Buckling and force propagation along intracellular microtubules

Moumita Das\textsuperscript{1,2(a)}, Alex J. Levine\textsuperscript{3} and F. C. MacKintosh\textsuperscript{1,2}

\textsuperscript{1} Department of Physics and Astronomy, Vrije Universiteit - Amsterdam, The Netherlands, EU
\textsuperscript{2} The Aspen Center for Physics - Aspen, CO 81611, USA
\textsuperscript{3} Department of Chemistry and Biochemistry, and University of California - Los Angeles, CA 90095, USA

California Nanosystems Institute, University of California - Los Angeles, CA 90095, USA

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Abstract – Motivated by recent experiments showing the compressive buckling of microtubules in cells, we study theoretically the mechanical response of and force propagation along elastic filaments embedded in a non-linear elastic medium. We find that embedded microtubules buckle when their compressive load exceeds a critical value $f_c$, and that the resulting deformation is restricted to a penetration depth that depends on both the non-linear material properties of the surrounding cytoskeleton, as well as the direct coupling of the microtubule to the cytoskeleton. The deformation amplitude depends on the applied load $f > f_c$ as $(f - f_c)^{1/2}$. This work shows how the range of compressive force transmission by microtubules can be tens of microns and is governed by the mechanical coupling to the surrounding cytoskeleton.

The mechanical response of most eukaryotic cells depends on their cytoskeleton, a composite network of filamentous proteins \cite{1}. Microtubules (MTs) are the stiffest of these cytoskeletal filaments, and they play an important role in organization of, and transport within the cell. Their mechanical rigidity allows them to support significant stresses in the cytoplasm. These stresses can be highly inhomogeneous, with compressive/tensile forces directed along stiff MTs, permitting directed force transmission and mechanical signaling over several microns within the cell. As with macroscopic elastic rods, however, even the comparatively rigid MTs cannot, on their own, withstand as large compressive loads as tensile loads. This is because of the classical Euler buckling instability limiting the compressive force to a maximum value, which actually vanishes for long rods. It was recently shown, however, that even long MTs can bear large compressive loads, as a result of their coupling to the surrounding elastic matrix of the cytoskeleton \cite{2}. This composite aspect of the cytoskeleton has important consequences for cell mechanics and mechanotransduction \cite{3–7} —the generation, transmission, and sensing of forces by the cell.

Here, we develop a model for compressively loaded elastic filaments such as MTs embedded in an elastic continuum. When their compressive load $f$ exceeds a critical force $f_c$, an oscillatory buckling of the filament is expected, with a wavelength depending on both the stiffness of the elastic filament and the shear modulus of the surrounding medium \cite{2,8,9}. In the classical Euler buckling problem, and even in the presence of a (linear) elastic background, an elastic rod becomes unstable for $f > f_c$. We include non-linear elastic properties expected for the cytoskeleton, and show that the system is stable to supercritical loads, with a buckling amplitude that increases above threshold as $(f - f_c)^{1/2}$. In addition, both the buckling amplitude and the compressive load decay away from the point of force application in a way that depends sensitively on the longitudinal mechanical coupling of the filament to its surroundings. This suggests that the range of force transmission in the cell can be effectively controlled by microtubule-associating proteins that couple MTs to the rest of the cytoskeleton. Surprisingly, the experiments in ref. \cite{2} (see fig. 6 therein) also found evidence that, despite the existence of a threshold force for buckling, the deformation of the microtubule was attenuated, suggesting a spatially varying force.

In studying the mechanical response of intracellular MTs, we extend the classical buckling theory of rods in an elastic medium \cite{2,8,9} to take into account the mechanical coupling of both longitudinal and transverse deformations.

\textsuperscript{(a)}E-mail: mdas@few.vu.nl
of the MT to the surrounding nonlinear elastic cytoskeletal network. We use a variational theory, numerical (conjugate gradient) minimization, and asymptotic analysis of the deformation of the MT. On applying a compressive load at one end, the microtubule bends and deforms the surrounding elastic network which resists this deformation. These competing effects determine the characteristic force and length scales associated with the buckling of the MT. The nonlinear elastic properties and longitudinal coupling of the MT to its surroundings determine the attenuation of both the force and deformation of the MT along its arclength away from the point of loading. While the details of the deformation profile depend on the precise form of the nonlinearity, the force penetration depth generically decreases with increasing strength of the longitudinal coupling between the MT and the medium.

