

Tunable corrugated patterns in an active nematic sheet

Anis Senoussi, Shunnichi Kashida, Raphaël Voituriez, Jean-Christophe Galas, Ananyo Maitra, André Estévez-Torres

▶ To cite this version:

Anis Senoussi, Shunnichi Kashida, Raphaël Voituriez, Jean-Christophe Galas, Ananyo Maitra, et al.. Tunable corrugated patterns in an active nematic sheet. Proceedings of the National Academy of Sciences of the United States of America, National Academy of Sciences, 2019, pp.201912223. 10.1073/pnas.1912223116. hal-02334253

HAL Id: hal-02334253 https://hal.archives-ouvertes.fr/hal-02334253

Submitted on 27 Nov 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Tunable corrugated patterns in an active nematic sheet

Anis Senoussi,[†] Shunnichi Kashida,[†] Raphael Voituriez,^{†,‡} Jean-Christophe

Galas,*^{,†} Ananyo Maitra,^{*,†} and André Estevez-Torres^{*,†}

1

†Sorbonne Université and CNRS, Laboratoire Jean Perrin, F-75005, Paris, France ‡Sorbonne Université and CNRS, Laboratoire de Physique Théorique de la Matière Condensée, F-75005, Paris, France

E-mail: jean-christophe.galas@upmc.fr; nyomaitra07@gmail.com; andre.estevez-torres@upmc.fr

2

Abstract

Active matter locally converts chemical energy into mechanical work and, for this 3 reason, it provides new mechanisms of pattern formation. In particular, active nematic 4 fluids made of protein motors and filaments are far-from-equilibrium systems that may 5 exhibit spontaneous motion, leading to actively-driven spatio-temporally chaotic states 6 in two and three dimensions and coherent flows in three dimensions (3D). Although 7 these dynamic flows reveal a characteristic length scale resulting from the interplay 8 between active forcing and passive restoring forces, the observation of static and large 9 scale spatial patterns in active nematic fluids has remained elusive. In this work, we 10 demonstrate that a 3D solution of kinesin motors and microtubule filaments sponta-11 neously forms a 2D free-standing nematic active sheet that actively buckles out-of-plane 12 into a centimeter-sized periodic corrugated sheet that is stable for several days at low 13 activity. Importantly, the nematic orientational field does not display topological de-14 fects in the corrugated state and the wavelength and stability of the corrugations are 15 controlled by the motor concentration, in agreement with a hydrodynamic theory. At 16 higher activities these patterns are transient and chaotic flows are observed at longer 17 times. Our results underline the importance of both passive and active forces in shap-18 ing active matter and demonstrate that a spontaneously-flowing active fluid can be 19 sculpted into a static material through an active mechanism. 20

Significance: To what extent can we engineer matter that shapes itself? To investigate 21 this question we study an aqueous solution containing molecular motors that walk on protein 22 filaments. When the filaments are long and attract each other, bundles of filaments are 23 parallelly oriented. We show that such a nematic solution in the presence of multimers 24 of motors has an unexpected behavior: it forms a fluid film that autonomously wrinkles. 25 The observed wrinkles have a well-defined wavelength that decreases with increasing motor 26 concentration. The wrinkles are either stable or break into a chaotic flowing state at high 27 motor concentration, providing insights into how to engineer static or dynamic materials 28 with this class of active matter. 29

Active matter is composed of subunits that convert free energy into mechanical work. 30 It comprises systems composed of objects with very different sizes, from flocks of animals¹ 31 and bacterial colonies² to gels of cytoskeletal proteins.^{3,4} Active matter has attracted much 32 attention, both theoretically and experimentally, because it displays phase transitions and 33 states that greatly differ from those observed at equilibrium, such as motile ordered states and 34 spontaneous coherent or incoherent flow.^{5–9} Among the active systems that can be studied 35 in the laboratory, those composed of the protein filaments and motors that constitute the 36 cytoskeleton of the eukaryotic cell are of special interest for three reasons: i) their biological 37 importance,^{10,11} ii) the possibility to make purified systems that can be easily controlled and 38 studied.^{3,4,12} and iii) their potential to make self-organising materials.¹³ 39

Depending on the conditions, cytoskeletal active systems display a wide array of dynamic 40 behaviors. Isotropic systems contract^{12,14–16} and buckle¹⁷ in three dimensions (3D). Polar 41 ones generate density waves¹⁸ and large scale vortices¹⁹ in 2D, and asters and vortices³ in 42 3D. Nematic systems display spatio-temporally chaotic flows both in 2D^{4,20,21} and in 3D,^{4,22} 43 and also coherent flow²² in 3D. This diversity of behaviors is qualitatively understood by a 44 hydrodynamic theory.^{5–9} However, we currently do not fully understand why one behavior 45 is observed in a given experimental system and not in another and which experimental 46 parameter has to be modified to switch from one state to another. This is due, on the one 47 hand, to the difficulty of measuring the phenomenological parameters of the hydrodynamic 48 theory and, on the other hand, to the use of two experimental systems, actin/myosin and 49 microtubule/kinesin, with very different microscopic properties. In this regard, the recent 50 demonstration that global contractions^{12,15} and chaotic flows in $2D^{4,21}$ were present in both 51 systems, and the understanding of the nematic to polar transition in microtubule/kinesin 52 systems²³ has clarified the design of these dynamic behaviors. Yet, the two aforementioned 53 difficulties remain, hindering the development of controllable active materials. 54

In this work, we report the observation of a novel static patterned state in an active nematic fluid, we provide a semi-quantitative interpretation to why this state is observed and

we show which experimental parameters need to be tuned to reach either this static state or 57 a previously-reported flow state. More precisely, we demonstrate that a microtubule/kinesin 58 nematic fluid that is known to flow in $3D^{4,22}$ can be rationally engineered to form a thin 59 static corrugated sheet in three dimensions, a behavior that has only recently been observed 60 in isotropic and cross-linked $\operatorname{actin}/\operatorname{myosin}$ gels¹⁷ that cannot flow. Essentially, the fluid 61 contracts anisotropically along its two shortest dimensions to form a thin sheet of gel that 62 freely floats in the aqueous solution, mainly due to passive depletion forces. Simultaneously, 63 the extensile active stress generated by the motors buckles the sheet along the direction 64 perpendicular to its plane, forming a corrugated sheet of filaments with a well-controlled 65 wavelength of the order of 100 μ m over an area of 10 mm². We demonstrate that this out-of-66 plane buckling differs both from classical Euler buckling in passive gels^{24,25} and from the flow-67 generating in-plane buckling that is common in 2D active nematic gels.^{4,22,26,27} Importantly, 68 nematic topological defects are not observed and we provide a theoretical prediction of the 69 dependence of the wavelength with the motor concentration and with the thickness of the 70 fluid that is in agreement with the experiments. Finally, we show that the transition between 71 static corrugations and chaotic flow can be experimentally controlled by two parameters, the 72 motor concentration and the attractive interactions between microtubule filaments. 73

