Fiber plucking by molecular motors yields large emergent contractility in stiff biopolymer networks

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The mechanical properties of the cell depend crucially on the tension of its cytoskeleton, a biopolymer network that is put under stress by active motor proteins. While the fibrous nature of the network is known to strongly affect the transmission of these forces to the cellular scale, our understanding of this process remains incomplete. Here we investigate the transmission of forces through the network at the individual filament level, and show that active forces can be geometrically amplified as a transverse motor-generated force “plucks” the fiber and induces a nonlinear tension. In stiff and densely connected networks, this tension results in large network-wide tensile stresses that far exceed the expectation drawn from a linear elastic theory. This amplification mechanism competes with a recently characterized network-level amplification due to fiber buckling, suggesting that fiber networks provide several distinct pathways for living systems to amplify their molecular forces.

I. INTRODUCTION

Living organisms use chemical energy to produce the mechanical forces required to move and control their shape. These forces originate at the molecular scale from the power strokes exerted by molecular motors, and are generically transmitted by fiber networks such as the actin cortex, thus resulting in large-scale stresses [1]. To develop a theory for active stress generation, it is thus important to understand both how active forces exerted on an individual filament transmit along this filament to the mesh size level, and are subsequently propagated through the network. Stress generation thus involves force transmission through local one-dimensional objects and higher-dimensional networks.

Tensile forces propagating through biopolymer networks can be much longer-ranged than would be expected from linear elastic force transmission [2–6], and have been proposed to enhance cell-cell mechanical communication [7]. This suggests that fiber networks do more than simply transmitting forces: they amplify them [8]. This amplification of contractile stresses can be accounted for by the large-scale mechanical response of the network surrounding the motors. There, buckling of filaments under compression increases the range of propagation of tensile forces, thus enhancing contractile stresses. In order to generate a significant amplification, this scenario requires active forces that are much larger than the buckling force of individual filaments, which induce buckling in a large neighborhood of the motor. As a result, this mechanism for stress generation is sensitive to the network’s mechanical properties and results in very limited amplification in stiff, densely connected networks.

Here we investigate another stress amplification mechanism that becomes dominant in such stiff networks. This mechanism involves force transmission at smaller scales than the previous one, and we thus consider how forces exerted by a motor on a filament are transmitted by this filament to the rest of the network (Fig. 1a-b). When subjected to a sufficiently large longitudinal motor force, the filament buckles, which at this single-filament level results in a limited force amplification (Fig. 1c). By contrast, a filament under a transverse active force de-
forms and tenses as a plucked string (Fig. 1d), an effect for which we introduce a single-filament model in Sec. II. In this geometry, the force transmitted to the mesh size level consists of the initial active force, plus an additional contractile force dipole due to this plucking tension. In Sec. III, we show that this plucking-induced contractility can become much larger than the one naively expected from the magnitude of the motor force, thus leading to force amplification at the level of a single filament. This geometrical effect is controlled by the deflection angle of the filament, which in turn depends on the stiffness of both the filament and the network: the stiffer the network, the smaller the deflection, the larger the amplification. By explicitly modeling fiber networks involving many filaments, we show in Sec. IV that this results in a strong dependence of the amplification on the network connectivity, which itself controls its stiffness [9]. This study complements our understanding of network-level force amplification to provide a comprehensive theory for force amplification in biopolymer networks.

II. A SINGLE-FILAMENT MODEL FOR FIBER PLUCKING

To account for the large tensions induced in a filament as a “plucking” force is exerted perpendicular to it, we introduce a single-filament model illustrated in Fig. 2a. The plucked fiber consists of two hinged rods acting under extension as Hookean springs with unit rest length and spring constant $\mu$. An active force $F$ is exerted perpendicular to the filament at the hinge. We penalize any bending of the filament by an angle $\theta$ with an energy $2\sin^2(\theta/2)$. The exact form of this penalty is unimportant as long as it goes as $\theta^2/2$ for small angles, which implies that we set the filament’s bending modulus to unity. The surrounding network is modeled as two additional zero-length Hookean springs with spring constant $k$ anchoring the filament to a fixed substrate. The state of the filament is thus represented by two geometrical coordinates $X$ and $Y$, which respectively characterize the filament bending deformation and the network deformation. The energy of the system reads:

$$E = \underbrace{-FX}_{\text{active force}} + \underbrace{\mu(\delta\ell)^2 + 2\sin^2(\theta/2)}_{\text{filament deformation}} + \underbrace{kY^2}_{\text{network response}}$$

with $\delta\ell = \sqrt{X^2 + (1-Y)^2}$ and $\tan(\theta/2) = X/(1-Y)$. In practice, actin filaments are much easier to bend than to stretch, implying $\mu \gg 1$.

