Elasticity of 3D networks with rigid filaments and compliant crosslinks

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Disordered filamentous networks with compliant crosslinks exhibit a low linear elastic shear modulus at small strains, but stiffen dramatically at high strains. Experiments have shown that the elastic modulus can increase by up to three orders of magnitude while the networks withstand relatively large stresses without rupturing. Here, we perform an analytical and numerical study on model networks in three dimensions. Our model consists of a collection of randomly oriented rigid filaments connected by flexible crosslinks that are modeled as wormlike chains. Due to zero probability of filament intersection in three dimensions, our model networks are by construction prestressed in terms of initial tension in the crosslinks. We demonstrate how the linear elastic modulus can be related to the prestress in these networks. Under the assumption of affine deformations in the limit of infinite crosslink density, we show analytically that the nonlinear elastic regime in 1- and 2-dimensional networks is characterized by power-law scaling of the elastic modulus with the stress. In contrast, 3-dimensional networks show an exponential dependence of the modulus on stress. Independent of dimensionality, if the crosslink density is finite, we show that the only persistent scaling exponent is that of the single wormlike chain. We further show that there is no qualitative change in the stiffening behavior of filamentous networks even if the filaments are bending-compliant. Consequently, unlike suggested in prior work, the model system studied here cannot provide an explanation for the experimentally observed linear scaling of the modulus with the stress in filamentous networks.

1 Introduction

The mechanical properties of biological cells are governed by the cytoskeleton, a viscoelastic composite consisting of three main types of linear protein polymers: actin, microtubules, and intermediate filaments. These filamentous polymers are cross-linked by various binding proteins and constitute a dynamic complex network that maintains the structural integrity of the cell with the capacity for dynamic reorganization needed for active processes. Many in vitro studies have focused on reconstituted networks with rigid crosslinks.1–11 In the cytoskeleton, however, many of the crosslinks are themselves extended and highly compliant proteins. Such flexible crosslinks can strongly affect the macroscopic network elasticity.11–21 Indeed, experimental studies show that composite networks can have a linear modulus as low as ~1 Pa, while being able to stiffen by up to a factor of 1000.11–16

Here we analyze 3-dimensional (3D) composite networks theoretically, and we offer physical simulations thereof. Our networks are composed of randomly oriented rigid filaments that are connected by highly flexible crosslinks, each of which is modeled as a wormlike chain (WLC),22,23 which has been shown to accurately describe flexible crosslinkers, such as filamin.24,25 In our approach we assume that the filaments are much more rigid than the crosslinks, meaning that the network elasticity is dominated by the entropic stretching resistance of the crosslinks.

In our theoretical analysis we adopt the widely employed assumption of affine deformations.16,19,25 Under this premise, the network is assumed to deform affinely on the length scale of the filaments, which in turn is assumed to be much longer than the contour length of the crosslinks. Using a single filament description in the limit of a continuous distribution of crosslinks along the filament, we obtain the asymptotic scaling behavior of the elastic modulus with the stress in the nonlinear regime. We show that in 1-dimensional (1D) networks, the elastic modulus scales with the second power of the stress, whereas it scales with the third power in 2-dimensional (2D) networks. Remarkably, there is no power law scaling in 3D—in fact, the elastic modulus of a 3D composite network increases exponentially with the stress. Numerical evaluation of the affine theory at finite crosslink densities—as opposed to a continuous distribution of crosslinks—shows that (i) the only asymptotic scaling is the...
modulus scaling with an exponent 3/2 with the stress and that
(ii) the dependence on dimensionality of the system is limited
to an intermediate-stress regime. These findings are in agree-
ment with our extensive physical simulations of 3D composite
networks. For all cases, the elastic modulus diverges at a finite
strain.

Our theoretical analysis is inspired by the mean-field model
proposed by Broedersz et al.19,26 In sharp contrast to our theo-
retical analysis and to the results of our physical simulations,
however, these authors predict linear scaling of the elastic
modulus with applied stress. In particular, for any finite strain,
the elastic modulus remains finite in their model. While this
linear scaling of the elastic modulus is in accordance with what
has been observed experimentally,13,26,21 we here argue that this
model does not adequately capture the elastic response of
networks with rigid filaments and permanent (i.e., non-
rupturing or rebinding) crosslinks of finite length.

In ref. 19, the authors ruled out that the experimentally
observed approximately linear scaling of the modulus with
the stress might be due to enthalpic (linear) stretching compliance
of the crosslinks or filaments. Here, we complement their analysis
by physical simulations that take into account bending of fila-
ments. Our results empirically show that the inclusion of bending
rigidity does not impact the nonlinear stiffening behavior of
composite networks either. We therefore conclude that the
theoretical explanation for the linear scaling of the modulus with
stress in experiments remains a challenging open problem.