74 Results and Discussion

⁷⁵ A 3D active nematic fluid forms a static corrugated sheet

The active fluid is constituted of a dense suspension of non-growing microtubules bundled together by a depletion agent and by clusters of kinesin-1 motors (Figure 1a). We use the word fluid, instead of gel, to underline the fact that the system does not present irreversible crosslinks, as actin gels do.¹⁷ It is supplemented with ATP and an ATP-regeneration system that drives the system out of equilibrium by keeping the motor active for at least 4 h. Additionally, the microtubule bundles are fluorescent because they bear a small fraction of

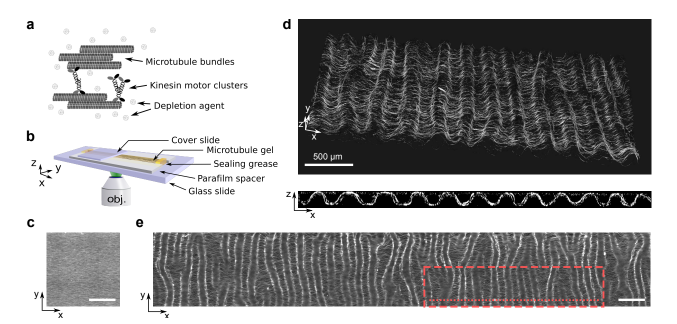


Figure 1: At low motor concentration a 3D active nematic fluid creates a thin corrugated sheet of well defined wavelength. **a** Scheme of the components of the active fluid formed by non-growing microtubules bundled together by a depletion agent and clusters of kinesin motors. **b** Scheme of the channel where the fluid (in yellow) is observed. **c** Epifluorescence image of the fluid at initial time. **d** Confocal images in 3D (top) and cross-section in the xz plane (bottom) of the fluid after 300 min. **e** Epi-fluorescence image of the same sample after one day and over a $9.5 \times 1.4 \text{ mm}^2$ area, the red dashed rectangle and the red dotted line respectively indicate the region where the top and bottom images in panel d were recorded. Scale bars are 500 μ m and motor concentration 0.5 nM.

fluorescent tubulin, allowing their observation by fluorescence microscopy. This system is 82 similar to previously published active nematic fluids⁴ but it differs in several important ways: 83 the microtubules are longer (8 \pm 6 μ m instead of 1 μ m, Figure S1), the kinesin used here,²⁸ 84 K430, is different from the standard K401 (it comes from a different organism and forms 85 non-specific clusters), and its typical concentration is two orders of magnitude lower (see SI 86 Section 1). The active fluid is prepared inside a long and shallow channel of rectangular 87 cross-section, with length L = 22 mm, width W = 1.5 mm and height H = 0.13 mm (see SI 88 Methods). Initially, the density of microtubule bundles is homogeneous in 3D but they are 89 aligned along the long axis of the channel, parallel to x (Figure 1b-c). This nematic order 90 arises spontaneously during the filling process of the channel by capillarity, the angle of the 91 director of the nematic with the x axis being $2 \pm 16^{\circ}$ (Figure S2). 92

In the presence of 0.5 nM of motors, confocal images recorded after 300 min show that 93 the active fluid has contracted along z and buckled in the xz plane to form a corrugated sheet 94 whose hills and valleys reach the top and bottom walls of the channel and whose grooves are 95 strikingly parallel to the y axis (Figure 1d). The thickness of the sheet is $\ell_z = 35 \pm 5 \ \mu \text{m}$ and 96 the wavelength of the corrugations is $\lambda = 285 \pm 15 \,\mu \text{m}$. This periodic pattern extends along an 97 area of at least $9.5 \times 1.4 \text{ mm}^2$, with dislocations corresponding to the junction of two valleys 98 or hills. Notably, these dislocations in the periodic undulatory pattern do not correspond to 99 defects in the nematic field. The pattern can also be visualized in epifluorescence, where it 100 appears in the form of focused and defocused bands (Figure 1e). 101

¹⁰² During the formation of the corrugations the fluid buckles along z¹⁰³ and contracts along z and y

To elucidate the mechanism of pattern formation we recorded confocal (Movie S1) and 104 epifluorescence time-lapse images of a buckling fluid at 0.5 nM motors (Figure 2). Two 105 processes are observed: buckling along the z direction and contraction along z and y. These 106 processes are quantified by the angle ϕ between the microtubule bundles and the x axis in 107 the xz plane, and by $\Delta \ell_z$ and $\Delta \ell_y$, the contracted lengths of the fluid along the z and y 108 axes, respectively. Buckling initially proceeds at a rate $\omega_{\phi} = 0.3 \text{ min}^{-1}$ but later slows down 109 until reaching a maximal buckling angle $\phi^{max} = 32.2 \pm 0.5^{\circ}$ (Figure 2c) and an amplitude 110 $h^{max} = 22 \pm 3 \ \mu \text{m}$ after 100 min. Contraction along z and y is significantly slower with onset 111 rates $\omega_z = 6.4 \times 10^{-2} \text{ min}^{-1}$ and $\omega_y = 1.5 \times 10^{-2} \text{ min}^{-1}$, respectively, to reach maximum 112 amplitudes $\Delta \ell_z^{max} = 40 \ \mu m$ and $\Delta \ell_y^{max} = 210 \ \mu m$ (Figure 2d). Note that the relative 113 contraction amplitudes $\Delta \ell_z^{max}/H = 0.40$ and $\Delta \ell_y^{max}/W = 0.14$ are significantly different, 114 indicating that the final contracted state does not correspond to a nematic liquid droplet at 115 equilibrium.²⁹ 116

