Title
Physical properties of the cytoplasm modulate the rates of microtubule growth and shrinkage

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Abstract

The cytoplasm represents a crowded environment whose properties may change according to physiological or developmental states. Although the effects of crowding and viscosity on in vitro reactions have been well studied, if and how the biophysical properties of the cytoplasm impact cellular functions in vivo remain poorly understood. Here, we probed the effects of cytoplasmic concentration on microtubule (MT) dynamics by studying the effects of osmotic shifts in the fission yeast *Schizosaccharomyces pombe*. Increasing cytoplasmic concentration by hyperosmotic shock led to proportionate reductions in the rates of interphase MT growth and shrinkage. Conversely, dilution of the cytoplasm in hypoosmotic shifts led to proportionately faster rates. Numerous lines of evidence indicate that these effects were due to biophysical properties of the cytoplasm. These effects were recapitulated in in vitro reconstituted MT assays by modulating viscosity, not by crowding. Our findings suggest that even at normal conditions, the viscous properties of cytoplasm modulate the dynamic reactions of MT polymerization and depolymerization.
Introduction

The properties of the cytoplasm and its effects on cellular processes remain poorly understood. Cytoplasm is composed of 100-300 mg/ml of macromolecules (proteins, nucleic acids, lipids, etc), which occupy 10-40% of the total cellular volume (Milo & Phillips, Cell Biology by the Numbers, 2018). These components range in size from small globular proteins to extended networks of organelles and cytoskeletal polymers. Ribosomes alone occupy ~20% of that volume (Delarue et al., 2018). Biophysical studies reveal that these constituents form a porous-elastic material that allows diffusion of small molecules but impedes movement of larger particles (Luby-Phelps et al., 1986; Moeendarbary et al., 2013; Xiang et al., 2020), and molecular simulations suggest a highly dense packing of macromolecules (Yu et al., 2016). Unlike the density of inert matter, recent findings show that cytoplasmic density is not constant. Rather, cells regulate and vary their density as part of normal cell physiology: during the cell cycle, among different cell types, aging, nutritional responses and disease states (Neurohr and Amon, 2020). There is evidence though that the density of cytoplasm may be near functional limits (Dill et al., 2011) as increasing its concentration can lead to large effects including solidification of the cytoplasm (Heimlicher et al., 2019; Joyner et al., 2016; Munder et al., 2016). These density changes, which potentially affect cytoplasmic concentration and crowding effects, further underscore the importance of understanding effects of cytoplasmic concentration. How molecular reactions are able to function effectively within this crowded intracellular environment remains a critical open question.

There are several conceptual models used to describe the influence of macromolecular crowding. Minton and colleagues have argued that bulky crowding agents “exclude volume”, which increases the effective concentration and activity of other enzymes (Minton, 2006; Shahid et al., 2017). Nonspecific interactions between macromolecules in the cytoplasm can affect the rates and equilibria on reactions in numerous ways including macromolecular crowding, confinement and adsorption (Minton, 2006). Macromolecular crowding can impede movement, produce forces that draw molecules together, for instance to promote phase separation, as well as produce osmotic pressure within cells (Mitchison, 2019; R.John, 2001; Shahid et al., 2017). Although effects of crowding have been studied largely in theory and in in vitro experiments, the extent to which these theories and experiments apply to cellular reactions in vivo remains debatable. For example, Mitchison used the concept of colloidal osmotic pressure to argue that cytoplasm is functionally dilute, such that reactions such as microtubule (MT) polymerization are unaffected by crowding effects like excluded volume, perhaps because many macromolecules are well organized in large complexes (Mitchison, 2019). To distinguish between these concepts, what is needed are experiments that perturb the properties of cytoplasm and measure the rates of cellular reactions.

MT dynamic instability represents an attractive case with which to probe the effects of the cytoplasm on a defined molecular reaction in vivo. First, the polymerization and depolymerization of single MTs are reactions that can be quantitatively measured in living cells as microtubule length changes using microscopy. Second, the effects of macromolecular crowding on MTs in vitro are known, as are the effects of viscosity alone: MTs grow significantly faster in the presence of bulky crowders like polyethylene glycol but significantly slower in the presence of small viscous agents like glycerol (Wieczorek et al., 2013). These in vitro measurements can be compared to in vivo measurements when considering mechanisms. Third, the mechanisms driving MT dynamics have been extensively studied, and the biological importance of these dynamics is well established. At 100 kDa in mass and 8 nm in length, the tubulin dimer represents a size
range typical for soluble proteins and enzymes. Finally, MT polymerization is dependent on tubulin concentration while MT depolymerization is not (Fygenson et al., 1994; Walker et al., 1988). Thus, changes in tubulin concentration (e.g., because of changes in cytoplasmic concentration) should impact polymerization alone, which is a testable prediction of some models. Taken together, microtubule dynamic instability is well-suited to discover which properties of cytoplasm are having the strongest impact on cellular reactions, and more generally, to inform biophysical models describing the physical properties of cytoplasm. As the cytoplasm may exhibit both crowding and viscous properties, studying which of these (or other properties) affect reactions such as MT growth and shrinkage in vivo may elucidate relevant properties of the cytoplasm.

Fission yeast is an excellent model organism with which to study the physical regulation of MT dynamics in vivo. We can readily image the interphase microtubule bundles and measure the dynamic behavior of individual microtubules with precision (Höög et al., 2007; Loiodice et al., 2019; Sawin and Tran, 2006). Importantly, we can readily manipulate the properties of their cytoplasm using osmotic shifts, which create robust and well characterized changes in cellular volume and cytoplasm concentration (Atilgan et al., 2015; Knapp et al., 2019).

Here, we study the effects of the properties of the cytoplasm on MTs in fission yeast by using osmotic shifts to vary cytoplasmic concentration. We show that hyperosmotic shifts, which increase cytoplasmic concentration, lead to dampening and “freezing” of MT growth and shrinkage. Conversely, hypoosmotic shifts, which decrease cytoplasmic concentration, lead to increased rates of MT polymerization and depolymerization. The observed MT effects, which were independent of the osmotic stress response and key microtubule regulators, correlated with global changes in cytoplasmic biophysical properties and were recapitulated in vitro through modulation of viscosity and in silico by variation the association rate constant. These findings demonstrate that cytoplasm is not functionally dilute but modulates MT dynamics through viscous effects even at normal concentrations of cytoplasm.