By physical simulations, we also study the role of prestress.
We show that in contrast to 1D and 2D networks, 3D networks
experience an initial tension due to non-intersecting filaments
resulting in initially stretched crosslinks, and are therefore
prestressed. The modulus in the linear deformation regime is
then governed by this prestress; indeed, it is higher than the
modulus expected from the affine theory. Our simulations
additionally indicate that if the network is allowed to relax
initial tension by unbinding and rebinding of crosslinks, the
impact of prestress on the elastic modulus in the linear regime
becomes insignificant, although the prestress does not relax all
the way to zero.

The remainder of the article is organized as follows. First, we
present the affine theory of composite networks in Section 2.
Under the assumption that deformations of the network are
affine on the length scale of the filaments, we derive expressions
for the stress and modulus in 1D, 2D, and 3D. We then present
our physical simulation model and describe our network
creation generation procedure in Section 3. We expand on the implica-
tions of our 3D simulation procedure in Section 4; in particular,
we explain the emergence of prestress. We then discuss the
results of our simulations in the linear deformation regime in
Section 5 and indicate which results from the affine theory are
still valid. Finally, we analyze the simulation results in the
nonlinear regime in Section 6.

2 Theory
In this section we analytically derive the stress and modulus of a
composite network under the assumption of affine
deformations on the length scale of the filaments. We consider
a collection of $N$ rigid filaments of length $L$ that are permanently
connected by $nN/2$ flexible crosslinks of contour length $l_0$, where
$n$ is referred to as the crosslink density, i.e., the number of
crosslinks per filament. The filaments are assumed to be
perfectly rigid, i.e., they neither bend nor stretch, and the
crosslinks are modeled via the WLC interpolation formula23

$$f_0(u) = k_B T \left( \frac{1}{l_p} \left( \frac{l_0}{1 - \frac{u}{l_0}} \right)^2 - \frac{1}{4} \left( \frac{u}{l_0} \right)^2 \right).$$ (1)

where $k_B T$ is the thermal energy, $l_p$ is the persistence length and
$u \geq 0$ is the end-to-end distance of the crosslink. Assuming $l_0 >> l_p$ this force-extension relation implements a crosslink rest-
length of zero and shows a characteristic stiffening with diver-
gence of force as $u \to l_0$. Eqn (1) can be integrated to yield the
energy† (up to a constant)

$$E_0(u) = k_B T \left( \frac{l_0}{4} \left( \frac{u}{l_0} \right)^2 - \frac{l_0}{4} \left( \frac{u}{l_0} \right)^2 \right).$$ (2)

Imposing affine deformations on the filament level fully
determines the deformation field $u$ on the subfilament level. In
the following analysis, we consider a single representative fila-
ment subject to an extensional strain of the surrounding
medium that it is embedded in and crosslinked to.

2.1 1D network calculation
We start with a one dimensional system, i.e., 1D extensional
strain $\varepsilon$, and advance in dimensionality by converting an
applied shear strain $\gamma$ into the orientation dependent extensional
strain $\varepsilon(\gamma)$ felt by the filament.

In the rest frame of the filament, the end-to-end distance of a
crosslink at distance $x$ from the center of the filament is given by
$|u(x,\varepsilon)| = |x|$ (see Fig. 1(a)). For notational convenience, we
consider positive $\varepsilon$ only. Under the assumption that the cross-
link density is high enough that one can consider the associated
distribution as uniformly continuous, the total energy of a
filament in 1D is given by

$$E_{1D}(\varepsilon) = 2 \frac{n}{T L} \int_0^{L/2} E_0(\varepsilon x) dx. \quad (3)$$

Substituting eqn (2) into eqn (3), this expression can be inte-
grated analytically (see Appendix A1).

Following the described approach for the linear regime of the
WLC force-extension relation, i.e., for $u < l_0$, the linear
modulus may be extracted as $G_0^{aff} = \frac{2E}{V}$, where $E/V$ is the energy
per unit volume $V$ in the network and $\varepsilon$ is a small strain.27 For a
1D system this yields $G_0^{aff} = \frac{1}{8} \pi n k_B L$, with $k_B = \frac{3}{2} \frac{k_B T}{l_p l_0}$ being the

† More precisely, it is a free energy, which includes both energetic (bending) and
entropic terms for the crosslinks (not for the filaments).
This scaling relation between the modulus and stress in 1D has also been derived in previous work. Next we consider scaling relations in 2D and 3D.

### 2.2 2D network calculation

We perform similar calculations as in 1D, while taking into account that the extensional strain $\epsilon$, which results from a shear strain $\gamma$ on a 2D system, depends on the orientation of the filament under consideration. In the small-strain limit one thus obtains

$$|\epsilon(\gamma, \theta)| = |(\gamma/2) \sin 2\theta|,$$

where $\theta \in [0, \pi]$ is the angle between the filament and the shear direction (see Fig. 1(b)).