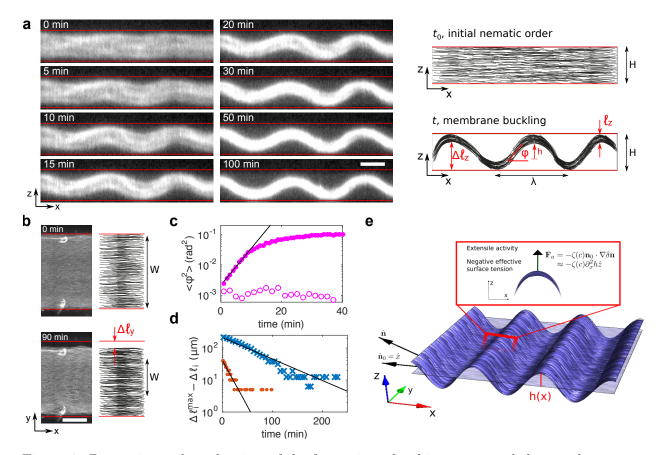


Figure 2: Dynamics and mechanism of the formation of a thin corrugated sheet at low motor concentration. **a** Time-lapse confocal fluorescence images of the active fluid in the xz plane (left) and sketch of the observations indicating the measured quantities $\Delta \ell_z$, ϕ , h, and ℓ_z . Scale bar is 100 μ m. **b** Epifluorescence images of the fluid at t = 0 and 90 min (left) and sketch indicating the measured quantity $\Delta \ell_y$. Scale bar is 500 μ m. Red lines in panels a and b indicate channel walls. **c** Average of ϕ^2 along the x direction vs. time in the presence (filled disks) and in the absence of motors (empty circles). **d** Offset to the maximum contracted length along the z (red disks) and y (blue crosses) directions. Black lines in panels c and d are exponential fits. **e** Sketch of the mechanism for the active buckling of a thin membrane through the negative tension F_a proportional to the active stress $\zeta(c)$ and the Laplacian of the height h(x) of the sheet above its fiducial plane. All data correspond to 0.5 nM motors except empty circles in panel c.

¹¹⁷ Buckling is active and contraction is passive

Passive gels of various compositions have been reported to form corrugations through Euler buckling when they are submitted either to an external contractile stress or to an extensile stress at constant length.^{24,25} To assess if this could explain our observations, we performed experiments that demonstrated, firstly, that buckling is principally an active process and, secondly, that an Euler mechanism is not compatible with the data.

In the absence of motors buckling is undetectable in confocal images with the same field of view in x as above (660 μ m) (Figure 2c and Table S1), although it is weak but detectable in images acquired over a wider field of view (Figure S3). In contrast, contraction is similar both in passive and active fluids (Figure S3). The passive origin of contraction is further supported by the fact that its amplitude is strongly dependent on the concentration of depletion agent (Figure S4).

In passive fluids, depletion forces induce the condensation of microtubules into a dense 129 nematic fluid phase, which, in the absence of confinement, would relax to a highly anisotropic 130 tactoid droplet.³⁰ In the geometry of our experiments, this results in the formation of a quasi 131 2D sheet that elongates along the nematic axis x, thereby leading to Euler buckling in the 132 presence of boundaries. Indeed, a membrane with an excess area, which in this case arises 133 from the excess length $\Delta \ell_x = \ell_x - L$, where ℓ_x is the length of the membrane along x, will 134 have a buckled state beyond a critical $\Delta \ell_x^c$ that depends on H. Note that this mechanism 135 could in principle also explain active buckling if $\Delta \ell_x$ depends on the activity. However, this 136 sort of passive mechanism, reminiscent of the classic Euler buckling, is only possible if the 137 membrane had a fixed projected area, i.e. if it were confined in the x direction. On the 138 contrary, if active buckling were generated by local extensile forces exerted by the motors, 139 it would be independent of whether the fluid is constrained in length or not. 140

To test these two hypotheses, we performed experiments where one end of the fluid was 141 in contact with an aqueous solution and thus free to change length. In this configuration 142 (Figure 3) the active fluid buckled everywhere except on the tip close to the free boundary, 143 while the passive one did not buckle at all. We attribute the lack of buckling on the tip of 144 the active fluid to a gradient of microtubule concentration across the free boundary arising 145 during the preparation of the fluid (see SI Methods). These results suggest that, while the 146 excess area mechanism explains passive buckling, it cannot fully account for the buckling 147 of active films. We thus conclude, firstly, that passive and active buckling happen through 148

different mechanisms and, secondly, that in active fluids buckling is principally an active mechanism while contractions in y and z are mainly passive.

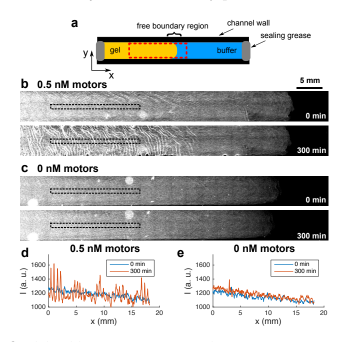


Figure 3: An active fluid buckles, in contrast with a passive one, in fluids with one free boundary. **a** Scheme of the experiment. The red dashed rectangle indicates the area where the images were recorded. Epifluorescence images of an active (**b**) and a passive (**c**) fluid at initial and final time. The black, dashed rectangles show the zones where the intensity profiles in **d** and **e** were extracted.

¹⁵¹ A hydrodynamic theory that predicts the wavenumber of the cor-¹⁵² rugations

The behavior of active fluids, including those composed of microtubules and kinesins, ^{4,22,31,32} has been successfully described with the hydrodynamic theory of liquid crystals supplemented with a stress term resulting from activity. We now demonstrate that this framework applied to a thin film that can buckle in the third dimension can provide an explanation and theoretical estimates of the wavenumber $q^* = 2\pi/\lambda$ and of the formation rate ω^* of the corrugated pattern (see Figure 2e and SI Section 2).

