $\tau_0$ in the center of the rod is found by summing up the forces exerted by the stretched cross-links on one side of the midpoint of the rod. Assuming a high uniform line density $n/L$ of cross-links along the rod, we can write the sum as an integral

$$\tau_0(\epsilon) = \frac{n}{L} \int_0^{L/2} dx' f_{cl}[u_{cl}(x', \epsilon)].$$

where $u_{cl}(x', \epsilon)$ is obtained by solving Eq. (1). The full tension profile $\tau(\epsilon, x)$ is found by replacing the lower limit of the integration by $x$,

$$\tau(\epsilon, x) = \frac{n}{L} \int_x^{L/2} dx' f_{cl}[u_{cl}(x', \epsilon)].$$

### A. Finely extensible Hookean cross-linkers

We can solve Eqs. (1) and (2) to compute the midpoint tension in a rod, as soon as a force-extension curve for the cross-links is specified. In the absence of unfolding or unbinding, we can describe the force-extension behavior of a flexible cross-linker such as filamin with the WLC model, as depicted with the solid line in Fig. 2(c). It is instructive to simplify the WLC force-extension curve by assuming a Hookean response with a spring constant $k_{cl}$ up to an extension $\ell_0$, which is the molecular weight of the cross-linker. The spring constant $k_{cl} = \frac{k_B T}{\ell_0^2}$ is found from the WLC model for small extensions in the limit $\ell_0 \ll \ell_0 [18]$, where $k_B T$ is the thermal energy. Beyond an extension $\ell_0$, the cross-linker becomes infinitely stiff. The force-extension curve of these finitely extensible Hookean (FEH) cross-links is shown as a dashed curve in Fig. 2(c). The finite extensibility of the cross-links implies a critical strain $\epsilon_c = \frac{\ell_0}{L}$ at which the cross-linkers at the boundaries of the rod reach full extension. For strains $\epsilon \ll \epsilon_c$

$$\tau_0(\epsilon) = \frac{n}{L} \int_0^{L/2} dx' \frac{k_{cl} K_{EM}}{k_{cl} + K_{EM}} \epsilon' \cdot$$

Thus, the midpoint tension depends linearly on strain for $\epsilon \ll \epsilon_c$. For larger strains, the expression for the midpoint tension in a hairy rod in Eq. (2) reads

$$\tau_0(\epsilon) = \frac{n}{L} \int_0^{\epsilon/\epsilon_c} dx' \frac{k_{cl} K_{EM}}{k_{cl} + K_{EM}} \epsilon' + \frac{n}{L} \int_{\epsilon/\epsilon_c}^{L/2} dx' \left[ \frac{k_{cl} K_{EM}}{k_{cl} + K_{EM}} \ell_0 + K_{EM}(\epsilon' - \ell_0) \right].$$

The expression has separated into two integrals, clearly representing a sum over the cross-links with an extension $<\ell_0$
and a sum over the cross-links that have already reached full extension. We also note that beyond $\varepsilon_c$, the midpoint tension depends nonlinearly on strain. Using Eq. (5) we compute the 1D modulus $G_{1D} = \tau_0 / \varepsilon_c$ as shown in Fig. 3. Below the critical strain, the response is dominated by the linear elasticity of the cross-links $G_{1D} = \frac{1}{2} n k_{cl} \ell_0$. The cross-links at the edge of the rod become rigid at a strain threshold $\varepsilon_c = 2 \ell_0 / L$. As the strain is further increased, the outer cross-links stiffen consecutively, resulting in a sharp increase in $G_{1D}$. At large strains, $G_{1D}$ asymptotically approaches a secondary linear regime $\sim \frac{1}{2} n K_{EM} L$.