Substituting this expression into eqn (4) and averaging over all orientations leads to

$$\langle E_{1D}^{\text{div}}(\gamma) \rangle \sim \frac{1}{\pi/2} \int_0^{\pi/2} \ln \left( \frac{1-\gamma L}{2L_0} \sin 2\theta \right) d\theta,$$

where we assume $\gamma \geq 0$ for notational convenience; the upper integration limit is reduced to $\pi/2$ because $|\sin 2\theta|$ is $\pi/2$-periodic. Note that we do not take into account a redistribution of filament orientations under the shear transformation. This approach, as well as the small-strain approximation for $\epsilon(\gamma, \theta)$, are justified if $L \gg L_0$, since then the strain $\gamma_0 = 4L_0/L$ at which the integrand diverges is small.

Differentiating eqn (7) with respect to $\gamma$ and neglecting the weaker (logarithmically) diverging part of the integrand lead to an expression for the stress, as $\gamma \rightarrow \gamma_0$:

$$\langle \sigma_{2D}(\gamma) \rangle \sim \frac{1}{\pi/2} \int_0^{\pi/2} \frac{d\theta}{1-\left( \frac{\gamma}{\gamma_0} \right) \sin 2\theta},$$

$$= \frac{\pi - \arccos \left( 1 - \frac{\gamma_0}{\gamma} \right)}{\sqrt{1-\left( \frac{\gamma_0}{\gamma} \right)^2}}.$$

The divergence of the stress is of the form $\sigma_{2D} \sim 1/(1-\gamma/\gamma_0)^{1/2}$ and hence $K_{2D} \sim 1/(1-\gamma/\gamma_0)^{1/2}$. Therefore, the asymptotic scaling behavior for the differential modulus in two dimensions is given by

$$K_{2D} \sim (\sigma_{2D})^3.$$

Note the difference of the scaling relations to the ones in the 1D scenario. Stress shows a weaker divergence with strain than in 1D but a stronger dependence on the differential modulus. Integration of the diverging part of the stress further shows that the energy at maximum strain is finite—in contrast to the 1D setting, where the energy diverges at the critical strain. This is an effect introduced by orientational averaging only.

### 2.3 3D network calculation

For a 3D network, the extensional strain on a filament in the small-strain limit is given by
\[ |e(\gamma, \theta, \phi)| = |(\gamma/2)\sin 2\theta \cos \phi|, \]  
\[ (\sigma_{3D})_{s_0}(\gamma) \sim \int_{0}^{\pi/2} \int_{0}^{\pi/2} \frac{\sin \theta \, d\phi d\theta}{1 - (\gamma/\gamma_0) \sin 2\theta \cos \phi}, \]  
with $\gamma \geq 0$; the upper integration limit for the $\phi$ integration is reduced to $\pi/2$ because $|\cos \phi|$ is $\pi$-periodic and symmetric about $\pi/2$ on $[0, \pi]$. If we carry out the $\phi$ integral and expand the integrand around $\theta = \pi/4$, in order to integrate over $\theta$ (see Appendix A.2 for details), we obtain $\sigma_{3D} \sim -\ln(1 - \gamma/\gamma_0)$ and hence $K \sim 1/(1 - \gamma/\gamma_0)$. Consequently, $K$ does not scale with $\sigma$ as a power law; instead, one obtains

\[ K_{3D} \sim e^{c_0\sigma_0}, \]  

with a real constant $c_0$. The absence of asymptotic power law scaling sets 3D networks apart from 1D and 2D networks. In 3D, we observe the weakest (logarithmic) divergence of stress with strain. Integrating the diverging part of the stress shows that the energy again remains finite for $\gamma \to \gamma_d$.

**Finite crosslink density.** By considering the limit of infinite crosslink density, we have derived theoretical scaling relations for strain stiffening by integrating along a filament’s backbone (see eqn (3)). For any real system, however, the crosslink density is finite and eqn (3) turns into a sum

\[ E = \sum_{i=1}^{n} E_0(x_i), \]  

where $\{x_i\}$ are the crosslink binding sites along the filament. Fig. 2 shows numerical results for the behavior of the corresponding differential modulus $K$ for finite $n$, obtained by numerical evaluation of eqn (14) and appropriate orientational averaging. Note that the asymptotic scaling behavior of $K$ in the limit of infinite crosslink density influences a finite network’s behavior in the intermediate-stress regime (see the inset of Fig. 2); however, near the critical strain, the differential modulus scales as $K \sim \sigma^{3/2}$, i.e., like the response of a single WLC. Furthermore, for 1D and 2D systems the theoretical scaling exponents in the limit of infinite crosslink densities can (in the intermediate regime) indeed be approached by increasing $n$. In contrast, as shown above, in 3D the theoretically derived scaling of $K$ is exponential in $\sigma$. Such an exponential increase is quantified by an (in principle) indefinitely increasing maximal slope with increasing $n$ in the $\ln K$ versus $\ln \sigma$ plots; e.g., for $n = 60$ the maximal slope is 3.49, for $n = 3000$ it is 5.82. However, for any finite $n$, eventually there is always universal scaling of $K \sim \sigma^{3/2}$, resulting from the single WLC force-extension relation, independent of the dimensionality of the system. Indeed, for any given $n$, the integral representation eqn (3) becomes invalid close to $\gamma = \gamma_0$ due to the divergence of the WLC energy.