We consider a linear elastic theory for the MT and the surrounding medium and include the leading nonlinearity for the elasticity of the surrounding matrix. The MT is subjected to a compressive load \( f \) at one end denoted by \( x = 0 \). The resulting elastic deformation energy can be written in terms of its transverse and longitudinal displacements, \( u(x) \) and \( v(x) \), respectively:

\[
\mathcal{E} = -fv(0) + \int_0^\infty \left[ \frac{\kappa}{2} u''(x)^2 + \frac{\alpha_\perp}{2} u(x)^2 \right] dx + \int_0^\infty \left[ \frac{\alpha_\parallel}{2} v(x)^2 + \frac{\beta}{4} v(x)^4 \right] dx, \tag{1}
\]

where \( v(x) = \int_x^\infty \frac{1}{2} u'(y)^2 dy \) as is required for an incompressible rod. The first term represents the axial compression energy released by bending. The second term corresponds to the bending energy of the rod; \( \kappa \) is the bending rigidity. Classical Euler buckling, in which a long-wavelength bend of a rod of length \( L \) occurs at a threshold force of \( \pi^2 \kappa / L^2 \), results from just the first two terms [8,10]. The elastic constants \( \alpha_\perp, \alpha_\parallel \) correspond to the linear elastic transverse and longitudinal mechanical couplings between the MT and the surrounding actin network. The transverse coupling is directly related to the (linear) shear modulus \( G \) of the surrounding medium: \( \alpha_\perp = 4\pi G \ln(\lambda/a) \), where \( \lambda \) is the characteristic wavelength of lateral displacement and \( a \) is a small wavelength cut-off. Adding just this third term leads to buckling on a short wavelength \( \lambda = 2\pi \ell_0 \), where \( \ell_0 = (\kappa / \alpha_\perp)^{1/4} [2,8] \), and at a force threshold \( f_c = 8\pi^2 \kappa / \lambda^2 = 2(\kappa \alpha_\perp)^{1/2} \) that is significantly larger than for Euler buckling. The longitudinal coupling, in contrast, is not directly related to the elasticity of the surrounding medium, but also depends on the strength/degree of attachment between the MT and the surrounding network. Thus, it falls in the range \( 0 \leq \alpha_\parallel \leq 2\pi G / \ln(\lambda/a) \). Here, the lower limit corresponds to an uncoupled MT that slides freely along its axis, while the upper limit corresponds to the strong coupling of the MT to the medium along its length. Finally, the term proportional to \( \beta > 0 \) represents the leading non-linear transverse coupling to the medium.

In fact, both terms in the second integrand of eq. (1) are non-linear in the transverse displacement \( u \) since the first of these is effectively fourth-order in \( u \), due to the quadratic relationship between \( u \) and \( v \).

We examine the effect of the nonlinear terms in eq. (1) by numerically determining the deformation of a MT in the surrounding matrix that is subjected to symmetric compressive loads applied at both ends: \( x = 0 \) and \( x = L \). In the limit of large \( L \), the solution near \( x = 0 \) is equivalent to that of a MT loaded at just one end, and the symmetrized loading is simply for computational convenience. Using a conjugate gradient technique [11], we minimize the energy of the loaded system given by eq. (1) with an additional term \( +fv(L) \) representing the load at \( x = L \) and subject to hinged boundary conditions. We find that the buckling threshold \( f_c \) and wavelength \( \lambda = 2\pi \ell_0 \) remain approximately the same as in the linear system, but that the amplitude of the undulatory deformation \( u(x) \) just above threshold (i.e., for small \( f - f_c > 0 \)) is highly attenuated (see fig. 1). The buckling amplitude scales with the applied force \( f \) as \( (f - f_c)^{1/2} \) reminiscent of a supercritical pitchfork bifurcation [12]; it also scales with the nonlinear elastic constant \( \beta \) as a power law with exponent \(-1/2\) as shown in the inset of fig. 2. At a given force, the oscillatory deformation field decays with an apparently exponential envelope as one moves along the axis of the MT away from the loading point, with a penetration depth or decay length that scales as \( (\beta / \alpha_\parallel)^{1/2} \) (fig. 3).