Results

Cytoplasm concentration tunes microtubule dynamics

The density of cellular components in a cell can be experimentally manipulated by varying its osmotic environment (Knapp et al., 2019). We treated live fission yeast cells with an external osmotic agent (addition of sorbitol in the growth medium), which led to an acute loss of cell volume in a dose-dependent, reversible manner (Fig. 1 A and B) (Sup. Fig. 1 and Sup. Fig. 2) (Atilgan et al., 2015; Knapp et al., 2019). For instance, 1 M sorbitol in YE (YE5S rich medium) caused cells to decrease 40% in volume, while 1.5 M sorbitol led to a 60% decrease in volume, all without loss in cell viability (Sup. Fig. 1). This volume loss presumably occurs through water loss. Increases in fluorescence intensity of GFP-labeled tubulin and a ribosomal protein confirmed that cytoplasmic proteins increased in concentration inversely proportional to volume changes (Sup. Fig. 1).

Having validated this approach to alter cytoplasmic concentration, we applied it to cells expressing GFP-tubulin. Hyperosmotic shifts caused striking effects on the dynamic behaviors of interphase MTs, as noted previously (Robertson and Hagan, 2008; Tatebe et al., 2005). We found that in acute response to sorbitol shifts, the interphase MT cytoskeleton was largely maintained (Fig. 1 A). Time lapse imaging showed that while interphase MTs in untreated cells were characteristically dynamic with little time in pause, increasing concentrations of sorbitol caused MTs to “freeze”, especially at high sorbitol
concentrations (1.5 M) (Fig. 1 B, C and D). In general, MTs were paused at various lengths and exhibited little or no growth or shrinkage. To determine whether the observed sorbitol effects were reversible, we treated cells with osmotic oscillations in which sorbitol-containing medium was introduced and then replaced with medium lacking sorbitol in repeated cycles at 5-min intervals. Upon each hyperosmotic shift to 1.5 M sorbitol, most of the MTs were “frozen” within 30 sec (Fig. 1 B and C). Upon shift back to sorbitol-free medium, within 30 s, all interphase MTs started to shrink toward the middle of the cell, and then regrew back in <10 min (Fig. 1 B and Sup. Fig. 2), consistent with previous observations (Alasdair and Hagan, 2008). This uniform shrinkage may be a consequence of the loss of the stabilizing MT GTP cap during the frozen state. This cycle of MT “freezing” and resumption of dynamics occurred repeatedly (Sup. Fig. 2), demonstrating that the effects of sorbitol were rapid (timescales <30 s) and reversible.

The effects on MT dynamics were dependent on the sorbitol concentration. We detected progressive increases in both the time that MTs spent in a pause state (Fig. 1 C) and the number of MTs in a pause state (Sup. Fig. 3). At 1 M sorbitol, ~20% of MTs were frozen, while at 1.5 M, 70% of MTs appeared frozen over the observation period (Sup. Fig. 3). Of the MTs that continued to be dynamic, their rates of polymerization and depolymerization decreased in a sorbitol dose-dependent manner (Fig. 1 C, E, and F, and Sup. Fig. 3). Importantly, sorbitol’s effects on MT polymerization and depolymerization were proportional; for instance, at 1 M sorbitol, both polymerization and depolymerization rates decreased by ~40% (Fig. 1 E and F). Treatment with high sorbitol also made MTs resistant to depolymerization by cold temperature (Sup. Fig. 4), further indicating that these MTs were in a highly stabilized state.

We next asked whether hypoosmotic shifts, which effectively dilute the cytoplasm, yield effects on MT dynamics opposite to those prompted by hyperosmotic shifts. Our initial attempts to swell intact fission yeast cells with hypoosmotic shifts were not successful: cells did not increase in volume very much and any effects were transient, possibly because the cell wall impeded swelling (Nakayama et al., 2012). However, protoplasts (cells with the cell wall removed) swelled substantially in response to hypoosmotic shifts without lysing (Fig. 2 A) (Lemière and Berro, 2018). Shifting protoplasts to hypotonic and hypertonic conditions led to predictable changes in cell volume over a ~2.5-fold range (Fig. 2 A). We determined that protoplasts prepared in 0.4 M sorbitol in YE medium were in isotonic conditions, as their average volume matched that of intact cells in YE medium without sorbitol, and the diffusion rates of GEMS (or nano) particles also matched those in intact cells (see below and Methods). MTs were detectable in protoplasts across all volumes (Fig. 2 B). Hyperosmotic shifts in protoplasts led to proportionate decrease in MT polymerization and depolymerization rates, similar to what was observed in cells with an intact cell wall (Fig. 2 C to F). In hypoosmotic shifts, MT polymerization and depolymerization rates increased proportionally over isotonic conditions (Fig. 2 C to F). Notably, when cell volume increased by ~40 % over isotonic conditions (corresponding to a ~30% dilution of the cytoplasm), both MT polymerization and depolymerization rates increased by ~50% (Fig. 2 E and F). This result implies that cytoplasm dampens the rates of MT polymerization and depolymerization at “normal” cytoplasmic concentration.

To evaluate whether these effects arise in other cell types, we performed similar osmotic shifts with the moss Physcomitrium (Physcomitrella) patens. Caulonema cells expressing mCherry-labelled tubulin and citrine-labelled EB1 displayed lower MT polymerization and depolymerization rates when the osmolarity of the medium was increased with sorbitol (Sup. Fig. 5). This conservation suggested that these effects arise from conserved properties of the cytoplasm.
Normalization of the MT polymerization and depolymerization rates to the isotonic conditions for both yeast cells and protoplasts revealed how changes in the concentration of the cytoplasm, above and below normal levels, caused similar graded responses for MT polymerization and depolymerization (Fig. 2E and F). This proportionate response suggests that the property of the cytoplasm that is changed by osmotic shocks affects both MT polymerization and depolymerization in a similar manner. Interestingly, MT dynamics decreased when the cytoplasm concentration and tubulin concentration increased (Fig. 1 and 2), opposite to what would be expected if crowding or tubulin concentration were responsible for MT dynamics.