### B. Wormlike chain cross-linkers

We now consider flexible cross-linkers described by the more realistic WLC force-extension curve, as depicted by the solid line in Fig. 2(c). The force-extension relation is well described by the interpolation formula [18]

$$f_{cl}(u) = \frac{k_B T}{\ell_p} \left( \frac{1}{4(1-\frac{u}{\ell_0})^2} - \frac{1}{4} + \frac{u}{\ell_0} \right),$$

where $k_B$ is Boltzmann’s constant and $T$ is the temperature. This interpolation formula captures the linear and asymptotic stiffening regimes. Additional theoretical work on WLC polymers in the semiflexible limit can be found in Refs. [30–34]. Using Eqs. (1) and (2) we can calculate the 1D modulus $G_{1D}$ for cross-linkers with this force-extension curve. The result of this calculation is shown in Fig. 4. The force-extension curve of the WLC cross-linker is linear up to extensions very close to $\ell_0$, upon which a pronounced stiffening occurs, as shown in Fig. 2(c). We can exploit this, together with the property that for a dense network the medium is much stiffer than the flexible cross-linkers $K_{EM} \gg k_{cl}$, to write an approximate expression for the tension in a hairy rod in a closed form analogous to Eq. (5),

$$\tau_0(\varepsilon) = \frac{n}{L} \int_0^{\ell_0} d\varepsilon' \int_0^{\varepsilon'} du \frac{k_{cl}(u)}{k_{cl}(u) + K_{EM}} + \frac{n}{L} \int_{\ell_0}^{\varepsilon} d\varepsilon' \int_0^{\varepsilon'} du \frac{k_{cl}(u)}{k_{cl}(u) + K_{EM}(\varepsilon' - \ell_0)},$$

where $k_{cl}(u)$ is the differential stiffness $df_{cl}/du$ of the WLC cross-linker. This equation states that an HR unit deforms essentially affinely up to the critical strain. Beyond $\varepsilon_c$, those cross-links that have reached full extension are no longer compliant and start to pull back on the surrounding medium. The approximate calculation of $d\tau_0/d\varepsilon$ is shown together with the exact calculation performed with Eq. (2) in Fig. 4. This graph demonstrates that the approximation captures the essential behavior of the exact curve, and results only in a minor quantitative difference in the crossover regime. Therefore, we will continue constructing our model using this approximation.

The 1D modulus calculated with Eq. (7) is shown for the WLC cross-links together with the results of the FEH cross-links in Fig. 3. Although the main behavior is very similar to that of the FEH cross-linker model, the use of the more realistic WLC force-extension curve has introduced a considerable smoothing of the crossover. The nonlinear behavior in the WLC force-extension curve initiates slowly well before full extension, resulting in a more gradual onset of nonlinear behavior of the HR with WLC cross-linkers. Remarkably, the characteristic strain $\varepsilon_c$ for the nonlinear behavior is proportional to $\ell_0 / L$, independent of the exact nonlinear response of the links.

For a calculation of network mechanics the average tension $\bar{\tau}$ in a filament is more relevant than the midpoint tension $\tau_0$ [35]. $\bar{\tau}$ is found by averaging the tension profile given by Eq. (3) along the backbone of the filament. The ratio $\bar{\tau} / \tau_0$...
is shown in the inset of Fig. 3. We find that over a broad range of strains \( \bar{\varepsilon} = 3/2 \tau_0 \). During the crossover regime the ratio exhibits a peak with an amplitude that depends on the exact ratio of \( K_{EM} \) and \( k_{cl} \).

### IV. Self-consistent-medium model

The linear treatment of the effective medium breaks down at large strains. The network, consisting of a collection of many HRs, will exhibit nonlinear response when the cross-linkers start to get extended into their nonlinear regime. Thus, it is no longer realistic to assume that the effective medium, which should reflect the network elasticity, remains linear. In this section we extend our model by requiring that the elasticity of the background medium self-consistently represents the nonlinear elasticity of the constituent HRs. The elasticity of the medium should therefore depend on the density of filaments and on the elasticity of an HR averaged over all orientations. Thus, we require the stiffness per cross-link of the effective medium \( K_{EM} \) to be determined by the stiffness of an HR,

\[
K_{EM} = \frac{\alpha d\tau_0}{nL \, d\varepsilon}.
\]

The proportionality constant \( \alpha \) depends on the detailed structure of the network. In IV A, we derive an expression for \( \alpha \) in the continuum elastic limit. The midpoint tension in a rod can be written down analogous to Eq. (7),

\[
\tau_0(\varepsilon) = \frac{nL}{\bar{\varepsilon}} \int_0^{L/2} dx' \int_0^\varepsilon d\varepsilon' \frac{\alpha d\tau}{nL \, d\varepsilon} \frac{k_c(x' \varepsilon')}{k_c(x' \varepsilon') + \frac{\alpha d\tau}{nL \, d\varepsilon}} \frac{x' \varepsilon'}{L/2},
\]