The numerical results in Fig. 2 have been obtained without the small-strain approximation for the extension of the filaments. However, the redistribution of the filament orientations under shear has not been taken into account in Fig. 2. Calculations including this effect show that it may both decrease and increase the maximum intermediate slope in the $\ln K$ versus $\ln \sigma$ plot and shift the peak to larger stress values depending on the maximum strain $\gamma_d$. In any case, the asymptotic scaling regime remains unchanged.

In the next section we introduce the simulation framework that we use to study 3D networks consisting of many filaments and crosslinks, relaxing the assumption of affine deformations.

### 3 Simulation model

We perform quasistatic simulations of 3D networks that consist of $N$ rigid filaments of length $L$, permanently crosslinked by a collection of $nN/2$ crosslinks of length $l_0$. All lengths are measured in units of the side length of the cubic periodic simulation box. A typical set of parameters is $N = 3000$, $L = 0.3$, $n = 60$, and $l_0 = 0.03$.

Each filament is modeled as perfectly rigid, implying that its configuration can be described by its two endpoints only, which are constraint to stay at distance $L$. The flexible crosslinks are modeled as a central force acting between the two binding sites. In particular, we use the WLC interpolation formula (eqn (1)) and the corresponding energy (eqn (2)). In all data that are presented, forces are measured in units of $(k_BT)/l_p$. There are no additional degrees of freedom introduced through the
crosslinks, since their configuration is represented via the endpoints of the filaments, in terms of barycentric coordinates.

In order to generate an initial network configuration we proceed as follows. We generate $N$ randomly distributed filaments by first randomly choosing their centers of mass in our simulation box and by then picking a random orientation for each filament. For crosslinking we apply the following iterative procedure. We randomly choose two points on distinct filaments and insert a crosslink if the corresponding point-to-point distance is shorter than a certain threshold $\alpha l_0$. Here $\alpha \in [0,1)$ serves as a parameter to vary the initially allowed crosslink lengths in the system. This procedure is repeated until the desired number of crosslinks is reached; see Fig. 3 for an illustration of the final configuration. Since we perform quasi-static simulations, the system must be at static equilibrium at all times. As practically all crosslinks will be stretched beyond their rest-length after the initial network generation, we minimize the energy (of the crosslinks) before subjecting the simulation box to any deformation.† For energy minimization we use the freely available external library IPOPT,28 which requires the gradient and the Hessian of the system’s energy function. It might happen during the optimization process that individual crosslinks reach extensions $u$ larger than their contour length $l_0$. Acceptance of these solutions is prohibited by setting the energy to infinity ($10^{19}$) for $u \geq l_0$ in eqn (2); without this modification it would become negative in that regime. The length constraints for the filaments are realized via Lagrange multipliers.

In order to extract elastic properties of the network we perform quasistatic shearing by applying an affine incremental shear strain $\delta \gamma$ to the network, with subsequent rescaling of filaments to length $L$ (see Fig. 1). We apply Lees–Edwards shearing periodic boundary conditions.29 The magnitude of $\delta \gamma$ is determined by calculating the maximum affine shear that leaves all crosslinks below their contour length. Due to the rescaling of filament lengths, a nonaffine deformation component is introduced. This nonaffinity may lead to crosslinks being overstretched after all. In this case, we iteratively halve the shear strain until the length of all crosslinks remains below their contour length. After each shear increment, the energy is minimized. We apply a fixed upper bound of 1% strain on $\delta \gamma$ in order to stay reasonably close to the previous solution. This increases numerical efficiency and accelerates convergence because it allows us to use a warm-start procedure that reuses Lagrange multipliers from one minimization as initial guesses for the next one. Moreover, the application of small shear steps reduces the likelihood of discontinuously jumping between local energy minima.

We stop shearing when the achievable increment in shear strain becomes smaller than a chosen threshold due to crosslinks that are very close to their maximum extension. During the entire simulation process, we record network parameters in the equilibrated states—in particular, the energy $E$ as a function of shear strain $\gamma$. This allows us to extract the shear stress $\sigma = \frac{1}{V} \frac{dE}{d\gamma}$ as well as the differential shear elastic modulus $K = \frac{d\sigma}{d\gamma} = \frac{1}{V} \frac{d^2E}{d\gamma^2}$. Derivatives are taken by first interpolating $E(\gamma)$ with a cubic spline. We define the linear shear elastic modulus as

$$G_0 := K|_{\gamma \rightarrow 0}. \quad (15)$$

In the following section we discuss the implications of our specific simulation model, in particular with respect to the network structure, and contrast it with previous studies that have been carried out mostly in 2D.