We propose a variational ansatz for the form of the transverse deformation field based on the numerical solutions shown in fig. 1:

\[
u(x) = u_0 \exp \left(-x/\ell_0\right) \sin (qx). \tag{2}\]

Substituting this into our energy expression, eq. (1), we determine \( u_0, \ell_0 \) and \( q \) so as to minimize that energy. For applied loads less than a threshold given approximately by \( f_c = 2(\kappa \alpha_\perp)^{1/2} \), the minimum requires a straight MT: \( u_0 = q = 0 \). Above this threshold, the minimum is at a finite value of \( u_0 \) and \( q \approx 1/\ell_0 \). Once again, the buckling amplitude grows with applied load as \( u_0 \sim (f - f_c)^{1/2} \) and decreases with an increasing nonlinear elastic constant \( \beta \) as \( \beta^{-1/2} \) as found numerically (see fig. 2). At a given load

Fig. 1: The deformation of a stiff elastic filament embedded in a nonlinearly elastic medium decays as an oscillatory exponential (from numerics). The material parameters are \( \kappa = \beta = \alpha_\perp = 1.0 \), and \( \alpha_\parallel = 0.1 \). The deformation \( u \) has been nondimensionalized by scaling it by \( \ell_0 = (\kappa / \alpha_\perp)^{1/4} \).
above threshold, $f > f_c$, the penetration depth of the oscillatory deformation field scales as $(\beta/\alpha_\parallel)^{1/2}$ (fig. 3). The results of the variational calculation, therefore, corroborate all of the observations from the numerics.

We nondimensionalize the energy in eq. (1) by scaling $x$ and all displacements $u, v$ by $\ell_0 - e.g., \tilde{u}(\tilde{x}) = u(x/\ell_0)/\ell_0$. We also scale the energy by $f_0\ell_0$, where $f_0 \equiv (\kappa\alpha_\perp)^{1/2}$. The resulting dimensionless energy $\tilde{E} = f_0\ell_0 u_0$ can be written as

$$\tilde{E} = -\phi \tilde{e}(0) + \int_0^\infty \left[ \frac{1}{2} \tilde{u}''(\tilde{x})^2 + \frac{1}{2} \tilde{u}(\tilde{x})^2 \right] d\tilde{x} + \int_0^\infty \left[ \frac{\epsilon}{2} \tilde{v}(\tilde{x})^2 + \frac{\eta}{4} |\tilde{u}(\tilde{x})|^4 \right] d\tilde{x},$$

where $\phi = f/f_0$ is the dimensionless applied force and $\tilde{x} = x/\ell_0$. We have also introduced dimensionless material parameters $\epsilon = \alpha_\parallel/\alpha_\perp$ and $\eta = \beta\ell_0^2/\alpha_\perp$. Based on the estimates above for $\alpha_\parallel$, we expect that $\epsilon < 1/2$. The local strain is approximately given by $u' \approx \tilde{u}$. For cytoskeletal networks, significant stiffening is expected for strains less than 1 [13,14]. Thus, we expect $\eta \ll 1$. Our calculations are all for $\eta/\epsilon \gg 10, 0.5 < \eta < 100, \text{and } 0.001 < \epsilon < 0.1$.

In terms of the dimensionless variables, the ansatz in eq. (2) becomes $\tilde{u}(\tilde{x}) = \psi \exp(-\tilde{x}/\Lambda) \sin(\tilde{q} \tilde{x})$, with $\Lambda = \ell/\ell_0$ and $\tilde{q} = q\ell_0$. Using this to minimize eq. (3), we find the dependence of $\Lambda$ and $\psi$ on the applied load near threshold, $(\phi - \phi_c)/\phi_c \ll 1$, in the case of weak longitudinal coupling ($\epsilon \ll 1$), as follows. In this limit, $\Lambda \gg 1$ so we expand the energy eq. (3) in $\Lambda^{-1}$ and keep the leading terms, which are of order $\Lambda$ and $\epsilon \Lambda^3$. Dropping corrections that are suppressed by a factor of either $\epsilon$ or $1/\Lambda^2$, we determine the minimum of the truncated, asymptotic expansion of the energy with respect to $\psi, \Lambda$, and $\tilde{q}$. We find $\tilde{q} = \tilde{q}_0 = 1$, and

$$\Lambda = \sqrt{\frac{12\eta}{\epsilon}}, \quad \psi = \pm \sqrt{\frac{4}{3\eta}} (\phi - \phi_c).$$