where \( k_c(u) \) is the derivative of the force-extension relation of the cross-linker. Note that we have applied the same approximation as we did in Eq. (7). However, we expect this approximation to hold even better here since the medium stiffens strongly as well as the cross-links. Equation (9) can be simplified to the following differential equation for \( \tau_0(\varepsilon) \):

\[
2 \frac{d\tau_0}{d\varepsilon} + \varepsilon \frac{d\tau_0}{d\varepsilon} = \left\{ \begin{array}{ll} 
\frac{nL}{4} \frac{\alpha d\tau_0}{k_c(eL/2)} + \frac{\alpha d\tau_0}{k_c(eL/2)} & \text{if } \varepsilon < \frac{\ell_0}{L/2} \\
\frac{\alpha d\tau_0}{k_c(eL/2)} & \text{if } \varepsilon \geq \frac{\ell_0}{L/2}
\end{array} \right.
\]

We find the following behavior of the model with WLC cross-linkers: Below the characteristic strain for nonlinear response \( \varepsilon_c = 2\ell_0/L \), the tension in a rod depends approximately linearly on strain. This linearity will be reflected in the self-consistent effective medium, and consequently, the model shows behavior similar to the linear medium model up to the critical strain. By solving Eq. (10) we find the midpoint tension \( \tau_0 \) in a rod as a function of extensional strain \( \varepsilon \). Beyond the critical strain the tension depends highly nonlinearly on strain, with a derivative that increases as

\[
\frac{d\tau_0}{d\varepsilon} \sim \varepsilon \alpha. 
\]

Note that unlike in the linear medium model, where the derivative asymptotes to a final value set by \( K_{EM} \), here \( d\tau_0/d\varepsilon \) increases indefinitely. For the FEH cross-linkers we find similar behavior, although in that case the crossover between the linear regime and the asymptotic stiffening regime is more abrupt.

#### A. Continuum elastic limit

Here we derive an expression for \( \alpha \) in the continuum elastic limit. Note that this will only be a good approximation for a dense, isotropic network. The modulus of the medium \( G_{EM} \) can be expressed in terms of the stiffness \( \frac{d\tau_0}{d\varepsilon} \) of a HR by averaging over rod orientations [35,36],

\[
G_{network} = \frac{1}{45} \frac{d\tau}{\rho \, d\varepsilon},
\]

where \( \rho \) is the length of filament per unit volume. \( \rho \) can also be expressed in terms of the mesh-size \( \rho = 1/E \). In the linear medium treatment in Sec. III we found that \( \bar{\varepsilon} = \frac{2}{3} \tau_0 \). Thus, the network modulus reads

\[
G_{network} = \frac{2}{45} \frac{d\tau_0}{\xi^2 \, d\varepsilon}.
\]

We proceed by relating \( K_{EM} \) to \( G_{network} \), which enables us to find an expression for \( \alpha \). Consider a rigid rod of diameter \( a \) and length \( L \), which we use as a microrheological probe in an effective elastic medium with a shear modulus \( G_{EM} \). If the rod is displaced along its axis, it will induce a medium deformation \( \delta \ell \) that leads to a restoring force acting along its backbone. The restoring force per unit length is given by \( 2\pi G_{EM} \log(L/a) \times \delta \ell \). Here we ignore the log term, which is of order \( 2\pi \). Thus, the stiffness of the medium per cross-link \( K_{EM} \) is related to \( G_{EM} \) by

\[
K_{EM} = \frac{L}{n} G_{EM}.
\]

By requiring \( G_{EM} = G_{network} \) we find \( \alpha \) from Eqs. (13) and (14)

\[
\alpha = \frac{2}{45} \frac{L^2}{\xi^2}.
\]

Note that for a dense network \( \alpha \approx 1 \).

### V. 3D network calculation

In this section we describe in detail how the macroscopic mechanical properties of a uniformly deforming network can be inferred from single filament properties. This procedure has been used to describe the viscoelastic [35,36] and nonlinear elastic properties [4,5,8] of semiflexible polymer networks with pointlike rigid cross-links, although a detailed derivation of this theory is still lacking. The main assumption
of this calculation is a uniform, or affine deformation of the network. An affinely deforming polymer chain of length ℓ will stretch or compress, depending on its orientation, by an amount that scales as ~ℓγ. The validity of the affine treatment of cross-linked semiflexible polymer networks has been subject to much debate. Interestingly, two-dimensional (2D) simulations in the zero-temperature limit have found that the deformation can be both affine and non-affine depending on the density of the network and filament rigidity [27,28]. Here we derive the affine theory for the case of a filamentous network with pointlike rigid cross-links. Then we show how this framework can be used together with the effective-medium approach to describe the mechanics of stiff polymer networks with flexible cross-links.