### 4 Initial tension and prestress

As mentioned in Section 3, our network generation results in a non-zero initial energy $E_0$ at zero strain. Indeed, by randomly placing (zero-diameter) filaments in a 3D container, filaments have zero probability to intersect; thus, crosslinks have finite initial extension with probability one. This is different from 2D, where randomly placed filaments mutually intersect with a probability approaching one as their number increases. Indeed, so-called Mikado models,19,30–32 where filaments are crosslinked at their intersection sites only, exhibit no forces at zero strain.

In contrast, the initial stretching of crosslinks in our networks results in an initial tension before any deformation. For a quantitative analysis we measure a global variant of this effect by what we call total prestress $\sigma_0$, which measures the

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† We neither take into account fluctuations of the filaments nor excluded-volume effects.
normal stress\(\) component orthogonal to the shear planes. More precisely, we measure the single sided (e.g., upward) normal component of the force that is acting on a given shear plane, by summing up the normal components of the forces exerted by each crosslink and filament passing through the given shear plane, see Fig. 4(a). The normal stress is then given by dividing by the surface area of the shear plane. Note that \(\sigma_0\) does not depend on the choice of a particular shear plane; indeed, if the total stress was changing during vertical movement of a shear plane, then this would immediately contradict force balance in the system.

Intuitively, one might expect negative normal stresses (pulling down on the upper face of the simulation box), since crosslinks are contractile. However, since filaments withstand compression, it is possible to construct systems that exhibit positive normal stress. This suggests the existence of configurations with zero normal stress.\(\)\(\) Indeed, so-called tensegrity structures,\(\) which are in static equilibrium in the absence of boundary conditions satisfy this criterion—while still being able to store arbitrary amounts of energy (see Fig. 4(b)). Empirically, our simulations show that the random networks generated by the procedure described in Section 3 exhibit negative initial normal stresses throughout. Their integrity is provided through the application of periodic boundary conditions. Note, in particular, that our setup enforces conservation of the volume of the simulation box. In general, it would be possible to relax the prestress by letting the volume of the simulation box change. However, we did not follow this approach in the study presented here, in order to ensure that the filament length remains significantly smaller than the size of the simulation box.

In the following, we relate total prestress to the linear elastic response of our networks.

## 5 Linear regime

In previous work,\(^{16,19,26}\) an expression for the linear modulus in 3D was derived under the assumption of affine deformations and in the absence of any initial tension in the network. Our simulations show that the linear elastic modulus depends on the initial tension in the network.

One scenario that clearly demonstrates the dependence of the linear modulus \(G_0\) (defined in eqn (15)) on the initial tension is shown in Fig. 5 where the admissible maximum initial crosslink length was varied.

For a more quantitative analysis we have designed a method that allows us to change initial tension for a network with a fixed set of simulation parameters. We first randomly generate a network as described above and let it relax into static equilibrium. We then remove a given amount (5%) of the most-stretched crosslinks in the system. Then we reconnect those crosslinks randomly again, and let the network relax. This procedure is repeated \(N_{\text{rel}}\) times. Thereby, we successively decrease the system’s initial tension, and therefore also its total energy, see the inset of Fig. 6. Not only does the total energy decrease, we also observe a change in the distribution of forces (see Fig. 6). As long as one performs the crosslink binding–unbinding procedure over a small enough fraction of crosslinks, the network remains nearly isotropic.

![Fig. 5](image_url)  
**Fig. 5** Differential elastic modulus \(K\) as a function of strain \(\gamma\) for different levels of initial tension. The initial tension in the network is varied by changing the initially admissible maximal crosslink length \(a_0\). The linear modulus \(G_0 = K|_{\gamma=0}\) increases with the initial tension in the network (initial tension increases with \(a\)). It is also evident that the divergence of \(K\) occurs at a strain \(\gamma_4\) that decreases with increasing \(a\). Here: \(N = 3000, n = 60, L = 0.3, I_0 = 0.03\).
It is apparent from the inset of Fig. 7 that the linear elastic modulus is reduced by increasing the number of relaxation steps, as expected. Fig. 7 also shows the dependence of linear modulus \( G_0 \) on the total prestress \( \sigma_0 \), which has been introduced in Section 4. We varied \( \sigma_0 \) via the described procedure, and measured \( G_0 \) with the shearing protocol described in Section 3. After a certain number of relaxation steps the empirical value for \( G_0 \) equals the value \( G_0^{\text{aff}} \) expected from affine theory (see Section 2.1). Relaxing initial tension further, we reach moduli even below \( G_0^{\text{aff}} \). This is possible because the network can rearrange nonaffinely, thereby softening its response. Over a certain range of total prestresses, we observe linear scaling of \( G_0 \) with \( \sigma_0 \), a phenomenon, which has been discussed in other contexts before (see for example ref. 35). We explain the linear regime as follows. For small strains the normal component \( \sigma_N \) of the stress acting on shear planes is close in magnitude to the total prestress \( \sigma_0 \), i.e., \( \sigma_N \approx \sigma_0 \). For small strains given by shear angles \( \vartheta = 0 \), total forces acting on the shear planes make an angle \( \varphi \) with the direction normal to the shear planes (see Fig. 8). Our simulations show that \( \tan \varphi \propto \tan \vartheta \) and that the constant of proportionality remains unchanged in the linear scaling regime. Therefore, shear satisfies
\[
\gamma = \tan \vartheta \times \frac{\sigma_S}{\sigma_0},
\]
where \( \sigma_S \) is the component of the stress acting on shear planes in the shear direction, see Fig. 8. Hence, the linear elastic shear modulus \( G_0 \) defined via \( \sigma_S = G_0 \gamma \) is proportional to the total prestress \( \sigma_0 \) via eqn (16). However, for very small total prestresses, i.e., after many relaxation steps, the modulus shows a steeper than linear dependence on \( \sigma_0 \). Indeed, in this regime the aforementioned constant of proportionality becomes larger.