III. FORCE AMPLIFICATION AT THE SINGLE FILAMENT LEVEL

To elucidate the magnitude of the contraction generated by our single-filament model, we compute the tensile force $T = kY$ that it transmits to the rest of the network. We first consider the regime of small displacements $(X,Y \ll 1)$. Minimizing over $Y$, we obtain the following force balance equation for $X$:

$$F = \frac{\mu k}{\mu + k} X^3 + X,$$

where we omit numerical prefactors for legibility (their precise values give rise to the numerical factors on the axes of Figs. 2c,d). The role of the filament stretching modulus $\mu$ and the network stiffness $k$ are combined in a single parameter

$$g = \frac{\mu k}{\mu + k} \approx \min(k,\mu)$$

FIG. 2: Single-filament model for filament plucking. a. Model notation, as introduced in Eq. 1. b. Scaling regimes for the nonlinear tension $T$ that a plucked filament transmits to the surrounding network, as a function of the plucking force $F$ and the effective rigidity $g$ of the system defined in Eq. 3. Filled areas indicate different scaling regions: smooth (green), kinked (blue) and folded (white), with inset depicting each situation. The hatched area represents the domain of parameters where the plucking tension $T$ exceeds the applied force $F$, indicating that plucking dominates the forces transmitted to the network. c. The plucking tension (blue line, obtained by minimizing the energy in Eq. 1) as a function of the active force for a soft network with $g = 1$; this tension never exceeds the applied force $F$ (gray line), and the kinked regime does not appear. d. In a stiff network, $k \sim g \gg 1$, all three regimes are present, with a wide regime of force amplification where the blue line is above the grey line.
which corresponds to an effective stiffness of the system, i.e. that associated with the softest mode of deformation, either that of the network or that of the filament. Due to the left-right symmetry of Fig. 2a, the tension in the filament is even in the force, and thus has the nonlinear dependence \( T \approx gX^2 \) in the small elongation limit \( \delta \ell \ll 1 \). We can thus rewrite Eq. 2 in terms of the tension:

\[
g^{1/2}F = T^{1/2}(1 + T)
\]

which relates the plucking tension to the applied force and the mechanical properties of the system. As a result, in the small-displacement regime the value of the plucking tension depends on the system parameters only through the combination \( g^{1/2}F \).

Now considering both small and large displacements, we distinguish between the three amplification regimes illustrated in Fig. 2b, the first two being directly described by Eq. 4:

- at small forces \( F \ll g^{-1/2} \), the filament’s response is dominated by its bending stiffness, and

\[
T \approx gF^2
\]

In this regime, the filament is bent smoothly. If our discrete filament mode were replaced by a continuum one, the same scaling would be obtained and the curvature would be spread over the whole length of the filament.

- at intermediate forces \( g^{-1/2} \ll F \ll g \), the response is dominated by the stretching properties of the filament, and

\[
T \approx g^{1/3}F^{2/3}
\]

note that this non-trivial scaling regime only occurs if \( g \gg 1 \), i.e. for a stiff filament in a stiff network. The filament is kinked at the hinge in this regime. In a continuum filament with a stretching modulus determined by entropic elasticity, this kink would manifest itself as a localization of the curvature in a region of size \( \propto (k_B T/F)^{2/3} \ell_p^{1/3} \) around the point of application of the force, where \( \ell_p \) is the filament persistence length [10].

- at very large force (or for a very soft network) \( F \gg g \), the filament folds around its hinge and bends completely; in that case the plucking tension saturates to a finite value

\[
T \approx g
\]

that is independent of the force \( F \).

To determine whether the effects considered here generate force amplification, we compare the magnitude of the tension \( T \) to that of the force \( F \) applied by the motor. Amplification occurs in both the smooth and kinked regimes, and specifically for \( 1/g \ll F \ll g \) as represented by the hatched area in Fig. 2b. The force amplification ratio \( T/F \) reaches its maximum \( \sqrt{g} \) at the crossover between these two regimes. At this crossover \( T = 1 \), meaning that the tension equals the longitudinal force required to buckle the filament.