Consider a segment of a filament between two cross-links with an initial orientation ̂ni. When subjected to a deformation described by the Cauchy deformation tensor Λij, this filament segment experiences an extensional strain directed along its backbone

$$\epsilon = |\Lambda ̂n_i| - 1. \quad (16)$$

As before, we denote a 1D extensional strain with ε and a 3D strain with γ. This extensional strain leads either to compression or extension in the polymer segment depending on its orientation, and thus results in a tension τ(Λ̂ni|−1). The contribution of this tension to the macroscopic stress depends also on the orientation of the polymer segment. By integrating over contributions of the tension over all orientations accordingly, we can compute the macroscopic stress tensor σij. We calculate the contribution of the tension in a polymer segment with an initial orientation ̂n as follows. The deformation Λij transforms the orientation of the segment into ̂nj = Λ̂ni/|Λ̂ni|. Thus, the length density of polymers with an orientation ̂ni that cross the j plane is given by $$\rho_{\det \Lambda} \Lambda_{ji} n_i$$, where the factor det Λ accounts for the volume change associated with the deformation. For the network calculations in this article we consider only simple shear, which conserves volume (det Λ = 1). The tension in the i direction in a filament with an initial orientation ̂ni, as it reorients under strain, is τ(Λ̂ni|−1)Λ̂ni/|Λ̂ni|. Thus, the (symmetric) stress tensor reads [5]

$$\sigma_{ij} = \frac{\rho}{\det \Lambda} \left( \tau(\Lambda ̂n_i) - 1 \right) \frac{\Lambda_{ji} n_i}{|\Lambda ̂n_i|}. \quad (17)$$

The angular brackets indicate an average over the initial orientation of the polymer chains.

One remarkable feature follows directly from Eq. (17). A nonlinear force-extension curve for the filaments is not strictly required for a nonlinear network response [37]. To demonstrate this we express the extensional strain of a filament explicitly in terms of the strain tensor γij

$$\epsilon = \sqrt{1 + 2 \gamma ̂n_i ̂n_i} - 1. \quad (18)$$

Thus the extensional strain of a filament depends nonlinearly on the macroscopic strain of the network. Additionally, the reorientation of the filaments under strain leads to an increasingly more anisotropic filament distribution. Remarkably, these geometric effects result in a stiffening of the shear modulus under shear strains of order 1, even in the case of Hookean filaments. At large strains all filaments are effectively oriented in the strain direction, which limits the amount of stiffening to a factor of 4 (2D networks) and 5 (3D networks) over the linear modulus at strains of order 10. Thus the stiffening due to this effect occurs only at large strains and is limited to a factor 5. Therefore we expect this mechanism to have a marginal contribution to the more dramatic stiffening that is observed in biopolymer gels at strains < 1 [4,5]. We would like to stress that the geometric stiffening discussed above has a different nature than the geometric stiffening discussed by [24,38,39]. In their case, the stiffening is attributed to a crossover between an elastic response dominated by soft bending modes in the zero strain limit and a stiffer stretching mode dominated regime at finite strains. In the affine calculation described here, only stretching modes are considered.

By limiting ourselves to a small strain limit, we can exclude the geometric stiffening effects discussed here. This is instructive, since it allows us to study network stiffening due to filament properties alone, and it is a very good approximation for most networks since the nonlinear response typically sets in at strains < 1. For a volume conserving deformation (det Λ = 1) in the small strain limit the stress tensor in Eq. (17) reduces to [36]

$$\sigma_{ij} = \rho(\tau(γ ̂n_i ̂n_i) ̂n_i ̂n_j), \quad (19)$$

In this limit the geometric stiffening mechanism discussed above is absent. Next we show explicitly how to calculate the shear stress σxz, in the z plane for a network, which is sheared in the x direction. A filament segment with an orientation given by the usual spherical coordinates θ and ϕ undergoes an extensional strain