This effect might be attributed to the fact that for small \( \sigma_0 \) tensegrity type elements (see Fig. 4(b)), which do not contribute to the total prestress but carry energy, contribute significantly to the measured shear stress, thereby increasing \( \varphi \) (see Fig. 8).

Furthermore, affine theory predicts linear scaling of the modulus \( G_0 \) with crosslink density \( n \). Fig. 9 shows that this linear scaling is indeed reproduced in our simulations, independent of the prestress. Moreover, by changing the prestress via our relaxation procedure it is possible to reach comparable slopes to what is predicted by the affine theory.

The next section deals with the nonlinear elastic response of the simulated networks, and relates it to the theoretical results that were derived in Section 2.
6 Nonlinear regime

6.1 Critical strain

The networks that we study are inherently nonlinear because crosslinks are WLCs with finite length \( l_0 \) (see eqn (1)), resulting in pronounced strain stiffening at a critical strain \( \gamma_c \). Stress diverges at a higher strain \( \gamma_d \). In our simulations, we define the critical strain \( \gamma_c \) to be the strain where \( K/G_0 \approx 3 \). In the affine theory, \( \gamma_d \) and \( \gamma_c \) scale linearly with the ratio of crosslink to filament length \( l_0/L \). In our simulations, we cannot conclusively report on this dependence because the accessible ranges for \( l_0 \) and \( L \) are quite limited. On the one hand, there exists an upper limit for \( L \) (therefore also for \( l_0 \) since \( l_0/L \ll 1 \) should hold) to be significantly smaller than the simulation box. On the other hand, \( L \) and \( l_0 \) are bounded from below due to computational limitations—this is because we need to increase the number of filaments in order to keep networks homogenous.

For ranges that are accessible to our simulations, we obtain the following results. If we fix \( l_0 \), then we observe linear scaling \( \gamma_c \propto 1/L \) for systems where no relaxation procedure has been applied (see Fig. 10(a)). Relaxed systems, however, sometimes show a less than linear dependence. This effect might be due to anisotropies induced by the relaxation procedure. If we fix \( L \), then the dependence of \( \gamma_c \) on \( l_0 \) is slightly less than linear (see Fig. 10(b)).

6.2 Differential modulus

It remains to discuss the dependence of the differential modulus on stress, the affine theory of which has been derived in Section 2. For finite crosslink densities, the only persistent scaling behavior is \( K \sim \sigma^{3/2} \), as \( \gamma \) approaches \( \gamma_d \)—due to the fact that eventually the single WLC response dominates. In an intermediate regime, above the critical stress \( \sigma_c = \sigma(\gamma_c) \), we observe slopes \( (d \ln K/d \ln \sigma) > 3/2 \). The majority of the simulations shows intermediate slopes around 2 or slightly above, mostly independent of simulation parameters, but there are also realizations that show maximum slopes up to 3.5 (see Fig. 11). These higher slopes and the final scaling \( K \sim \sigma^{3/2} \) are in accordance with the predictions of affine theory. Indeed, a slope of 3.5 is the maximum slope predicted by the affine theory when using the same crosslink density as in the simulation (Fig. 2). There are, however, differences between theory and simulation in terms of slope profiles since various assumptions are made by the theory that do not hold in the simulations: a randomly generated network does not have a uniform crosslink density along the filaments, these systems are prestressed, and there is

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**Fig. 9** Linear elastic modulus \( G_0 \) versus crosslink density \( n \) for systems with different numbers of relaxation steps: \( N_{rel} = 0 \) (diamonds) and \( N_{rel} = 50 \) (squares). The solid line indicates values expected from affine theory: \( G_0^a = \text{const} \). Parameters: \( N = 5000, L = 0.3, l_0 = 0.06 \), and \( \alpha = 0.5 \).

**Fig. 10** (a) Critical strain \( \gamma_c \) versus inverse filament length \( 1/L \) for \( N_{rel} = 0 \) and \( N_{rel} = 50 \). Other parameters: \( N = 5000, n = 60, l_0 = 0.04 \), and \( \alpha = 0.7 \). We observe linear scaling \( \gamma_c \propto 1/L \) for \( N_{rel} = 0 \); systems in which relaxation has been applied show deviations from this behavior (see \( N_{rel} = 50 \) here). (b) Critical strain \( \gamma_c \) versus crosslink contour length \( l_0 \) for a system with \( N = 3000, n = 50, L = 0.3 \), and \( \alpha = 0.5 \).