To lowest order in $1/\Lambda$ and $\epsilon$, the critical force $f_c$ is unchanged from that of the linear theory. Specifically, $\phi_c = 2 + 4\Lambda^{-2} + \cdots$. In fig. 4 we show results of the numerics (filled symbols), and variational analysis (open symbols).
as well as from the asymptotic analysis (lines) demonstrating the excellent agreement of these approaches.

The $u^4$ term in eq. (3) represents only the initial nonlinear correction to the elasticity of the matrix. We expect a stronger nonlinear response for crosslinked cytoskeletal networks. Prior theory and experiment suggest an inverse quadratic divergence of the stress versus strain [13–15] for F-actin networks. Motivated by this, we replace the transverse elastic terms in eq. (4) by a term of the form $(\bar{u}^2/4)[(\bar{u}/\bar{u}_s) + 1]^{-1} + [(\bar{u}/\bar{u}_s) - 1]^{-1} - 2].$ This is chosen to reproduce the $u^2$ term in eq. (3) to leading order. For small $\bar{u}$, in fact, this should be equivalent to eq. (3) with $\eta = 2/\bar{u}_s^2.$ With this more physical nonlinearity, we find a clear departure from the exponential envelope for buckling oscillations observed earlier. In fig. 5 we show the amplitude of the buckled MT for $\bar{u}_s = 1.4$; there appears to be a transition from a regime of slow amplitude decay near the point of loading to an exponential decay with the same rate as for the model in eq. (3) with $\eta = 1.$ The inset compares the actual decay envelopes for the simple and higher-order nonlinearities. The latter decreases the amplitude of buckling near the point of loading, however, the envelope asymptotes to exponential decay with a decay length equal to that for the simple nonlinearity of eq. (3).

Finally, we note that we have assumed the MT to be incompressible. Finite compressibility introduces another decay length; the compressive strain on an elastic rod with compression modulus $\mu$ and longitudinal coupling to the network of $\alpha_\parallel$ decays over a length scale $\sqrt{\mu/\alpha_\parallel}$ in the unbuckled limit. Treating a MT as an elastic rod of radius $a = 10 \text{ nm}$ and Young’s modulus $1 \text{ GPa}$ [16], we estimate this decay length to be $\sim 10 \mu \text{ m}$ for the largest $\alpha_\parallel \simeq 1 \text{ kPa}.$ At smaller $\epsilon,$ where our model is applicable to intracellular MTs, the actual decay length will be larger than this, likely spanning the whole cell.

Motivated by observations of MT buckling in the cytoskeleton, we have explored force propagation and compressive buckling of stiff rods embedded in model nonlinear elastic matrices. Consistent with prior experiments [2], we find oscillatory buckling with a decaying amplitude and finite spatial extent $\ell.$ We have focussed on the limit of weak longitudinal coupling $\alpha_\parallel,$ where the penetration depth $\ell$ is large compared with the buckling wavelength $\lambda.$ We estimate these lengths using the experimental values from ref. [2]. From $\kappa \simeq 2 \times 10^{-23} \text{ Nm}^2$ and $G \simeq 1 \text{ kPa},$ we estimate that $\lambda \simeq 2 \mu \text{ m}$ and $\ell \simeq 0.3 \mu \text{ m}.$ The local strain in the network induced by a dimensionless displacement $\bar{u}$ is of order $\bar{u}.$ Thus, for a typical cytoskeletal network that becomes nonlinear for strains of order $10–20\%,$ we estimate that $\eta$ is no less than unity, and may be as large as $10.$ Thus, $\Lambda = \ell/\ell_0 \gtrsim 5$ since $\epsilon \ll 1/2.$