$$\epsilon = \sqrt{1 + 2 \gamma \cos(ϕ) \sin(θ) \cos(θ) + \gamma^2 \cos^2(θ)} - 1$$

$$= \gamma \cos(ϕ) \sin(θ) \cos(θ), \quad (20)$$

where we have used a small strain approximation in the second line. The tension in this segment contributes to the xz component of the stress tensor through a geometric multiplication factor $$\cos(ϕ) \sin(θ) \cos(θ)$$, where the first two terms are due to a projection of the forces in the x direction and the second term is due to a projection of the orientation of the filament into the orientation of the z plane. The stress in the xz direction is thus given by

$$\sigma_{xz} = \frac{\rho}{4\pi} \int_{0}^{\pi} \int_{0}^{2\pi} dθ dϕ \sin(θ)$$

$$\times \{ \tau(\gamma \cos(ϕ) \sin(θ) \cos(θ)) \cos(ϕ) \sin(θ) \cos(θ) \}. \quad (21)$$

Since we limit ourselves to the small strain limit, we do not account for a redistribution of the filament orientations by the shear transformation in this equation.

A. Semiflexible polymer networks with rigid pointlike cross-links

In this section we show how the affine framework can be used to compute the elastic response of a network with inex-
tensile semiflexible polymers connected by pointlike rigid cross-links.

Consider a segment of an inextensible semiflexible polymer of length \( \ell_c \) between two rigid cross-links in the network. Thermal energy induces undulations in the filament, which can be stretched out by an applied tension. By adopting the WLC model in the semiflexible limit \( \ell_c \approx \ell_p \), the force-extension relation of this segment has been shown to be given implicitly by \[ 23 \]

\[
\delta \ell = \frac{\ell_c^2}{\pi^2} \sum_{p=1}^{\infty} \frac{\phi}{n^2(n^2 + \phi)}
\]

where \( \phi \) is the tension \( \tau \) normalized by the buckling force threshold \( \kappa_0^2 \). This relationship can be inverted to obtain the tension as a function of the extension \( \delta \ell \),

\[
\tau = \frac{\pi^2}{\ell_c^4} \phi (\delta \ell / \delta \ell_{\text{max}}),
\]

where \( \delta \ell_{\text{max}} = \frac{1}{6} \ell_c^2 / \ell_p \) is the total stored length due to equilibrium fluctuations. This is also the maximum extension, which can be found from Eq. \( 22 \) as \( \phi \to \infty \). For small extensions \( \delta \ell \) this reduces to

\[
\tau = 90 \frac{k_B T \ell_c^2}{k_p} \delta \ell.
\]

This result can be inserted into Eq. \( 19 \) to find the linear modulus of the network

\[
G_0 = 6 \rho \frac{k_B T \ell_c^2}{k_p}.
\]

For a network in either two or three dimensions, the maximally strained filaments under shear are oriented at a 45 degree angle with respect to the shear plane, meaning that the maximum shear strain is

\[
\gamma_{\text{max}} = \frac{1}{3} \ell_c / \ell_p.
\]

Using the small strain approximation [as in Eq. \( 19 \)], we can calculate the nonlinear network response

\[
\frac{\sigma}{\sigma_c} = \frac{1}{4 \pi^2} \int_0^\pi \int_0^{2\pi} d\theta d\phi \sin(\theta) \left( \phi \left( \bar{\gamma} \cos(\varphi) \sin(\theta) \cos(\theta) \right) \cos(\varphi) \sin(\theta) \cos(\theta) \right),
\]

where we define the critical stress to be \( \sigma_c = \rho \ell_c^2 \). We have also defined \( \bar{\gamma} = \gamma / \gamma_c \), where the critical strain for the network is

\[
\gamma_c = \frac{1}{6} \ell_c / \ell_p.
\]

Equation \( 27 \) demonstrates that the nonlinear response of a network of inextensible semiflexible polymers with rigid cross-links is universal for small strains, as discussed in Ref. [4]. We note, however, that this would not hold if we were to use the full nonlinear theory from Eq. \( 17 \), valid for arbitrarily large strains; geometric effects lead to small departures from universality. This also implies geometric corrections for \( \gamma_c \) [see Eq. \( 28 \)] at high strains. In addition, universality may break down as a result of enthalpic stretching of the polymer backbone [5].