**Fig. 11** Differential modulus as a function of shear stress, rescaled by linear modulus and critical stress \( \sigma_c = \sigma(\gamma_c) \), respectively. Parameters: \( N = 3000, n = 60, L = 0.3, l_0 = 0.06 \), and \( \alpha = 0.5 \), with \( N_{rel} = 150 \) and without \( N_{rel} = 0 \) relaxation. The inset shows the local slope \( d \ln K/d \ln \sigma \) from the main plot. For large stresses, we observe power law scaling \( K \sim \sigma^{3/2} \) (solid straight line). For intermediate stresses we recover slopes in the range of those derived from affine theory.
no perfect isotropy. Moreover, the networks do not deform perfectly affinely.

6.3 Nonaffinity

In order to study to what extent simulation results deviate from affine theory, apart from prestress, nonuniform crosslink density, and anisotropy, we analyze the nonaffinity of the network deformation under shear. For a single filament, we define its differential nonaffinity with respect to the center of mass by

$$\frac{||\delta r_{\text{aff}} - \delta r||^2}{||\delta r||^2},$$

where $\delta r_{\text{aff}}$ and $\delta r$ are the 3D coordinates of filament’s center of mass after applying an incremental shear strain $\delta \gamma$ without and with relaxation, respectively.

We let $\delta \Gamma$ denote the average of the differential nonaffinities over all filaments. Affine approximations imply $\delta \Gamma = 0$. Fig. 12 shows that center of mass deformations are mostly affine for small strains. However, the differential nonaffinity increases starting at a strain around $\gamma_c$ and eventually diverges as $\gamma \rightarrow \gamma_d$. This can be understood, since the networks are strain stiffening, such that small incremental strain can induce a large increase in the forces of individual crosslinks, thereby inducing local rearrangements during energy minimization.

While increasing shear strain, there are force chains developing in the network, which carry most of the tension, and which cannot reduce their strain due to the fact that they span the entire system (see the inset of Fig. 13). We quantify this effect by considering tension profiles along filaments. The tension $\tau$ at position $x$ along a filament is given via $\tau(x) = \sum_{|x_i| > |x|} f_{\text{el}}(u_i)$, where $x_i$ are the crosslink binding sites and $u_i$ are their extensions ($u_i = e x_i$ in affine theory). Fig. 13 shows tension profiles averaged over all filaments for both, theoretical and simulated systems at various strains. In the simulations there is non-zero tension at zero strain due to prestress. With increasing $\gamma$, the simulations resemble the profiles expected from affine theory. However, when approaching the maximum strain $\gamma_d$, the emergence of selective paths (force chains) that carry most of the tension becomes evident. The highly stretched crosslinks dominate the averaged tension profiles and therefore lead to jumps in the tension curves at the respective binding sites along the filament (green solid curve in Fig. 13).

6.4 Bending

Thus far we have restricted our theory and simulations to rigid filaments that can neither bend nor stretch. In ref. 19, the authors considered finite stretching compliance of filaments, while bending compliance was assumed to be zero. They report that finite stretching stiffness does not impact the nonlinear stiffening regime of a composite network apart from the expected convergence (to some constant value) of the modulus at high strains. Here we complement this analysis by considering filaments that have finite bending but no stretching compliance. We performed simulations on a 2D network because of the relative computational ease compared to the 3D case. In addition to the energy stored in the crosslinks, we consider bending energy of the form $E_b = \kappa \theta^2/(2l_w)$, where $\kappa$ is the bending rigidity, $\theta$ is the angle through which the filaments bend locally, and $l_w = (l_1 + l_0)/2$ is the average distance between two adjacent pairs of crosslinks (Fig. 14). We show the results in
models at present cannot explain the composite networks. These large strains.

Thus, in isolation, neither bending nor stretching compliance of filaments impacts the nonlinear stiffening regime of composite networks. These findings suggest that the theoretical models at present cannot explain the $K \sim \sigma$ scaling observed in experiments.

7 Conclusions

We have studied the elastic properties of composite crosslinked filamentous networks in 3D analytically and numerically. We modeled such networks as a collection of rigid filaments connected by WLC crosslinks.

Based on the affine theory introduced in ref. 19 we derived asymptotic power law scaling exponents for the differential elastic modulus with stress, in the limit of infinite crosslink density. In this case, the scaling exponents depend on the dimensionality of the system. In particular, we showed that 3D systems no longer exhibit a power law. Furthermore, we showed that for finite crosslink densities, the only persistent regime (over several orders of magnitude of stress) is the $\sigma^{3/2}$ scaling, as it is derived from the single WLC force–extension relation, eqn (1). This is in sharp contrast with the model proposed in ref. 16 and 26, where linear scaling was suggested, independent of the dimensionality of the system. This model implies finite stress at any strain and therefore does not apply to composite networks of rigid filaments with flexible crosslinks of finite length.