**Effect of osmotic shifts on MT dynamics is independent of stress response and regulatory proteins at MT plus ends**

We next investigated several plausible mechanisms for these effects of the cytoplasm on MT dynamics. We first considered indirect mechanisms that operate through osmotic shock response pathways (such as regulation through phosphorylation) and/or regulation by MT regulators at the MT plus end.

First, we tested whether MT stabilization is a downstream effect of an osmotic stress response pathway. Cells exhibit osmotic stress responses that allow them to adapt and survive stresses; the MAP kinase Sty1 (Hog1, p38 ortholog) is a master integrator of multiple stress pathways (Perez and Cansado, 2011). Sorbitol-mediated osmotic shifts applied to sty1Δ cells caused a decrease in MT dynamics (Sup. Fig. 6) very similar to the decrease observed in wild-type cells (Fig. 1) as previously observed (Robertson and Hagan, 2008). Other triggers of the Sty1 stress pathways, such as latrunculin A, do not produce MT “freezing” (Daga et al., 2006). Thus, “freezing” of the MT network is not a downstream response of Sty1-dependent stress pathways.

We next explored the role of MT regulatory proteins at MT plus ends (+TIPs). These proteins could cap and stabilize MTs during hyperosmotic shifts and/or drive more rapid dynamics during hypo-osmotic shifts. Numerous +TIP proteins have been characterized in fission yeast, with the major classes represented by the EB-family +TIP protein Mal3 (EB1) that binds the GTP-cap and the XMAP215-family polymerase Alp14 (XMAP215) (Akhmanova and Steinmetz, 2010; Al-Bassam et al., 2012; Busch and Brunner, 2004). In hyperosmotic shifts, mal3Δ and alp14Δ mutant cells exhibited similar dampening of MT dynamics as wild-type cells (Sup. Fig. 7). As most other +TIP proteins are dependent on Mal3 for localization these two mutants effectively capture the function of most of the +TIPs; therefore, the stabilization is not dependent on these +TIPs. Imaging during hyperosmotic shifts showed that Alp14-GFP was maintained at the MT plus ends during the shift, but Mal3-GFP localization to plus ends decreased (Sup. Fig. 8), consistent with hydrolysis of the GTP-cap (Guesdon et al., 2016) which explains the rapid collapse of the MT following reversal of the osmotic shock (Fig. 1B and Sup. Fig. 2).

Taken together, these observations demonstrate that the acute effect of increased cytoplasm concentration on MT dynamics observed here is independent of the osmotic stress response and the presence of +TIPs. An attractive alternative model is that the assembly and disassembly of MTs are affected directly by biophysical properties of the cytoplasm that scale with its concentration.

**Cytoplasm concentration modulates nanoparticles diffusion**

To characterize how osmotic shifts affect the biophysical properties of the cytoplasm, we carried out microrheology experiments with GEMs. GEMs are spherical nanoparticles of defined sizes composed
of fluorescently labeled non-eukaryotic proteins (Delarue et al., 2018). We expressed gene fusions encoding GEMs fused with the Sapphire fluorescent protein in fission yeast (Methods) and imaged them at rapid frame rates (~100 fps) to analyze their motion (Fig. 3 A). We analyzed GEMs of a range of sizes (20, 40, and 50 nm in diameter) similar in scale to the widths of a MT (24 nm), a ribosome (~25 nm), a proteasome (~15 nm), and a XMAP215/Alp14-tubulin complex (19 nm diameter), but larger than a tubulin dimer (4 by 8 nm). Mean squared displacement (MSD) plots revealed that the movements of the GEMs were sub-diffusive (α < 1) (Sup. Fig. 9), as observed in other cell types (Delarue et al., 2018). The effective diffusion coefficients (D_{eff}) extracted from the MSD plots showed that GEMs motion was size-dependent: larger particles diffused more slowly than the smaller GEMs (Fig. 3 B), consistent with previous rheological observations in other cell types (Luby-Phelps et al., 1986) (Moeendarbary et al., 2013).

Having characterized GEMs motion in yeast cells, we measured the impact of osmotic shocks on their diffusion. At 1 M sorbitol, 40-nm GEMs and 50-nm GEMs were effectively immobile, while 20-nm GEMs still moved (Fig. 3 B). At 1.5 M sorbitol, GEMs of all three sizes largely stopped moving in the cytoplasm (Fig. 3 B).

We also analyzed the motion of the 40-nm GEM in protoplasts to investigate the effect of hypoosmotic shocks, which decrease cytoplasm concentration on GEMs diffusion. We confirmed that the D_{eff} values for GEMs in protoplasts exposed to 0.4 M sorbitol were similar to those in intact cells (Fig. 3 D), consistent with this concentration being the isotonic point of the normal cytoplasm. GEMs mobility increased in hypoosmotic shifts (Fig. 3 C), consistent with decreased cytoplasm concentration and decreased in hyperosmotic shifts, consistent with increased cytoplasm concentration. These observations on yeast cells and protoplasts suggest that osmotic shifts caused substantial changes in cytoplasm biophysical properties.

Normalization of GEMs diffusion to the isotonic condition in yeast and protoplasts revealed that osmotic shifts affect MT dynamics and the diffusion of nanoparticles within the cytoplasm in roughly similar manners (Fig. 3 D), further supporting the hypothesis that physical properties of the cytoplasm affect both GEMs and MTs. Both followed the general inverse trend of slower dynamics with increasing cytoplasmic concentration. Thus, the GEMs data show that general biophysical properties of the cytoplasm (encountered by particles the size of GEMs) correspond with the changes in MT dynamics. during osmotic shifts. The slower diffusion of GEMs throughout the cytoplasm in hyperosmotic shifts suggests that diffusion of even smaller proteins like tubulin may also be slower in these conditions.