To do so, we consider the periodic undulation of the thin sheet made of microtubules and motors in the xz plane, supposing that passive forces have already collapsed the 3D fluid

into a thin 2D sheet. The nematic active fluid sheet has bending modulus K and its director 161 $\hat{\mathbf{n}}$ is on average parallel to the x axis: $\hat{\mathbf{n}}_0 = \hat{x}$. The fluctuation of the membrane about a 162 fiducial plane parallel to the xy plane (here, taken to be the mid-plane of the channel) is 163 denoted by h(x, y). The deflection of the director in the xz plane, $\delta \mathbf{n}_z$, leads to a buckling 164 of the membrane in the z direction: $\delta \mathbf{n}_z \approx \partial_x h \hat{z}$. The passive elasticity of the nematic fluid 165 $\propto (\nabla \mathbf{n})^2$ then yields a bending energy $\propto (K/2)(\partial_x^2 h)^2$ for the buckling of the thin sheet in 166 the z direction. The standard active force⁵⁻⁹ is $-\zeta(c)\nabla \cdot (\mathbf{nn})$, where $\zeta(c) > 0$ is the strength 167 of the extensile activity that is a function of motor concentration c. This leads to a force 168 $\propto -\zeta(c)\partial_x^2 h\hat{z}$ that tends to destabilise the flat membrane and that is similar to an effective 169 negative surface tension.³³ The interplay between the negative surface tension, arising from 170 activity, and the stabilizing bending modulus, due to nematic elasticity, leads to the selection 171 of a pattern with wavenumber 172

$$q^* \sim \sqrt{\zeta(c)/K}.\tag{1}$$

The pattern arises with a rate ω whose exact expression is provided in SI Section 2. Note that in the absence of confinement, we expect the pattern to be unstable.

The theory thus shows that an out-of-plane buckling instability compatible with our 175 observations results from the interplay of active forcing $\zeta(c)$ and passive elastic restoring 176 forces, K; the same ingredients that in previous microtubule/kinesin active fluid provided 177 dramatically different patterns.^{4,31,32,34,35} Here, the out-of-plane buckling of the active sheet 178 precedes any planar pattern formation, in contrast to those experiments. In addition, the 179 instability described here does not result in coherent or incoherent flow, of either the active 180 or the embedding fluid, in contrast with theories describing 2D or 3D active fluids that do 181 not form sheets. 26,27,36 182

This qualitative interpretation has two advantages. Firstly, it is parsimonious because a single feature, activity, explains the 3D out-of-plane buckling observed here and the 2D inplane buckling^{4,32} and 3D chaotic flows²² observed previously in a similar system. Secondly,

it predicts that decreasing depletion forces precludes the formation of the thin sheet and 186 thus the emergence of out-of-plane buckling in favor of 3D chaotic flows. In the rest of 187 the paper we analyze these two questions in more detail. However, although hydrodynamic 188 theories, such as the one just described, provide an informative qualitative description of 189 the physics of active fluids, they feature phenomenological parameters, such as $\zeta(c)$ and K, 190 that are difficult to measure experimentally. To our knowledge, the only quantitative test 191 of such theories in the kinesin-microtubule system was recently performed by Martinez-Prat 192 and colleagues,³² where they obtained $\zeta(c) \sim c^2$. Using their scaling, our semi-empirical 193 prediction reads 194

$$q^* \sim c/\sqrt{K}.\tag{2}$$

Increasing the motor concentration linearly increases the wavenum ber of the corrugations and destabilizes the patterns

To test the prediction $q^* \sim c$ we investigated the behavior of the fluid over a range of motor 197 concentrations c spanning more than two orders of magnitude (Figure 4 and Movie S2). 198 Below 0.5 nM motors, the fluid behaves as described in Figure 2: buckling in the xz plane 199 and contractions in the z and y directions. As c increases, between 1 and 2.5 nM motors, 200 buckling in the xz plane is initially observed and followed by buckling in the xy plane that 201 distorts the corrugated pattern without breaking it. Finally, between 5 and 50 nM motors, 202 buckling in the xz plane is still observed at early times but the pattern breaks into a 3D 203 active chaotic state similar to the one already reported in this active fluid 4,22 (Movies S3 204 and S4). However, the velocity of this flow state is significantly lower in our case, possibly 205 because the solution is more viscous. 206

The transition to the chaotic state happens qualitatively through two processes: the accumulated tension on the hills and valleys of the corrugations breaks the microtubule bundles and the frozen fluid locally flows (Movie S2, 5 nM channel) or the dislocations

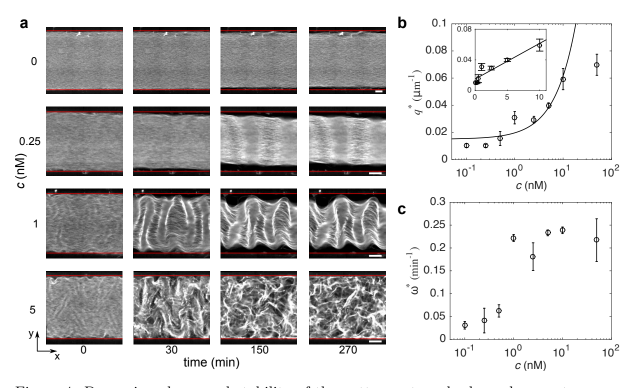


Figure 4: Dynamics, shape and stability of the patterns strongly depend on motor concentration. **a** Time-lapse epifluorescence images of fluids with different motor concentrations c. Red lines indicate channel walls. Scale bars are 200 μ m. **b** Wavenumber q^* of the corrugations vs. motor concentration. The inset is the lin-lin representation of the main plot and the line corresponds to a linear fit to the data in the range c = 0.1 - 10 nM with regression coefficient $r^2 = 0.9$. **c** Growth rate vs. motor concentration. Error bars indicate the standard deviation of a triplicate experiment where a single motor/filament mix was distributed into three different channels.

in the corrugations become motile leading to a shearing of the pattern and its consequent 210 destruction (Movie S2, 10 nM channel). Note that, in our experiments, the chaotic state 211 was never observed before the buckled state. However, if the characteristic time of active 212 transport is much shorter than the time of passive contraction, one would observe only the 213 spontaneous flow instability and would not observe the buckling instability (which happens 214 in ref. 4). Nevertheless, first observing spontaneous flow instability and then the buckling 215 instability is unlikely because the first one would destroy the nematic order that allows 216 passive buckling. 217

Importantly, the measured wavenumber of the corrugations is in agreement with the predicted linear scaling (Figure 4b), in particular in the range 0.5 - 10 nM. A linear fit $q^* = a_1 + a_2 c$ of the data yields $a_1 = 5 \times 10^{-3} \ \mu \text{m}^{-1}$ and $a_2 = 1.4 \times 10^{-3} \ \mu \text{m}^{-1} \text{nM}^{-1}$, where the constant term a_1 results from the weak contribution of Euler buckling in the absence of motors. Indeed, activity controls the wavenumber only if the active wavenumber is larger that the one selected by passive Euler buckling, a crossover that in our experiments happens at c = 0.5 nM.