In this section we have assumed that at zero strain all filament segments are at their equilibrium zero-force length. However, cross-linking of thermally fluctuating polymers will result in cross-linking distances both smaller and greater than their equilibrium length. This effect, which is ignored in our discussion here, leads to internal stresses build into the network during the gelation [5].

The universal nonlinear elastic response for a semiflexible polymer network with rigid cross-links is shown in Figs. 5 and 6. The divergence of the differential modulus beyond the critical strain is of the form \( \sim (1 - \gamma_c)^{1/2} \), as depicted in Fig. 5. This results into a power-law stiffening regime of the form \( K \sim \sigma^{1/2} \), as shown in the inset of Fig. 6. This prediction is consistent with experiments on actin gels with the rigid cross-linker scruin [4].

**B. Stiff polymer networks with highly flexible cross-links**

For a network with flexible cross-links we do not consider the tension in filament segments, but rather the average tension \( \bar{T} \) in the whole filament. By using the effective-medium approach we can compute the average tension in a filament as a function of the orientation of the rod and the macroscopic shear strain \( \gamma \). Contributions to the stress from the average tension in the rods are integrated over all orientations according to Eq. \( 21 \). In our description we thus assume affine deformation of the network on length scales \( \gg \ell \). Note, however, that we do not assume that the cross-links deform affinely.
emerges as a consequence of the finite extendability of the cross-links and the nonuniform deformation profile along the backbone of the filament. Remarkably, the power-law exponent is not universal. However, in the dense limit we consider in our model, the deviation to an exponent of 1 is ≲1 and depends only weakly on the ratio \( L/\xi \). As an example, we consider a typical in vitro network for which \( \xi \approx 0.3 \, \mu m \) and the average filament length is \( L=15 \, \mu m \). For this case we find an exponent of 0.98. The asymptotic power-law regime with an exponent \( \approx 1 \), as predicted by our model is consistent with recent experimental data on actin networks cross-linked by filamin [11,16].

The inset of Fig. 6 shows the rigid linker model together with the self-consistent model for a network with flexible cross-links. In this case the stress is normalized by a stress \( \sigma_0 \), which marks the knee of the curve.

VI. TENSION PROFILES AND SINGLE CROSS-LINKER FORCE ESTIMATE

Recently, there has been much debate on the mechanical response of actin binding proteins such as filamin. Specifically, it is discussed whether the cross-links stiffen, unfold, or unbind under tension in both physiological and in vitro conditions. This issue has major implications for the dynamical and mechanical properties of the cytoskeleton. The discussion has been partially resolved recently by single-molecule [22] and bulk rheology [16] experiments on the actin-filamin system. These experiments indicate that cross-links unbind at forces well below the force required for domain unfolding. It is crucial for the bulk rheology experiment, to be able to infer the forces experienced by a single cross-linker from the measured mechanical stress. In this section we show that by using the shape of the tension profile, we can relate a macroscopic quantity such as the stress to the maximum force experienced by a single cross-linker in the network.

The tension along a single filament is not uniform in networks of stiff finite length filaments and incompliant cross-links [27,40]. It was found in simulations that in the affine regime the tension profile is flat close to the midpoint and the tension decreases exponentially toward the boundaries of the filament. In the non-affine regime a different tension profile has been reported, in which the tension decreases linearly toward the ends [39]. In the case of a flexibly cross-linked network of stiff polymers we also expect a nonuniform tension profile, although in this case the underlying physics is different. The deformation of a cross-linker at a distance \( x \) from the midpoint of the rod is \( u_x \approx x/2 \) and, consequently, cross-links further away from the midpoint exert larger forces on the rod, resulting in a nonuniform tension profile.

We can calculate the tension profile for a given rod using Eq. (3). In the limit of highly flexible cross-linkers, the tension profile in the linear elastic regime is given by

\[
\tau(\epsilon, x) = \frac{n}{L k_c} \frac{k_c K_{EM}}{1 + k_c K_{EM}} \left[ \frac{L}{2} - \frac{x^2}{2} \right] \epsilon.
\]
Fig. 7. For low strains we find a parabolic profile, which flattens out toward the edges for larger strains.

We now proceed to estimate the force experienced by a single cross-linker. For an affinely deforming network in the linear-response regime Eq. (17) simplifies to

$$\sigma = \frac{1}{45} p \bar{\tau}(\gamma).$$

(33)

Fig. 7. (Color online) The reduced tension profile along the rod, normalized by the midpoint tension $\tau_0$. This profile is calculated with the self-consistent model with WLC cross-links.