**Microtubule dynamics scale with tubulin diffusion in cells**

As MT dynamics depend on the motion of tubulin subunits, we next directly assessed the effects of osmotic shifts on the diffusion of tubulin dimers. We measured the diffusion of soluble tubulin by fluorescence loss in photobleaching (FLIP) experiments. Fluorescence intensity in the cell is measured while a small region of the cell is repeatedly photobleached. The rate at which fluorescence decays over time is proportional to the diffusion of the fluorescent probe used (Ishikawa-Ankerhold et al., 2012). To our knowledge, the rate of tubulin diffusion has been measured only in sea urchin (Salmon et al., 1984) and in PTK2 cells (Wang et al., 2004). As a marker for soluble α/β tubulin dimers, we used Atb2-GFP (α-tubulin 2) expressed from the native chromosomal locus (Sato et al., 2009). Cells were treated with the MT inhibitor methyl benzimidazol-2-yl-carbamate to depolymerize MTs, and then we repeatedly photobleached a ~1 μm region of the cytoplasm (Fig. 4 A and B). We then measured the decay in GFP fluorescence in the cell (Methods).
To estimate the tubulin diffusion rate, we designed a 1D stochastic model of diffusion in which the only free parameter is tubulin diffusion (Methods). Comparison of the model predictions with our experimental data (Fig. 4 B) yielded an estimated diffusion rate of GFP-tubulin of 7 μm$^2$ s$^{-1}$ in cells in YE without sorbitol; very close to the value of ~6 μm$^2$ s$^{-1}$ obtained previously in other cell types (Salmon et al., 1984) (Wang et al., 2004). According to our model, diffusion rates decreased to 4 μm$^2$ s$^{-1}$ in YE + 1 M sorbitol and 1.5 μm$^2$ s$^{-1}$ in YE + 1.5 M sorbitol (Fig. 4 C); the latter value represents a decrease of up to 5x in tubulin dimer diffusion. Interestingly, these changes in tubulin diffusion were linearly correlated with the rates of MT polymerization and depolymerization (Fig. 4 D). Taken together, these data suggest that tubulin diffusion may constitute a rate-limiting factor in these circumstances (Discussion).

**Viscosity limits microtubule dynamics**

In order to determine whether changes in viscosity could explain the changes in tubulin diffusion and MT growth and shrinkage rates resulting from osmotic shifts, we measured the viscosity of the fission yeast cytoplasm. We estimated viscosity of the fission yeast cytoplasm using time-resolved Fluorescence Anisotropy Imaging (tr-FAIM) (Puchkov, 2012). This method measures the movement of a fluorescent dye (fluorescein) in the cytoplasm of living cells, assessing molecular movements of the size scale of the dye and water (see Methods; Sup. Fig. 10). Fitting and extraction of the rate constants and comparison to the calibration curve yielded a viscosity value of ~ 9 cP, which is in line with the broad range of previous viscosity measurements for eukaryotic cytoplasm (range from 1 to 50 cP) (Obodovskiy, 2019). This value suggests that the fission yeast cytoplasm has a viscosity similar to that of 53% (v/v) glycerol (in water). For cells treated with 1.5M sorbitol, we obtained a slightly higher value of 12 cP, corresponding to 57% (v/v) glycerol. This result suggests that the viscosity of the cytoplasm increases with its concentration following hyperosmotic shocks. As diffusion scales inversely with viscosity, such changes in viscosity may explain the effects of the osmotic shift on GEMs and tubulin diffusion rates.

In order to isolate the effects of viscosity alone on MT dynamics, we repeated these experiments in a reconstituted system. (Fig. 5 A). Although it was previously shown that viscosity decreases MT polymerization rates in vitro (Wieczorek et al., 2013) the constrains of the fluorescence-based MT dynamic assays used limited the time-lapse recording length and frequency and did not allow its effect on MT shrinkage rate to be quantitatively studied. We therefore took advantage of Interference Reflection Microscopy (IRM) to image MTs at 2 fps, allowing a more accurate quantification of shrinkage rates. We reconstituted MT dynamics in the presence of glycerol as a viscous agent using a well-established assay using purified tubulin (Fig. 5 A) (Gell et al., 2011). A range of glycerol concentrations was added to the reconstitutions (Wieczorek et al., 2013) to produce a range of viscosities from 0.7 to 1.5 cP. MT polymerization and depolymerization rates both decreased proportionally in a dose dependent manner with increasing viscosity (Fig. 5 B and C). Using viscosity of the buffer and the Stokes-Einstein equation we estimated tubulin diffusion in the reconstituted assay (see methods) and saw a linear relationship between both rates and tubulin diffusion (Fig 5. D). Thus, the proportional influence of cytoplasmic properties on MT growth and shrinkage rates that we observed in vivo may be reproduced in vitro by modulating a single parameter, viscosity.

To gain insight into how viscosity influences MT growth and shrinkage rates, we used a simple, 6 parameters model for MT dynamic instability (Hsu et al., 2020) that reproduces the core behaviors of polymerization, catastrophe, and depolymerization (Odde, 1997). In order to model the effect of viscosity, we varied the tubulin association rate constant $k^+$ (Fig 5. E), which defines the rate at which tubulin binds
to protofilament ends. All other parameters, notably bond energies of tubulin-tubulin bonds, remained constant. The model predicts that growth rates decrease with $k^*$, as expected from fewer binding events (Fig. 5 F). However, the model also predicts that depolymerization rates scale linearly with $k^*$ (Fig. 5 G), because when associate rate constants change, dissociation rate constants must also change if the bond energies ($\Delta G_{\text{long}}$ and $\Delta G_{\text{lat}}$) are to remain constant. In other words, the model predicts that both rates scale with the association rate constant.

Thus, our in vitro and in silico experiments paint a consistent picture: that both MT growth and MT shrinkage scale linearly with viscosity. Remarkably, our purified components recapitulate the observations from fission yeast, protoplasts, and moss cells. In all models, rates of MT polymerization and depolymerization are linearly correlated, indicating a conserved ratio of growth to shrinkage (Fig. 5 H). The ratio is different in each case, presumably due to the specific conditions of each case (presence of MAPs, tubulin isoform properties, tubulin concentration, temperature, etc.). Nonetheless, the ratio is maintained when MT dynamics are perturbed, either by changes in cytoplasm concentration in vivo, changes in viscosity in vitro, or changes in the association rate constant in silico.