The growth rate of the patterns, ω^* , increases slightly with c in the range 0.1 - 0.5 nM, then drastically between 0.5 and 1 nM, and saturates at higher c (Figure 4c and Figure S5), resulting in ω^* also increasing and then saturating with q^* . For the hydrodynamicsdominated approximation, the theory predicts $\omega^* \sim q^3$ for $q^*H \gg 1$ and $\omega^* \sim q^6$ for $q^*H \ll 1$, while our experiments correspond to $q^*H = 1.3 - 4$. In the range c = 0.1 - 0.5 nM, the data are compatible with the scaling $\omega^* \sim q^3$, although their precision does not allow to rule out other scaling laws (Figure S6).

Comparing the results of our out-of-plane instability with recent measurements of the in-232 plane buckling instability of a related system, 32 we find similar wavenumbers $(1-7 \times 10^{-2} \ \mu m)$ 233 in our case vs. $0.5 - 3 \times 10^{-2} \ \mu m$) but significantly slower dynamics $(1 - 4 \times 10^{-2} \ min^{-1})$ 234 vs. $6-240 \text{ min}^{-1}$, respectively). In addition, topological defects seem to play no role in the 235 emergence of our patterns, in contrast with what happens in 2D active nematic systems.^{21,34} 236 We do observe dislocations in the corrugations that rarely move along the y axis, although 237 they do so too slowly to play a significant role. In contrast, defects in the nematic field would 238 create non-periodic buckled shapes in the z direction 37,38 and we never observe this in our 239 conditions. 240

²⁴¹ The thickness of the nematic fluid influences the corrugations

To test the prediction $q^* \sim 1/\sqrt{K}$ we varied the thickness and the aspect ratio of the confinement of the active fluid at low motor concentration, with the hypotheses $K \sim \ell_z$ and $\ell_z \sim H$. Firstly, we measured ℓ_z and q^* for H in the range 70 – 540 μ m and confirmed that the data are in agreement with $\ell_z \sim H$ and with $q^* \sim 1/\sqrt{\ell_z}$, with the exception of this last scaling for the thinnest fluid (Figure 5). Secondly, reducing the aspect ratio of the channel section resulted in some portions of the fluid buckling in xz and others in the xy plane at W/H = 4.6 (Figure S8) and no preferential direction of buckling at W/H = 1 (Figure S9). In addition, in all the cases where both xy and xz buckling was observed, the wavenumbers in the two planes were in qualitative agreement with the aforementioned argument that essentially yields $q^* \sim 1/\sqrt{H} > q_{xy}^* \sim 1/\sqrt{W}$ when H < W and $q^* = q_{xy}^*$ when H = W.

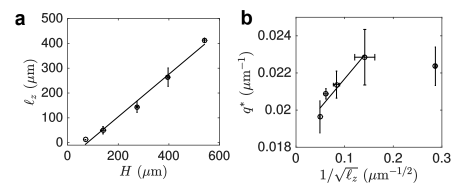


Figure 5: Increasing the thickness of the fluid reduces the wavenumber. **a** Final thickness ℓ_z vs. channel height *H*. **b** Wavenumber vs. $1/\ell_z^{1/2}$. Error bars indicate the standard deviation of a triplicate experiment where a single motor/filament mix was distributed into three different channels. Solid lines correspond to linear fits. 1 nM motors.

Another way to influence K is to change the microtubule concentration μ . Increasing μ in the range 0.5 - 2 mg/mL decreased q^* in agreement with the expectation that K should increase with μ . In contrast, at $\mu = 0.25$ mg/mL, global contraction, instead of corrugations, was observed, possibly because the initial nematic order was reduced (Figure S10).

²⁵⁶ Strong attractive interactions between the microtubules are crucial

²⁵⁷ to form a corrugated sheet

To the best of our knowledge, neither stable nor unstable out-of-plane buckling has been reported in nematic active fluids. We performed control experiments to determine which of the factors that differentiate our experiments from previously published 3D microtubule/kinesin nematics^{4,22,39} was responsible for the observed phenomenology: the type of motor or the length of the microtubules. We obtained both stable and unstable xz buckling with the kinesin K401 used in previous reports⁴ (Figure S11). This means that, although the motor K430 is not designed to form specific multimers, in contrast with K401, it forms non-specific ones. Our efforts to eliminate these non-specific multimers by size exclusion chromatography did not change the observed patterns (Figure S12), suggesting that these clusters either form rapidly or do so in the working buffer.

In contrast with the nature of the motor, the length of the microtubules had a strong 268 impact on the observed structures. When, instead of 8 μ m-long taxol-stabilized microtubules, 269 1.5 μ m-long GMPCPP-stabilized ones were used, no contraction of the fluid was observed 270 along z or y, with or without motors, precluding the formation of a thin sheet that could 271 buckle out of plane (Figure S13). In this case, chaotic flow was observed at high activity, in 272 agreement with previous reports.⁴ These observations are consistent with the expected linear 273 dependence of the depletion free energy on filament length⁴⁰ which, in our geometry, makes 274 long microtubules condense into a thin sheet. To further test this hypothesis we reduced the 275 attractive force between negatively charged 8 μ m-long microtubules by either lowering the 276 concentration of the depletion agent or the ionic strength of the buffer. In agreement with 277 this interpretation, neither fluid contraction, nor buckling in the xz plane, were apparent in 278 these conditions, although the fluid remained active (Figures S14 and S15). 279

²⁸⁰ Comparison with other out-of-equilibrium membrane buckling

The active buckling instability described here needs three ingredients: a thin film, nematic 281 order and extensile activity. We thus expect any system displaying these properties, such 282 as thin films that may be composed of living liquid crystals⁴¹ —which are suspensions of 283 living bacteria in an inert nematic solution— to buckle in a similar manner. Monolayers of 284 polarized living cells could potentially buckle in the same way, although so far only buckling 285 due to growth has been reported.⁴² Interestingly, a different type of active buckling has 286 recently been observed in a contractile isotropic film made of a crosslinked actin gel and 287 myosin.¹⁷ Despite their differences, for this contractile film to buckle, filaments also need to 288

form bundles, motors have to make aggregates and the observed thickness and wavelength were of the same order of magnitude as those reported here.