Filaments at a 45° angle with respect to the stress plane bear the largest tension $\bar{\tau}\text{max}$ and experience a strain along their backbone of $\gamma/2$. Assuming linear response we find $\bar{\tau}\text{max}(\gamma) = \bar{\tau}(\gamma)/2$. In the case of a parabolic tension profile, the average tension $\bar{\tau}$ in a filament is related to the largest force $f_0$ experienced by a cross-linker at the boundary of the rod by $\bar{\tau} = \frac{1}{2} \gamma f_0$. Thus, we can express the macroscopic stress in terms of the maximum forces experienced by cross-linkers on the filaments under the greatest load

$$\sigma = \frac{1}{45} pf_{\text{max}}.$$

(34)

For the derivation of this equation we have assumed to be in the linear-response regime. In the nonlinear regime we expect the expression to still hold approximately, although the prefactor will change.

Kasza et al. [16] found that the failure stress of the network $\sigma_{\text{max}}$ is proportional to the number of cross-links per filament $n$ in actin networks with the flexible cross-linker filamin. This suggests that filamin failure, rather than rupture of single actin filaments is the cause for network breakage. In contrast, for actin networks with the rigid cross-linker scruin, which binds more strongly to actin than filamin, rupture of actin was found to be the mechanism for network failure [4].

On the basis of our model and the experimental data from Ref. [16] we estimate filamin failure forces of order 1–5 pN, far below the unfolding force 100 pN. This suggests that network failure is due to filamin unfolding. This is consistent with recent single molecule experiments, which show that filamin unfolding is favored over unfolding of the Ig domains for low loading rates [22].

These numerical estimates for the force experienced by a single filamin cross-linker are for in vitro conditions. Under such conditions actin is present with a concentration of $\sim 1$ mg/ml and filamin is present at an actin to filamin ratio of $\sim 100$. In vivo the concentration of both actin and cross-linkers are believed to be up to an order of magnitude larger [41]. Living cells, however, experience stresses that are of order 1000 pa [42], an order of magnitude larger than the maximum stresses in the in vitro systems in Refs. [10,11]. Hence, the forces experienced by an individual cross-linker in vivo may well be of the same order of magnitude as under in vitro conditions. We note, however, that the actin cytoskeleton is a heterogeneous structure in which the stress is internally generated by motor proteins, which may result in nonuniform stresses. This is not accounted for in our estimate.

VII. IMPLICATIONS AND DISCUSSION

We have studied the nonlinear elasticity of stiff polymer network with highly flexible cross-links. We find that the mechanics of such a network is controlled by network connectivity expressed in the number of cross-links per filament $n$. This was found earlier in experiments on actin-filamin gels [16], providing strong experimental evidence for cross-link dominated mechanics in these networks. Within this picture, stiffening occurs at a strain where the cross-links are stretched toward their full extension. As a result, we expect $\gamma_c$ to be proportional to the molecular weight of the cross-link $\ell_0$. This prediction is consistent with the results of Wagner et al. [12], where cross-link length was varied, while keeping the average filament length fixed. Interestingly, they observed larger values of $\gamma_c$ than expected either from our model or based on Refs. [11,12,16].

In addition, we find here that the filament length $L$ plays an important role in the nonlinear elasticity of these networks. In particular, the onset of nonlinear response $\gamma_c \sim \ell_0/L$ depends crucially on filament length. This has been confirmed by recent experiments on actin-filamin gels, showing an approximate inverse dependence of the $\gamma_c$ on actin filament length [43]. The sensitivity of network response to filament length, both in experiments and in our model, appears to be one of the hallmarks of actin-filamin networks. On the one hand, this may explain the apparent difference between the critical strains reported in Refs. [11,12,16]. On the other hand, this also suggests that it may be even more important in such flexibly cross-linked networks to directly control and measure the filament length distribution than for other in vitro actin studies [44]. Our model does not account for filament length polydispersity. A distribution in filament length is expected to smooth somewhat the sharp stiffening transition predicted by our model.