In addition, normalizing the rates from each model to the value in the unperturbed condition collapsed all the data onto a line which matches the in silico model prediction (Fig. 5 I). By varying a single parameter, $k^*$, the model is able to recapitulate the observations both qualitatively and quantitatively. This suggests that increasing viscosity of the environment is equivalent to reducing the association rate constant.

**Discussion**

Here we probed the effect of cytoplasm on MT dynamics in vivo using osmotic shifts. During hyperosmotic shifts that increase cytoplasm concentration, MTs paused more frequently and polymerized and depolymerized more slowly; at high concentrations, MTs appeared to largely “freeze” (Figure 1). Conversely, in hypoosmotic shock, which decrease cytoplasm concentration, MT polymerization and depolymerization rates speed up (Figure 2). We provide numerous lines of evidence to indicate that these effects act via the physical properties of the cytoplasm such as viscosity, as opposed to regulation of osmotic stress pathways or MT plus end regulators. The effects were acute (Fig. 1), reversible (Sup. Fig. 2) and scaled linearly with cytoplasm concentration in hypo- and hyperosmotic shocks (Fig. 3), strongly indicative of a physical response. The effects on MTs scaled with effects on the diffusive-like movement of GEMs and tubulin dimers. While MT “freezing” in response to hyperosmotic shocks has been noted previously in fission yeast (Tatebe et al., 2005)(Robertson and Hagan, 2008), we obtained two particularly significant results; MT dynamics scale linearly with cytoplasm concentration and, hypoosmotic shifts speed up MT polymerization and depolymerization rates. A significant implication is that even the “normal” concentration of cytoplasm dampens MT dynamic behaviors.

How might the cytoplasm influence MT dynamic behaviors? That MT polymerization and depolymerization are proportionally affected seems like an important clue. This coordinated effect on polymerization and depolymerization is unusual, as known perturbations such as changing tubulin concentration typically affect MT polymerization but not depolymerization rates (Fygenson et al., 1994). Although temperature (a purely physical concept) affects polymerization and depolymerization similarly (Fygenson et al., 1994). Modifying cytoplasmic concentration could alter many parameters including the concentration of molecules such as tubulin, their effective diffusion, viscosity and crowding effects. Our results are the opposite of the expected effects of altered tubulin concentration, which changes MT
polymerization rate without changing depolymerization rate (Fygenson et al., 1994). Similarly, our results are not consistent with crowding playing a decisive role, as addition of bulky crowding agents increase polymerization rate, not slow it down (Wieczorek et al., 2013) and prevent depolymerization (Bachand et al., 2018). This is consistent with a proposal in which macromolecular crowding effects may not be as dominant in the cytoplasm as previously perceived (Mitchison, 2019). Rather, our data support a hypothesis that the predominant effects of the cytoplasm on MT dynamics are via changes in viscosity and diffusion. Importantly, we show that changes in viscosity slow down MT polymerization and depolymerization linearly in a reconstituted system with purified components, similar to what we see in vivo.

Viscosity potentially affects multiple steps in MT polymerization and depolymerization. Viscosity inhibits diffusion of the tubulin dimer, which we show correlates with MT rates (Fig. 4 and 5). MT polymerization has been modeled as a two-step process depending on the rate of collision of tubulin dimer to the end of the protofilaments and the rate of incorporation into the tubule (Hsu et al., 2020; Wieczorek et al., 2013). Viscosity may also inhibit rate of conformational molecular changes, for instance in protofilament dynamics and in compaction of the tubulin dimer within the lattice (Brouhard and Rice, 2018). Our findings are in agreement with recent studies that suggest that similar conformational changes in the protofilaments occur in MT polymerization and depolymerization (Richard McIntosh et al., 2018) (Gudimchuk et al., 2020) (Brouhard and Rice, 2018).

In summary, we use MTs as an example to study how cytoplasmic properties affect biochemical reactions in vivo and discovered that viscosity plays a key role in regulating MTs dynamics. As the concentration and the density and other properties of the cytoplasm are known to vary during the cell cycle, in development, aging and diseases (Neurohr and Amon, 2020), it will be important to consider how cytoplasmic properties such as viscosity and crowding globally affect cytoskeletal and other cellular reactions.
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Material and methods

Fission yeast, media, and growth conditions

Standard methods for growing and genetically manipulating *Schizosaccharomyces pombe* were used (Moreno et al., 1991). For most experiments, cells were grown overnight in Yeast Extract (YE5S) medium to exponential phase (0.2<OD<0.8 at 600nm) with shaking at 30 °C. See Sup. Table. for reagents and strains details.

Strain construction

Cassettes for expressing GEMs (20 nm, AqLs-Sapphire; 40 nm, PfV-Sapphire; 50 nm, Vuldi-Sapphire), were cloned into the fission yeast pREP41X expression vector from pRS306 budding yeast expression vectors. The Vuldi-sapphire pRS306 plasmid was a kind gift from the Holt lab. Briefly, GEMs expression cassettes were amplified via PCR and inserted into pREP41X via Gibson assembly at the XhoI site. The primers used for PCR are described in the supplementary table.

Protoplast preparation

*S. pombe* cells were grown in YE5S liquid culture at 30 °C to exponential phase, harvested, and washed with SCS buffer (20 mM citrate buffer, 1 M D-sorbitol, pH 5.8), then resuspended in SCS buffer supplemented with 0.1 g/mL Lallzyme (Lallemand, Montreal, Canada) (Flor-Parra et al., 2014). Cells were digested for 10 min at 37 °C with gentle shaking in the dark. The resulting protoplasts were gently washed twice in YE5S medium with 0.2-1 M D-sorbitol, using gentle centrifugation (2 min at 0.4 rcf).

Imaging of fission yeast

*S. pombe* cells and protoplasts were imaged in commercial microchannels (Ibidi µ-slide VI 0.4 slides; Ibidi 80606, Ibiditreat #1.5). Channels were pre-treated with 50 µl of 100 µg/ml lectin solution for 5 min at room temperature. The lectin solution was removed by pipetting and 50 µl of cell culture were introduced. After incubation for 3 to 10 minutes to allow cells to adhere to the lectin, the cells were washed with YE5S. For hyper-osmotic shocks, the medium was manually removed from the channel via pipetting and quickly replaced with hyper-osmotic media as indicated.