Finally, we have described our system as an active fluid and not a gel to stress that 291 there are no permanent crosslinks between the microtubule filaments. This is supported by 292 the low concentration of motors relative to tubulin (1:1000 w/w at 1 nM motors) and by 293 the flows observed by us and others $^{4,20-22,32}$ at high motor concentration. We believe that 294 the apparent contradiction of our observations with rheological measurements that reported 295 the presence of an effective cross-linking in pure microtubule solutions⁴³ could be explained, 296 either by the presence of microtubule bundles or by the longer timescale probed here (10^3 s) 297 compared with 10^2 in ref. 43). Ultimately, precise rheological measurements of this type of 298 active fluids will provide a definite answer. 290

300 Conclusion

In summary, we demonstrate that in vitro active fluids can be designed to form static or 301 transient suspended sheets with periodic corrugated patterns of tunable wavelength. The 302 mechanism of pattern formation that we propose combines passive and active processes that 303 can be controlled physicochemically. Passive depletion forces, which depend on depletion 304 agent concentration, filament length and ionic strength, induce the spontaneous condensation 305 of a 3D nematic fluid into a thin 2D nematic sheet, and active stresses buckle the fluid sheet 306 out of plane to form corrugations with well-defined wavelength that can be controlled by 307 activity. 308

In addition, we use an active gel theory to demonstrate that the observed patterns result from an out of plane buckling instability induced by active extensile stresses along the nematic axis of the fluid sheet, in contrast with in-plane buckling patterns that have been observed in pre-stressed nematic fluids of either non-growing F-actin⁴⁴ and growing microtubules⁴⁵ in the absence of motors. Our theory is appealing since it relies on the same essen-

tial physics that leads to 2D patterning and 2D and 3D spatio-temporal chaos. However, the 314 buckling instability that we report does not involve filament flows and therefore fundamen-315 tally differs from both contractile instabilities in anisotropic active fluids and spontaneous 316 flow transitions in nematic active fluids that have been described theoretically, ^{6,8,26,27,36,46} 317 and shown to be characterised by hydrodynamic flows and in-plane buckling of the director 318 field in the case of 2D systems. Such spontaneous flows have been observed in various active 319 matter systems, ^{4,31,32,34,35,41,47–50} which in practice yield either chaotic or large scale coherent 320 flows, but so far no static spatial patterns. In contrast, our results show that active matter 321 can be shaped into long-lived static 3D patterns that can be tuned by activity, which may 322 open the way to the design of 3D biomimetic materials capable of morphogenesis.^{51,52} 323

³²⁴ Material and Methods

325 Kinesin and microtubule preparation

The K430 truncated kinesin-1 from *Rattus norvegicus*, containing a C-terminal SNAP tag, 326 was the homodimer version of the kinesin-1 described in ref. 28. K430 was expressed in E. 327 *coli*, purified using a Nickel affinity column thanks to a His-tag, dialyzed and flash frozen. 328 K401 was purified as described.⁵³ Tubulin and TRITC-labeled tubulin (Cytoskeleton) were 329 dissolved at 10 mg/mL in 1X PEM buffer (80 mM PIPES pH 6.8, 1 mM EGTA, 1 mM 330 MgSO₄) supplemented with 1 mM GTP, flash-frozen and stored at -80 °C. They were 331 polymerized in 1X PEM, 1 mM GTP, 10 % (w/v) glycerol and 5 mg/mL tubulin at (including 332 2.5 % fluorescent tubulin). Taxol-stabilized microtubules were incubated at 37 °C for 15 min 333 followed by the addition of 20 μ M paclitaxel, and stored at room temperature for few days. 334 GMPCPP-stabilized microtubules were polymerized in the presence of 0.5 mg/mL GMPCPP 335 (Jena Bioscience) from tubulin at 37 °C for 30 min and left at room temperature for 5 hours. 336 They were used within the same day. These procedures are described in more detail in SI 337 Section 1. 338

339 Active mix

³⁴⁰ The active mix consisted in 1X PEM buffer, 10 mM K-acetate, 10 mM KCl, 5 mM MgCl₂, 2

 $_{341}$ %(w/v) Pluronic F-127, 5 μ g/mL creatine kinase, 20 mM creatine phosphate, 20 μ M taxol,

 $_{342}$ $\,$ 2 mM ATP, 1 mg/mL BSA, 1 mM trolox, 20 mM D-glucose, 3 mM DTT, 150 $\mu g/mL$ glucose

 $_{343}$ oxidase, 25 μ g/mL catalase and 0.5 mg/mL taxol-stabilized microtubules.

344 Channel assembly

³⁴⁵ Channels were assembled using a microscope glass slide (26 x 75 x 1 mm) and a coverslip ³⁴⁶ (22 x 50 x 0.17 mm) separated by strips of Parafilm cut with a Graphtec Cutting Plotter ³⁴⁷ CE6000-40. Both microscope glass slides and coverlips were passivated using an acrylamide ³⁴⁸ brush.⁵⁴ The active mix was filled in the flow cell (22 x 1.5 x 0.130 mm) by capillarity and ³⁴⁹ sealed with vacuum grease.

350 Imaging

Epifluorescence images were obtained with a Zeiss Observer 7 automated microscope equipped with a Hamamatsu C9100-02 camera, a 10X objective, a motorized stage and controlled with MicroManager 1.4. Images were recorded automatically every 3 min using an excitation at 550 nm with a CoolLED pE2. Confocal images were obtained with a Leica TCS SP5 II confocal microscope with a 25x water-immersion objective or a X-Light V2 Spinning Disk Confocal system mounted on an upright Nikon Eclipse 80i microscope with a 10x objective. Images were recorded automatically every 1 to 10 min.

358 Image analysis

Fluorescent images were binarized to obtain $\Delta \ell_z$ and $\Delta \ell_y$. To measure ϕ the binarized xz confocal cross-sections were averaged over x, smoothed along x by applying a moving average filter with a 30-pixel window, that was then differentiated. ϕ was the arctangent of 362 this derivative.

³⁶³ Note added in proof

³⁶⁴ During the revision process a similar observation was reported in the arXiv.⁵⁵

365 Acknowledgements

K. Furuta for providing the expression plasmids coding for the K430 kinesin, Z. Gueroui 366 for a kind gift of the K401 plasmid, F. Lam from the microscopy platform at IBPS and 367 L.L. Pontani for providing access to a spinning disk microscope, T. Surrey for insightful 368 discussions and C. del Junco and Y. Vyborna for comments on the manuscript. This work 369 has been funded by the European Research Council (ERC) under the European's Union 370 Horizon 2020 programme (grant No 770940, A.E.-T.) and by the Ville de Paris Emergences 371 programme (Morphoart, A.E.-T.). The data that support the findings of this study are 372 available from the corresponding author upon reasonable request. 373

374 References

- (1) Cavagna, A.; Cimarelli, A.; Giardina, I.; Parisi, G.; Santagati, R.; Stefanini, F.;
 Viale, M. Scale-free correlations in starling flocks. *Proceedings of the National Academy* of Sciences 2010, 107, 11865–11870.
- (2) Dombrowski, C.; Cisneros, L.; Chatkaew, S.; Goldstein, R. E.; Kessler, J. O. SelfConcentration and Large-Scale Coherence in Bacterial Dynamics. *Phys. Rev. Lett.*2004, 93, 098103.
- (3) Nédélec, F. J.; Surrey, T.; Maggs, A. C.; Leibler, S. Self-organization of microtubules
 and motors. *Nature* 1997, *389*, 305–308.