The dependence of the critical strain for networks with flexible cross-links observed in experiments and predicted by our model is in striking contrast with the behavior found for rigidly cross-linked networks. In the latter case theory predicts $\gamma_c \sim \ell_0/\ell_c$ [see Eq. (28)], which is consistent with experimental observations [4]. The insensitivity of the nonlinear elasticity of dense networks cross-linked with rigid linkers to filament length would suggest that network me-
stretching out of thermal fluctuations of the actin filaments. Consistent with theory for an affine response governed by the nonuniform deformation profile along a filament and the finite extendability of the cross-links, although it is independent of the exact shape of the force-extension behavior of the cross-links. The power-law stiffening $K \sim \sigma^y$ with $y \approx 1$ is consistent with recent experiments on actin-filamin gels [11,16]. This stiffening behavior is very different from the nonlinear response observed for actin gels with rigid cross-links for which a power-law exponent of 3/2 is observed [4], consistent with theory for an affine response governed by the stretching out of thermal fluctuations of the actin filaments. Interestingly, in vivo experiments show that cells also exhibit power-law stiffening with an exponent of 1 [45].

In this article we have examined a limit in which the stiffness of the cross-links is small compared to the stiffness of an F-actin segment between adjacent cross-links. For a large flexible cross-linker such as filamin this is clearly a good approximation in the linear regime. However, as the cross-links stiffen strongly they could, in principle, become as stiff as the actin segment. This would have a dramatic consequence for the nonlinear response of the network. To investigate this we have calculated the differential stiffness $df/du$ as a function of force $f$ for a filamin cross-linker and an actin segment with a length 0.5 to 2 μm, spanning the range of typical distances between cross-links in dense and sparse networks, respectively. This result is shown in Fig. 8. We find the differential stiffness of a filamin cross-link is always smaller than for an F-actin segment, even at large forces in the nonlinear regime. This justifies our approach, in which we have ignored the compliance of the actin, for a broad range of experimentally accessible polymer/cross-linking densities. However, at sufficiently high filamin concentrations, it may be possible that individual network nodes involve multiple cross-linkers, in which case the actin filament compliance may also become relevant. Thus the effect of the compliance of F-actin remains an interesting topic for further research.

We also use our model to study these networks on a more microscopic level, such as the nonuniform tension profiles along the filament backbone. These profiles can be used to establish a relation between the macroscopic stress and the largest force experienced by a single cross-linker in the network. This allows us to estimate the forces experienced by filamin cross-links under typical in vitro and in vivo conditions. We find that the load on these cross-links is not sufficiently high to lead to significant domain unfolding of the filamin Ig domains, even at stresses large enough to rupture the network. Indeed both rheology experiments on actin-filamin gels and single molecule experiments indicate that unbinding occurs well before domain unfolding.

In other large flexible cross-links such as spectrin [46], domain unfolding occurs at lower, more relevant forces. In this case the domain unfolding could have a dramatic effect on the nonlinear viscoelasticity of such networks. In previous work, DiDonna and Levine simulated 2D cross-linked networks, where they have assumed a sawtooth force-extension curve for the cross-linkers to mimic domain unfolding [13]. Their model, however, does not include the dramatic stiffening that is known to occur before unfolding in filamin cross-links. They report a fragile state with shear softening when an appreciable number of cross-linkers are at the threshold of domain unfolding. Our model is based on the stiffening of the cross-linkers, which initiates at forces far below those required for domain unfolding. This leads to strain stiffening at a point where only a fraction of cross-linkers are at their threshold for nonlinear response. Thus in both our model and that of Ref. [13] the network responds strongly to small strain changes, though in an opposite manner: stiffening in the present case vs softening in Ref. [13].

In related work, Dalhaimer et al. showed that isotropic networks linked by large compliant cross-linkers exhibit a shear-induced ordering transition to a nematic phase [14]. It would be interesting to investigate the effect of the nonlinear behavior of the cross-links on this transition. In the present calculation we have assumed an isotropic network. An ordering transition, which results in a strong alignment of filaments will dramatically affect the nonlinear elasticity of the network.

In this article we have studied networks of stiff polymers linked by highly flexible cross-links. Both experiments [11,16] and our model [15] show that these networks have intriguing nonlinear rheological properties. We find that the network mechanics is highly tunable. By varying filament length, cross-linker length and network connectivity we can sensitively regulate the linear and nonlinear elasticity over orders of magnitude. These unique properties can be exploited in the design of novel synthetic materials.
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