Microtubule dynamics. Yeast cells expressing GFP-tubulin (see Table) were observed with a 488 nm excitation laser at 100 ms of exposure per z-slice (1 µm spacing, 7 slices) at 0.1 Hz, with a 60x objective (CFI Plan Apochromat VC 60XC WI) on a Nikon TI-E equipped with a spinning-disk confocal head (CSU10, Yokogawa) and an EM-CCD camera (Hammamatsu C9100-13).

Protoplast volume. Protoplasts were resuspended YE5S medium with 0.2-1 M D-sorbitol, then imaged with a 561 nm excitation laser at 100 ms of exposure per z-slice (0.5 µm spacing), with the 60x objective, microscope, and camera used for MT dynamics.

GEMs diffusion. Yeast cells were imaged with a 60x TIRF objective (Nikon, MRD01691) on a Nikon TI-E equipped with a Nikon TIRF system and a SCMOS camera (Andor, Ixon Ultra 888). Protoplasts were imaged on a Nikon TI-2 equipped with a Diskovery Multi-modal imaging system from Andor and a SCMOS camera (Andor, Ixon Ultra 888) using a 60x TIRF objective (Nikon, MRD01691). Cells were imaged at 100 Hz, in TIRF, for 10 s with a 488 nm excitation laser.
**Tubulin diffusion.** Cells were imaged at 0.2 Hz, with a 60x TIRF objective (Nikon, MRD01691) on a Nikon TI-2 equipped with a Diskovery Multi-modal imaging system from Andor and a SCMOS camera (Andor, Ixon Ultra 888). Cells were imaged using spinning disk with pinhole of 100 μm. Cells were imaged at 488 nm laser excitation; the bleaching laser was a 473 nm laser controlled by a UGA-42 Firefly (Rapp OptoElectronic).

**Osmotic shock experiments**

**Hyperosmotic shifts.** Cells were grown in rich YE5S medium, mounted into microchannels, and then treated with YE5S containing various concentrations of sorbitol while on the microscope stage at room temperature. Loss of water after the switch of medium is almost instantaneous (<30 s). To minimize adaptation responses (Tatebe et al., 2005), imaging was initiated as soon as possible (<1 min) after adding sorbitol.

**Osmotic oscillations.** Cells were introduced into a microfluidic system (Cell Asics) as described in (Knapp et al., 2019) then the media in the chamber was oscillated; YE for 5 minutes then YE with 1.5 M sorbitol for 3 minutes. During the oscillations cells were imaged for MT dynamic measurement as described in above.

**Cold treatment.** Yeast cells expressing GFP-tubulin were pelleted gently and resuspended in YE containing 0, 0.5, 1, or 1.5 M sorbitol. Each culture was split into two tubes; one was incubated for 5 min at room temperature and the other was incubated on ice for 5 min. Cells were fixed by adding 16% paraformaldehyde to the medium for a final concentration of 4%. Cell were then imaged in lectin-treated Ibidi chambers.

**Osmotic shifts of protoplasts.** After cell-wall digestion, protoplasts were gently washed twice in YE5S with 0.4 M D-sorbitol using gentle centrifugation (2 min at 0.4 rcf), then placed in the Ibidi chamber for imaging. Medium was exchanged manually with hypo- or hyper-tonic medium right before imaging. YE + 0.4 M sorbitol was close to isotonic conditions, as determined by comparing volumes and GEMs dynamics to yeast cells. Thus, protoplasts resuspended in YE + 0.2, 0.25, or 0.3 M sorbitol were in hypo- tonic conditions, while protoplasts in YE + 0.5 or 1 M sorbitol were in hyper-tonic conditions.

**Measurements of MT dynamic parameters and cell volume in yeast and protoplasts**

**MT dynamics.** Measurements of MT dynamic parameters were obtained using analyses of kymographs of GFP-tubulin expressing cells. Images of individual cells were cropped and multiple MT bundles per cell were selected. Kymographs were made and analyzed with the KymoToolBox plugin of ImageJ (Schneider et al., 2012).

**Volume.** The effects of sorbitol on cell volume were determined from brightfield images (Atilgan et al., 2015). Protoplast volume was measured in 3D from Z-stack fluorescence images of cells expressing markers for the plasma membrane (mCherry-Psy1) using LimeSeg, a Fiji plugin (Schindelin et al., 2012) (Machado et al., 2019).

**FLIP experiments**

Cells expressing GFP-tubulin were mounted in microchannels and treated with 25 μg/ml methyl benzimidazol-2-yl-carbamate (MBC) for >1 min to depolymerize MTs. Cells were then subjected to repeated photobleaching with a focused 473 nm laser in a 1-μm region near the cell tip using a UGA-42 Firefly system from RapOpto. Cells of similar size (~ 12 μm long) were photobleached in order to reduce variability in the resulting data. Fluorescence decay was followed in the half of the cell submitted to the
bleaching sequence to avoid the effects of diffusion around/through the nucleus. Fluorescence decay curves were normalized and aligned to the time point preceding the activation of the bleach sequence.

A calibrated 1D Brownian model of diffusion was used to simulate the fluorescence decay in a 6 μm tube (half of a cell 12 μm in length). Bleaching rate, region size, and position were matched to the experimental setup. Particles positions were updated every 0.01 s. The decay in the number of unbleached particles in the model was read out every 5 s (matching the imaging frequency) and normalized to the total number of particles. The insensitivity of the model to the total number of particles and to the time interval used was established by changing these parameters across three orders of magnitude, without significant effect on the outputs. Decay plots for various diffusion rates in the model were compared to the experimentally measured values to obtain the estimated tubulin diffusion rates.