- (4) Sanchez, T.; Chen, D. T.; DeCamp, S. J.; Heymann, M.; Dogic, Z. Spontaneous motion
 in hierarchically assembled active matter. *Nature* 2012, 491, 431–4.
- (5) Toner, J.; Tu, Y.; Ramaswamy, S. Hydrodynamics and phases of flocks. Annals of Physics 2005, 318, 170 244.
- ³⁸⁷ (6) Jülicher, F.; Kruse, K.; Prost, J.; Joanny, J.-F. Active behavior of the Cytoskeleton.
 ³⁸⁸ Physics Reports 2007, 449, 3 28.
- (7) Ramaswamy, S. The Mechanics and Statistics of Active Matter. Annual Review of
 Condensed Matter Physics 2010, 1, 323–345.
- (8) Marchetti, M. C.; Joanny, J. F.; Ramaswamy, S.; Liverpool, T. B.; Prost, J.; Rao, M.;
 Simha, R. A. Hydrodynamics of soft active matter. *Reviews of Modern Physics* 2013,
 85, 1143–1189.
- ³⁹⁴ (9) Prost, J.; Julicher, F.; Joanny, J. F. Active gel physics. *Nat Phys* **2015**, *11*, 111–117.
- (10) Dogterom, M.; Surrey, T. Microtubule organization in vitro. Current Opinion in Cell
 Biology 2013, 25, 23–29.
- (11) Blanchoin, L.; Boujemaa-Paterski, R.; Sykes, C.; Plastino, J. Actin Dynamics, Archi tecture, and Mechanics in Cell Motility. *Physiological Reviews* 2014, 94, 235–263.
- (12) Bendix, P. M.; Koenderink, G. H.; Cuvelier, D.; Dogic, Z.; Koeleman, B. N.;
 Brieher, W. M.; Field, C. M.; Mahadevan, L.; Weitz, D. A. A quantitative analysis
 of contractility in active cytoskeletal protein networks. *Biophysical journal* 2008, *94*,
 3126–3136.
- (13) Needleman, D.; Dogic, Z. Active matter at the interface between materials science and
 cell biology. *Nature reviews Materials* 2017, *2*, 17048–17048.

- (14) Alvarado, J.; Sheinman, M.; Sharma, A.; MacKintosh, F. C.; Koenderink, G. H. Molecular motors robustly drive active gels to a critically connected state. *Nature Physics* **2013**, *9*, 591.
- (15) Foster, P. J.; Fürthauer, S.; Shelley, M. J.; Needleman, D. J. Active contraction of
 microtubule networks. *eLife* 2015, 4, e10837.
- (16) Torisawa, T.; Taniguchi, D.; Ishihara, S.; Oiwa, K. Spontaneous Formation of a Globally
 Connected Contractile Network in a Microtubule-Motor System. *Biophysical journal* **2016**, *111*, 373–385.
- (17) Ideses, Y.; Erukhimovitch, V.; Brand, R.; Jourdain, D.; Hernandez, J. S.; Gabinet, U. R.; Safran, S. A.; Kruse, K.; Bernheim-Groswasser, A. Spontaneous buckling of
 contractile poroelastic actomyosin sheets. *Nature Communications* 2018, *9*, 2461.
- (18) Schaller, V.; Weber, C.; Semmrich, C.; Frey, E.; Bausch, A. R. Polar patterns of driven
 filaments. *Nature* 2010, 467, 73.
- (19) Sumino, Y.; Nagai, K. H.; Shitaka, Y.; Tanaka, D.; Yoshikawa, K.; Chaté, H.; Oiwa, K.
 Large-scale vortex lattice emerging from collectively moving microtubules. *Nature* **2012**, *483*, 448.
- (20) Doostmohammadi, A.; Ignés-Mullol, J.; Yeomans, J. M.; Sagués, F. Active nematics.
 Nature Communications 2018, 9, 3246.
- (21) Kumar, N.; Zhang, R.; de Pablo, J. J.; Gardel, M. L. Tunable structure and dynamics
 of active liquid crystals. *Science Advances* 2018, *4*.
- (22) Wu, K.-T.; Hishamunda, J. B.; Chen, D. T. N.; DeCamp, S. J.; Chang, Y.-W.;
 Fernández-Nieves, A.; Fraden, S.; Dogic, Z. Transition from turbulent to coherent flows
 in confined three-dimensional active fluids. *Science* 2017, 355.

- (23) Roostalu, J.; Rickman, J.; Thomas, C.; Nédélec, F.; Surrey, T. Determinants of Polar
 versus Nematic Organization in Networks of Dynamic Microtubules and Mitotic Motors. *Cell* 2018, 175, 796–808.e14.
- ⁴³¹ (24) Matsuo, E. S.; Tanaka, T. Patterns in shrinking gels. *Nature* **1992**, *358*, 482–485.
- (25) Islam, M. F.; Alsayed, A. M.; Dogic, Z.; Zhang, J.; Lubensky, T. C.; Yodh, A. G.
 Nematic Nanotube Gels. *Phys. Rev. Lett.* **2004**, *92*, 088303.
- (26) Kruse, K.; Joanny, J.-F.; Jülicher, F.; Prost, J.; Sekimoto, K. Asters, vortices, and rotating spirals in active gels of polar filaments. *Physical review letters* 2004, *92*, 078101.
- (27) Voituriez, R.; Joanny, J. F.; Prost, J. Spontaneous flow transition in active polar gels.
 Europhysics Letters (EPL) 2005, 70, 404–410.
- (28) Furuta, K.; Furuta, A.; Toyoshima, Y. Y.; Amino, M.; Oiwa, K.; Kojima, H. Measuring
 collective transport by defined numbers of processive and nonprocessive kinesin motors.
 Proceedings of the National Academy of Sciences 2013, *110*, 501–506.
- (29) de Gennes, P. G.; Prost, J. The physics of liquid crystals (second edition); Oxford
 university press, 1993.
- (30) Kaznacheev, A. V.; Bogdanov, M. M.; Taraskin, S. A. The nature of prolate shape of
 tactoids in lyotropic inorganic liquid crystals. J. Exp. Theo. Phys. 2002, 95, 57–63.
- (31) Keber, F. C.; Loiseau, E.; Sanchez, T.; DeCamp, S. J.; Giomi, L.; Bowick, M. J.;
 Marchetti, M. C.; Dogic, Z.; Bausch, A. R. Topology and dynamics of active nematic
 vesicles. *Science* 2014, *345*, 1135–1139.
- (32) Martínez-Prat, B.; Ignés-Mullol, J.; Casademunt, J.; Sagués, F. Selection mechanism
 at the onset of active turbulence. *Nature Physics* 2019,
- (33) Maitra, A.; Srivastava, P.; Rao, M.; Ramaswamy, S. Activating Membranes. *Phys. Rev. Lett.* 2014, *112*, 258101.