Even for an isolated elastic rod, there are intrinsic nonlinearities that arise from the geometry of a buckled shape [17,18]. These nonlinearities are best described in terms of the angle $\theta(s)$ that the rod makes with respect to the $x$-axis as a function of the arc length $s$ along the rod, $u' = \tan \theta.$ In terms of $\theta,$ for instance, $v(s) = \int_s^\infty (1 - \cos \theta) \, ds,$ while the curvature is $d\theta/ds.$ We have evaluated the corresponding energy for our problem of a rod in an embedded medium in the asymptotic limit of $\epsilon \to 0,$ or $\Lambda \to \infty.$ Near the critical force at which buckling sets in, where the deflection $u$ is small, a consistent expansion in $u$ is still possible, and the leading quadratic order energy terms that arise from the first three terms in eq. (3) are unaffected. The nonlinearities, however, generate additional terms of order $u^3$ that also depend on $q,$ in general. Near the critical point, however, where such an expansion is valid, the equilibrium $\bar{q} = 1$ up to order $u^2,$ and the leading-order nonlinear effects can be incorporated in our phenomenological parameter $\eta.$ Specifically, we find that the combined effects of the nonlinearities arising from the longitudinal displacement $\bar{v},$ the curvature $d\theta/ds,$ and performing the integrals with respect to the measure $ds$ are equivalent to a simple replacement $\eta \to \eta - 1/3$ near the critical point. Thus, for realistic systems, where $1 \leq \eta \leq 10,$ we expect the extrinsic nonlinearities of the surrounding medium to dominate the geometric nonlinearities of the buckled shape. Therefore, these geometric nonlinearities will change our estimates of the amplitude $\psi$ and decay length $\Lambda$ by only a small amount for realistic values of $\eta \gtrsim 1.$ Furthermore, it is interesting to note that the existence of a decay length for the buckling amplitude still depends crucially on the existence of finite longitudinal coupling $\alpha_\parallel.$

Fig. 5: The transverse displacement $u(x)$ (scaled by $\ell_0$) in the buckled state of the MT in the presence of higher-order nonlinearity (from numerics). The solid line (black) envelope corresponds to an asymptotic exponential $(1.4 \exp(-x/28.8))$ fit, with the same decay length as for the simple $u^4$ nonlinearity. The material parameters are $\epsilon = 0.01, \eta = 1, \phi = 3.5.$ Inset: The circles (blue) show $u(x)$ at odd multiples of $\lambda/4$ (corresponding to the peaks in $u(x)$) as a function of axial length $x$ on a semilog scale for the higher order nonlinearity, while the solid line (black) shows the exponential decay with the simple $u^4$ nonlinearity.
We find that extrinsic nonlinearities of the embedding medium can result in a continuous force – extension/compression relation, much as intrinsic geometric nonlinearities can do for isolated elastic rods [17,18]. But, within this model only longitudinal coupling of an elastic rod to a surrounding medium can account for a decaying buckling amplitude, such as observed in the experiments of ref. [2]. This work also shows how force transmission by MTs can be controlled by the direct coupling of the MTs to the rest of the cytoskeleton, which may be regulated by microtubule-associating proteins. This can allow an embedded MT to bear and transmit supercritical forces over a long range within the cell. Although we have only considered here the effects of the confinement of MTs by and coupling of MTs to a bulk elastic medium, it is also possible that MTs near the plasma membrane may couple to the cell substrate. Such coupling can also increase the load-bearing capacity of MTs. Force transmission such as we consider here is different qualitatively from stress propagation in a homogenous and isotropic elastic continuum in that the force is focussed primarily along a single load-bearing element. We find that this force decays with a decay length $\ell$ to an asymptotic value $f_c$. For typical parameter values of $\eta = 1$ and $\epsilon = 0.01$, the lengthscale $\ell \sim 35 \times \ell_0 \gtrsim 10 \mu m$. Finite compressibility or elastic inhomogeneities in the surrounding medium can also lead to finite-range force transmission, although here we consider the case of a single stiff filament in an elastic continuum. In this case we show that mechanical loads applied at a cell’s periphery can propagate along MTs throughout the cell body without significant diminution. This elucidates the importance of MTs in force propagation in a network of F-actin with embedded MTs and agrees very well with recent experiments [19] that show that presence of MTs can significantly enhance the elastic response of such a network. While our study focusses on the mechanical response of intracellular microtubules, our results are applicable to composite elastic media with a wide separation of scales in stiffness of its constituents such as biopolymer networks with bundles and even nanotube-polymer composites.

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