**Microrheology with GEMs nanoparticles.**

GEM fusion proteins were expressed from pREP41X-based expression vectors from the thiamine-regulated nmt1* promoter (Maundrell, 1990). Transformants containing these plasmids were maintained on EMM-leu medium. The day before imaging, cells were inoculated in EMM-leu medium containing 0.05 μg/mL thiamine to allow a low level of construct expression. These conditions generally produced a few tens of GEM nanoparticles per cell. Overexpression of the GEMs commonly produced cells with single, bright, non-motile aggregates. Cells expressing GEMs were selected for sparse numbers of labeled motile nanoparticles and imaged at 100 Hz intervals. Individual cells were cropped for analysis. Nanoparticles in each cell were tracked using the MOSAIC plugin (Fiji ImageJ), and the effective diffusion rate was determined from mean squared displacement (MSD) plots as previously (Delarue et al., 2018). Briefly, tracks shorter than 10 timepoints were excluded from the MSD analysis. The following fit was used on the first 100 ms to extract the diffusion value for trajectories longer than 10 timepoints: MSD = 4Dt. The following parameters were used for the 2D Brownian dynamics tracking in MOSAIC: radius = 3, cutoff = 0, per/abs = 0.2-0.3, link = 1, and displacement = 6.

**MT dynamics in moss cells**

Moss cells were grown as described previously (Yamada et al., 2016). Caulonemal cells were mounted in microfluidic devices at room temperature as described previously (Kozgunova and Goshima, 2019). For osmotic shock, observation medium was removed manually via aspiration with a syringe, and medium with sorbitol was subsequently introduced with a syringe. Cells were imaged on a Nikon TI-E TIRF system with a 60x TIRF objective (Nikon, MRD01691) and a SCMOS camera (Andor, Ixon Ultra 888).

**MT dynamics in vitro**

*Tubulin preparation.* Tubulin was purified from juvenile bovine brains via cycles of polymerization and depolymerization, as described previously (Ashford and Hyman, 2006). GMPCPP-stabilized MT seeds were prepared by polymerizing a 1:4 molar ratio of tetramethylrhodamine (TAMRA, ThermoFisher Scientific) labeled:unlabeled tubulin (Hyman, 1991) in the presence of GMPCPP (Jena Biosciences) in two cycles, as described previously (Gell et al., 2011).

*MT reconstitution assay.* Dynamic MTs were imaged in a reconstitution assay from surface-bound, stabilized MT seeds (Gell et al., 2011). Cover glass was cleaned, as previously described (Helenius et al., 2006). Two silanized cover glasses (22 x 22 mm and 18 x 18 mm) were separated by multiple strips of double-sided tape on custom-machined mounts to create channels for solution exchange. Channels were prepared for experiments by flowing in anti-TAMRA antibodies (ThermoFisher Scientific) and blocking with 1% Pluronic F-127 for 20 min. Channels were rinsed three times with BRB80 before flowing in seeds
and placing the chamber on the microscope stage, where the objective was heated to 32 °C with a CU-501 Cham(cid)l(e) lens warmer (Live Cell Instrument).

Dynamic MTs were grown from GMPCPP seeds by filling the channel with 10 µM tubulin in reaction buffer: BRB80 (80 mM PIPES-KOH [pH 6.9], 1 mM EGTA, 1 mM MgCl₂) plus 1 mM GTP, 0.1 mg/mL bovine serum albumin, 10 mM dithiothreitol, 250 mM glucose oxidase, 64 nM catalase, and 40 mM D-glucose. Reaction buffer was prepared on ice before being flowed into the channel with a piece of filter paper. A 60% (v/v) glycerol stock solution in BRB80 was added to the indicated final concentrations. For consistency, a large aliquot of tubulin was thawed on the day of each experiment, sub- aliquoted, and stored in liquid nitrogen. A separate sub-aliquot was thawed for each individual experiment. Glycerol concentrations from 0-15% (v/v) were tested. We note that the effects of glycerol concentrations ≥20% were not measurable as no shrinkage events were observed because of the high levels of spontaneous nucleation and inhibition of catastrophes at those glycerol concentrations.

*Interference reflection microscopy.* Dynamic label-free MTs were imaged with interference reflection microscopy (IRM) as described previously (Mahamdeh and Howard, 2019).

*MT dynamics.* MT dynamics was analyzed using kymographs as described above.

*Tubulin diffusion.* Tubulin diffusion rate *in vitro* was estimated using the Stoke-Einstein equation $D = \frac{K_BT}{6\pi\eta r}$, where $T$ is the experimental temperature (32 °C), $\eta$ is the viscosity of the buffer, and $r$ is the radius of a sphere equivalent to the volume of a tubulin dimer (3 nm). Pairwise correlations among MT polymerization rate, depolymerization rate, and tubulin diffusion were analyzed using Pearson’s test performed with GraphPad Prism.

*Cytoplasm viscosity measurements*

Wild-type fission yeast cells were grown in YE overnight at room temperature with agitation to below saturation.

*FDA labeling.* Cells were labeled with 100 µM fluorescein diacetate (Sigma, F7378) for 30 min at room temp with agitation.

*Time-resolved fluorescence anisotropy.* Cells were imaged in a Ibidi μ-slide VI 0.4 (Ibidi 80606, Ibidiitreat #1.5) treated with lectin on an inverted TCS-SP2 microscope (Leica Microsystems, Germany) while the time- and polarization-resolved intensity decay was measured on SPC-150 TCSPC boards (Becker & Hickl, Germany). The emission was split using a polarizing beam splitter. Fluorescein diacetate (FDA) was excited using a picosecond-pulsed (90 ps optical pulse width, 20 MHz repetition rate) 467 nm diode laser (Hamamatsu, Japan) through a 63X 1.2 NA water objective, with a 485 nm dichroic mirror and a 500 nm long pass filter. All measurements were performed at room temperature.