- (34) DeCamp, S. J.; Redner, G. S.; Baskaran, A.; Hagan, M. F.; Dogic, Z. Orientational
 order of motile defects in active nematics. *Nature Materials* 2015, 14, 1110 EP –.
- (35) Opathalage, A.; Norton, M. M.; Juniper, M. P. N.; Langeslay, B.; Aghvami, S. A.;
 Fraden, S.; Dogic, Z. Self-organized dynamics and the transition to turbulence of confined active nematics. *Proceedings of the National Academy of Sciences* 2019, 116,
 4788–4797.
- (36) Edwards, S. A.; Yeomans, J. M. Spontaneous flow states in active nematics: A unified
 picture. EPL (Europhysics Letters) 2009, 85, 18008.
- 460 (37) Seung, H. S.; Nelson, D. R. Defects in flexible membranes with crystalline order. *Phys.*461 *Rev. A* 1988, *38*, 1005–1018.
- (38) Frank, J. R.; Kardar, M. Defects in nematic membranes can buckle into pseudospheres. *Phys. Rev. E* 2008, 77, 041705.
- (39) Henkin, G.; DeCamp, S. J.; Chen, D. T. N.; Sanchez, T.; Dogic, Z. Tunable dynamics of
 microtubule-based active isotropic gels. *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences* 2014, 372.
- (40) Braun, M.; Lansky, Z.; Hilitski, F.; Dogic, Z.; Diez, S. Entropic forces drive contraction
 of cytoskeletal networks. *BioEssays* 2016, *38*, 474–481.
- (41) Zhou, S.; Sokolov, A.; Lavrentovich, O. D.; Aranson, I. S. Living liquid crystals. Proceedings of the National Academy of Sciences 2014, 111, 1265–1270.
- (42) Nelson, M. R.; Howard, D.; Jensen, O. E.; King, J. R.; Rose, F. R. A. J.; Waters, S. L. Growth-induced buckling of an epithelial layer. *Biomechanics and Modeling in Mechanobiology* **2011**, *10*, 883–900.
- (43) Lin, Y.-C.; Koenderink, G. H.; MacKintosh, F. C.; Weitz, D. A. Viscoelastic Properties
 of Microtubule Networks. *Macromolecules* 2007, 40, 7714–7720.

- (44) Gentry, B.; Smith, D.; Käs, J. Buckling-induced zebra stripe patterns in nematic F-476 actin. Physical Review E 2009, 79, 031916. 477
- (45) Liu, Y.; Guo, Y.; Valles, J. M.; Tang, J. X. Microtubule bundling and nested buckling 478 drive stripe formation in polymerizing tubulin solutions. Proceedings of the National 479 Academy of Sciences **2006**, 103, 10654–10659. 480
- (46) Bois, J. S.; Jülicher, F.; Grill, S. W. Pattern Formation in Active Fluids. *Physical* 481 Review Letters 2011, 106, 028103. 482
- (47) Wensink, H. H.; Dunkel, J.; Heidenreich, S.; Drescher, K.; Goldstein, R. E.; Löwen, H.; 483 Yeomans, J. M. Meso-scale turbulence in living fluids. Proceedings of the National 484 Academy of Sciences **2012**, 109, 14308–14313. 485
- (48) Kumar, A.; Maitra, A.; Sumit, M.; Ramaswamy, S.; Shivashankar, G. V. Actomyosin 486 contractility rotates the cell nucleus. Scientific Reports 2014, 4, 3781 EP -. 487
- (49) Duclos, G.; Erlenkämper, C.; Joanny, J.-F.; Silberzan, P. Topological defects in confined 488 populations of spindle-shaped cells. Nature Physics 2016, 13, 58 EP -. 489
- (50) Duclos, G.; Blanch-Mercader, C.; Yashunsky, V.; Salbreux, G.; Joanny, J.-F.; Prost, J.; 490 Silberzan, P. Spontaneous shear flow in confined cellular nematics. *Nature Physics* 491 **2018**, 14, 728–732. 492
- (51) Zadorin, A. S.; Rondelez, Y.; Gines, G.; Dilhas, V.; Urtel, G.; Zambrano, A.; Galas, J.-493 C.; Estevez-Torres, A. Synthesis and materialization of a reaction-diffusion French flag 494 pattern. Nature Chemistry 2017, 9, 990.

495

(52) Furuta, K.; Furuta, A. Re-engineering of protein motors to understand mechanisms bi-496 asing random motion and generating collective dynamics. Current Opinion in Biotech-497 nology **2018**, *51*, 39–46. 498

- ⁴⁹⁹ (53) Subramanian, R.; Gelles, J. Two distinct modes of processive kinesin movement in
 ⁵⁰⁰ mixtures of ATP and AMP-PNP. *The Journal of general physiology* **2007**, *130*, 445–
 ⁵⁰¹ 455.
- ⁵⁰² (54) Sanchez, T.; Dogic, Z. *Methods in Enzymology*; Elsevier, 2013; Vol. 524; pp 205–224.
- 503 (55) Strübing, T.; Khosravanizadeh, A.; Vilfan, A.; Bodenschatz, E.; Golestanian, R.;
- ⁵⁰⁴ Guido, I. Wrinkling instability in 3D active nematics. arXiv:1908.10974