We further developed a simulation framework that allows us to measure the elastic response of random filamentous networks with WLC crosslinks. One important property of these 3D networks is that, by construction, they are prestressed due to initial extensions of the crosslinks. In addition to geometrical constraints, active elements such as motors can induce prestress as well. We showed that the prestress in a network can dominate the linear response and might therefore be a feature that is worthwhile analyzing in experimental systems.

Regarding the nonlinear response, we observed divergence of stress (and differential modulus) at finite strain. Close to this strain we measured power law scaling of the differential modulus with stress, with an exponent 3/2, just as expected for a single WLC. In an intermediate-stress regime we observed local exponents that span the entire range of theoretically derived values for systems of differing dimensionality. The fact that our simulation results do not always resemble the predictions of a 3D affine theory, in this intermediate regime, may be attributed to nonaffine deformations. Extracting the exact set of assumptions—such as uniform crosslink density, isotropy, or zero prestress—that are responsible for these discrepancies is left for future investigation.

Experiments (see, e.g., ref. 13, 20, and 21) have shown that in the nonlinear regime the differential modulus scales approximately linearly with the shear stress. We did not find such a regime in our simulations—neither when working with rigid filaments nor when incorporating finite bending stiffness (or enthalpic stretching as done in ref. 19). Therefore, we argue that none of the currently available theories can adequately explain the linear scaling of the differential modulus observed experimentally. It could possibly be that the WLC model does not accurately describe the elastic response of a single crosslink throughout the whole experimentally accessible regime. We speculate, however, that the linear scaling might be due to thermal fluctuations of the filaments, which have not been considered so far.

A Derivation of scaling relations for the shear modulus

A.1 1D network

The integral eqn (3) for the total energy of a single filament can be solved to give

$$ E_{1D}(\varepsilon) = 2n \frac{L^2}{L_0^2} \left[ \frac{L^2}{48L_0} - \frac{L^2}{32} - \frac{L^2}{8} - \frac{L^2}{6} \ln \left( 1 - \frac{L}{2L_0} \right) \right]. \quad (18) $$
The divergence of the energy for $\varepsilon \rightarrow \varepsilon_d = 2l_0/L$ stems from the term $\sim \frac{1}{\varepsilon} \ln \left(1 - \frac{\varepsilon}{\varepsilon_d}\right)$, which is therefore the only one that we need to consider for the asymptotic scaling analysis in 2D and 3D.

### A.2 3D network

To approximate the solution of the integral in eqn (12) we first carry out the $\phi$ integration analytically and obtain

$$\langle \sigma_{3D} \rangle_{\theta, \phi}(\gamma) \sim \frac{1}{\gamma} \left[ \frac{1 + (\gamma/\gamma_d) \sin 2\theta}{1 - (\gamma/\gamma_d) \sin 2\theta} \right] \times \sin \theta \, d\theta. $$  \hspace{1cm} (19)

The integral diverges for $\gamma = \gamma_d$ due to a pole at $\theta = \pi/4$. We can approximately consider $\tan^{-1} \left[ \frac{1 + (\gamma/\gamma_d) \sin 2\theta}{1 - (\gamma/\gamma_d) \sin 2\theta} \right] \times \sin \theta$ as a constant because it takes finite values around the pole. Since we are interested in the regime close to the divergence of the integrand, we expand $\sin^2 2\theta$ up to the second order in $\nu : = \theta - \pi/4$. We arrive at

$$\int_{-\pi/4}^{\pi/4} \frac{d\nu}{1 - (\gamma/\gamma_d)^2(1 - 4\nu^2)}. $$  \hspace{1cm} (20)

Approximation errors close to the boundary of the interval of integration that are made by expanding $\sin^2 2\theta$ are negligible, regarding the asymptotics, because the integrand diverges right at the center of the interval. Now we define $\mu : = 1 - \gamma/\gamma_d$ and drop all terms higher than the first order in $\mu$, since we are interested in the behavior close to $\gamma = \gamma_d$. With $\mu^2 : = 4\nu^2$ and $\delta : = 2\mu$, we obtain

$$\int_{-\pi/4}^{\pi/4} \frac{d\eta}{\sqrt{\eta^2(1 - \delta) + \delta}}. $$  \hspace{1cm} (21)

This can be integrated, with the diverging part being

$$\sim \ln \left(2 \sqrt{\eta^2(1 - \delta) + \delta(1 - \delta) + 2(1 - \delta)\eta} \right)_{-\pi/4}^{\pi/4}, $$  \hspace{1cm} (22)

$$\sim -\ln \delta, $$  \hspace{1cm} (23)

$$\sim -\ln(1 - \gamma/\gamma_d), $$  \hspace{1cm} (24)

which is what has been proposed in Section 2.3.

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