**IV. TRANSMISSION OF PLUCKING FORCES THROUGH FULL NETWORKS**

While we have introduced the network rigidity \( k \) as independent from the characteristics of the filament of interest, the filaments constituting the network itself are of the same nature as that filament, which strongly constrains the value of the rigidity parameter \( g \). There is some freedom in setting the value of \( g \) however, as the elasticity of fiber networks is not only controlled by the rigidity of its components, but also by its architecture, and in particular its connectivity. Indeed, densely connected networks primarily deform through stretching modes and have \( k \sim \mu \), while loosely connected networks deform through bending modes, and thus have \( k \sim 1 \) in our units [11]. Here we investigate how these different architectures impact force amplification through plucking.

Going back to our single-filament model, we predict that in soft bending-dominated networks with \( k \sim 1 \) (and thus \( g \sim 1 \)), the plucking tension is small compared to the applied force, regardless of the value of \( F \) (Fig. 2c). In contrast, for stretching-dominated networks with \( k \sim \mu \) (and thus \( g \sim \mu \gg 1 \)), there is a vast regime of force values for which amplification occurs (Fig. 2d).

To put this simple picture to the test, we simulate full networks using a numerical model where filaments are positioned at the edges of a regular two-dimensional lattice. Similar to the model of Fig. 2a, here all filaments are modeled by hinged Hookean springs of unit rest length, with a stretching energy \( \mu (\delta \ell)^2/2 \) associated with an elongation \( \delta \ell \) and a bending penalty \( 2 \sin^2(\theta/2) \) for creating an angular deflection \( \theta \) between two consecutive, initially aligned springs. In addition to their hinges at the lattice nodes, filaments have an additional hinge in between lattice nodes similar to that pictured in Fig. 2a, implying that they buckle if subjected to a longitudinal compressive force larger than the critical force \( F_{\text{buckling}} = 1 \). Distinct filaments are connected at each lattice vertex by crosslinks that do not constrain their relative angles. The energy is minimized with respect to the position of all nodes using the BFGS algorithm. The critical connectivity delimiting the bending- and stretching dominated regimes discussed above can be estimated through a simple constraint counting argument, which indicates that networks where each vertex has more than \( z = 4 \) neighbors in two dimensions are stretching-dominated, where those with lower connectivity are bending-dominated [9]. Here we simulate one network characteristic of each regime: a high-connectivity \((z = 6)\), stretching-dominated triangular lattice (Fig. 3a,b), and a low-connectivity \((z = 3)\),
Plucking is distinct from buckling as a stress amplification mechanism, although they share some characteristics. Indeed, both are nonlinear single-filament effects that rectify all local forces towards contraction while strongly amplifying far-field stresses. These effects are
domination network illustrated in Fig. 3c,d and further described in Appendix A.
To quantify the amount of contractile tension induced by a single motor in a circular piece of network with fixed boundaries, we measure the coarse-grained macroscopic contractile stress (corresponding to the “active stress” of active gel theories [12, 13]), which reads [14]

\[
\sigma = -\frac{1}{V} \sum_i f_i \cdot r_i \tag{8}
\]

where the sum runs over the vertices that belong to the boundary of the network, and \(f_i\) is the force exerted on the boundary at node \(i\) located at position \(r_i\). To assess the contribution of plucking to this contractile stress, we compare situations with and without plucking. The former situation is illustrated in Figs. 3a,c, the motor forces are exerted at the midpoint of each filament. The latter corresponds to forces being applied to the lattice vertices as shown in Figs. 3b,d. We plot the ratio \(\sigma_{\text{pluck}}/\sigma_{\text{nonpluck}}\) of contractile stresses in the two situations in Fig. 3e. For bending-dominated networks, it is roughly equal to one for all force values, indicating that plucking is of little relevance in these systems, as predicted by our local calculation (Fig. 2c). In contrast, in stretching-dominated networks the plucking contribution is large, with a stress ratio culminating at a large value \(\approx 17\). This value is compatible with the expectation drawn from our one-filament model, which predicts an amplification of the order \(\sqrt{\mu}\) with \(\sqrt{\mu} \approx \sqrt{\mu} = 100\) here. This strong amplification is evident when comparing Figs. 3f and g, showing the force transmission pattern in the stretching-dominated network with forces applied in a non-plucking and plucking geometry, respectively. The macroscopic stress of the latter system is 17 times larger than that of the former.

The plucking tension is intrinsically contractile in simple networks, as plucking a filament always tends to shorten its end-to-end distance. As a result, the far-field plucking stress does not depend on the orientation of the force perpendicular to the filament, but only on its magnitude. We illustrate this by reversing the active forces of Fig. 3g in Fig. 3h without any significant alteration of the force transmission pattern. As a result, plucking results in a reversal of locally extensile forces into far-field contractile forces, reminiscent of the buckling-induced rectification reported in Ref. [8].

V. DISCUSSION

Our results show that the generation of large-scale active stress \(\sigma\) in stiff biopolymer networks depends not only on the magnitude \(F\) of the local active forces, but also crucially on the geometry of force exertion. Forces exerted at filament intersections or along filaments induce a linear stress response \(\sigma \propto F\) if the force is smaller than the filaments' buckling threshold. In contrast, forces exerted perpendicularly to a filament far from filament intersections can induce a strongly nonlinear stress response even for forces well below this threshold. The tension resulting from this geometrical effect, which we term plucking, can dominate the far-field response, strongly amplify contractile stresses and reverse locally extensile forces. This plucking effect is generic and independent of spatial dimension.

While plucking can amplify stresses by over an order of magnitude for biologically relevant parameters, it involves several requirements. First, it requires moderate forces, sufficient to bend the filament that they are applied to but not large enough to fold it or buckle its neighbors (Fig. 3e). Second, plucking necessitates a stiff, densely connected networks of flexible filaments, and is negligible in softer, bending-dominated gels. This point was missed in previous studies of local motor-induced network deformations [10]. Beyond the linear networks considered here, stiffening under stress both of individual biopolymers and whole gels could thus enhance stress amplification by plucking.

FIG. 3: Full-network model of plucking. a–d. Local force application geometries. e. Ratio of the stresses generated in the presence vs. absence of plucking in a circular stretching-(bending-)dominated network with radius \(R = 11\). In the gray area buckling occurs, resulting in large displacements and a complex network response involving the effects described in Ref. [8]. f–g. Force propagation in a stretching-dominated network in the non-plucking and plucking geometries, respectively, at force \(F = 0.2\). Blue (red) bonds are under tension (compression). Black arrows indicate forces at a fixed boundary. h. Reversing forces compared to the previous panel leaves the stress pattern almost unchanged. Active force and stress scale are the same in panels f–h. Boundary forces are magnified 5 times in panel f. for legibility. In all simulations \(\mu = 10^4\).
each characterized by specific signatures that could permit to experimentally assess their individual pertinence. Plucking amplifies forces at a purely local level, making it highly sensitive to the local geometry of force exertion. The resulting amplification culminates at intermediate force and increases in magnitude with network stiffness. Buckling, in contrast, is essentially insensitive to local geometry. At larger scales however, buckling nonlinearly amplifies stress through an increased range of force transmission – as in a rope network, rather than an elastic medium. This amplification always increases with increasing force, but decreases with increasing network stiffness as filament buckling becomes more difficult. Despite their differences, buckling and plucking can cooperate: the large local forces emerging from plucking in the kinked regime exceed the buckling threshold, and are thus further amplified by buckling the surrounding network through force transmission.

Plucking and buckling thus provide distinct routes to amplify biological forces. Buckling amplifies forces at any strain, allowing cells in the extracellular matrix to enhance stress transmission and modify the mechanical properties of the matrix over large distances [15]. Plucking, in contrast, generates large stresses only at small strains. It could thus help build tension in the cell’s actomyosin cortex, and has been observed in reconstituted actomyosin networks [16].

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Appendix A: A regular bending-dominated network geometry

As described in the main text, fiber networks explore different elastic regimes depending on their average connectivity. While these different regimes have previously been studied in disordered networks [9], here we avoid the technical difficulties associated with disorder by designing a minimal regular, bending-dominated sublattice of the triangular lattice. Its local structure is illustrated in Fig. 3c,d of the main text. To further characterize the structure and elasticity of this network, we describe its elastic and geometrical response to shear and dilation strain in Fig. 4. All deformation modes presented show non-affine deformations characteristic of bending-dominated networks. As expected for a bending-dominated network, the network’s elastic moduli are of order one, i.e., on par with the filament bending rigidity. By contrast, the moduli of a highly coordinated, stretching-dominated triangular network are of order $\mu \gg 1$.

FIG. 4: Elasticity and geometry of a minimal bending-dominated network. a. Macroscopic stress-strain curves under bulk dilation and shear. b. The network in the undeformed configuration. c. Response to shear. d. Response to uniform dilation. e. Response to uniform compression. Red: compressed bonds, blue: tense bonds. Filament stretching modulus $\mu = 10^3$. 