Parallel ($I_\parallel$) and perpendicular ($I_\perp$) intensity decays were used to calculate the anisotropy decay, according to

$$r(t) = (I(t)_\parallel - G \ast I(t)_\perp)/(I(t)_\parallel + G \ast I(t)_\perp)$$

where $z = 1$ to account for the depolarization of the high magnitude objective and $G$ was 0.96. $Z$ was determined by comparing the calculated total intensity decay $I_{tot} = I_\parallel + z \ast G I_\perp$ of fluorescein measured on the 63x objective and measured on a 5x objective, altering the value of $z$ between 2 and 1 until the lifetime was an exact match (Suhling et al., 2014). $G$ is a correction factor to account for the sensitivity of the two detectors and is calculated by taking the tail value of
after tail-matching of the parallel and perpendicular decays by adjustment of the polarizing beam splitter. The resulting time-resolved anisotropy decay was fit to a monoexponentially decaying model according to

\[ G = \frac{I(t)}{I(t)} \]

where \( \theta \) is the rotational correlation time, \( A \) is the amplitude of the decay (in terms of the anisotropy decay, this value is equal to \( r_0 - r \)), and \( y_0 \) is the level to which the anisotropy decays (properly known as \( r \)). The rotational correlation time was converted into a viscosity value in cP using a calibration plot in the relevant viscosity region.

**Fluorescein calibration curve.** Methanol solutions containing various amounts of glycerol (0-30%) were prepared. Solution viscosities at room temperature (22 °C) were established previously using a rheometric expansion system rheometer (ARES) (Kuimova et al., 2008) (Steinmark et al., 2020). Fluorescein (10 μM) was added to the solutions and fluorescence was measured at room temperature to extract the rotational correlation times as described above. Note that the calibration was carried out on a 5x air objective; thus, \( z \) was taken as 2.

**Anisotropy decay.** Anisotropy data were fitted using GraphPad Prism. Fits used 1/STD\(^2\) weighting and were limited to the first 10 ns of recording. Best-fit values for each parameter and condition were compared via two-way ANOVA with Sidak’s multiple comparison test. Decay rate were different with p-value < 0.0001.
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Competing Interests

The authors declare no financial or competing interests.

Author Contributions

Conceptualization: A.T.M. and F.C.; Methodology: A.T.M., J.L, C.E., C-T.H., I.E.S., K.S., G.G., L.H., G.J.B., and F.C.; Formal analysis: A.T.M., J.L, I.E.S; Investigations: A.T.M., C.E., I.E.S; Writing: A.T.M., G.J.B., and F.C.; Review editing: A.T.M., C.E., G.J.B., and F.C.; Visualization: A.T.M.; Supervision: A.T.M.; Project administration and funding acquisition: F.C.

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Supplemental Information

Refers to the webpage for description and access to the supplemental material.
Figure 1: MT growth and shrinkage rates decrease upon hyperosmotic shock in yeast cells.

(A) Brightfield (BF; left) and fluorescence (right) images of interphase MT bundles in two fission yeast cells expressing GFP-tubulin (GFP-tub) upon sequential treatment with YE (medium alone), YE + 1 M sorbitol, and YE + 1.5 M sorbitol.

(B) MT dynamics in a cell treated with oscillations of YE (5 min) and YE + 1.5 M sorbitol (3 min). In the kymograph, this representative cell is expressing GFP-tubulin; the cell image has been collapsed onto a line. MTs exhibit decreased dynamics acutely in 1.5 M sorbitol. Upon sorbitol washout, MTs first depolymerize (yellow arrows) and then resume dynamic behaviors.

(C) Kymographs of MTs in cells at the indicated sorbitol concentrations. Lines highlight tracks of single growing MT plus-ends. Colored lines highlight growth events.

(D) Percentage of time that interphase MTs spent growing (green circles), shrinking (red squares), or paused (blue triangles) in YE and after hyperosmotic shocks (AVG +/- standard deviation). Cell and MT values of n are as in (E).

(E) MT polymerization rates (green circles) in cells treated acutely with the indicated sorbitol concentrations. Values are AVG +/- standard deviation. Data come from (left to right) n = 58 / 51 / 32 / 27 / 38 / 23 / 15 / 26 cells and 118 / 99 / 60 / 65 / 72 / 34 / 22 / 44 MT, respectively, from at least 2 experiments.

(F) MT depolymerization (red squares) rates in cells treated acutely with the indicated sorbitol concentrations. Values are AVG +/- standard deviation. Cell and MT values of n are as in (E).
Figure 2: MT growth and shrinkage rates scale with cell volume in yeast protoplasts.
(A) Normalized protoplast volume. Data were normalized to osmolarity conditions of 0.4 M sorbitol, in which the average volume distribution of protoplasts matched that of intact cells (AVG +/- standard deviation). Data come from at least two experiments; (left to right) n = 18 / 45 / 109 / 446 / 200 / 440 protoplasts.
(B) Representative images of GFP-labelled MTs in protoplasts in medium supplemented with the indicated sorbitol concentrations.
(C) MT polymerization (Pol.) and (D) depolymerization (Depol.) rates in yeasts protoplasts in medium supplemented with the indicated sorbitol concentrations. Values are AVG +/- standard deviation. Left to right, n = 10 / 13 / 64 / 29 / 25 / 13 polymerization events and n = 7 / 57 / 25 / 13 / 12 depolymerization events from three experiments.
(E) MT polymerization rate and (F) depolymerization rates, normalized to the isotonic condition for yeast cells (circles) and yeast protoplasts (diamonds), as a function of the normalized volume. Both rates are increase in hypo-tonic conditions (blue shading) and decrease in hyper-tonic conditions (orange shading).
Figure 3: Nanoparticles diffusion rate scales with cytoplasm concentration.

(A) Representative images of GEMs in fission yeast cells. Scale bar, 5 μm.

(B) Hyperosmotic shifts decrease the effective diffusion coefficients of GEMs of indicated sizes in yeast cells. Values are AVG +/- standard deviation. Data come from three experiments, n > 1000 trajectories, and n > 49 cells. Concentrations reflect sorbitol concentrations in the medium.

(C) Effective diffusion coefficients of 40-nm GEMs in yeast protoplasts as a function of sorbitol. AVG +/- standard deviation. Data come from at least two experiments and (left to right) n = 643 / 411 / 304 / 488 / 162 trajectories.

(D) Diffusion rate of the 40-nm GEMs in yeast cells (circles) and yeast protoplasts (diamonds) normalized to the iso-tonic condition as a function of the normalized volume. The rate of diffusion through the cytoplasm is faster in hypo-tonic conditions (blue shading) and slower in hyper-tonic conditions (orange shading).
Figure 4: Hyperosmotic shifts decrease the diffusion rate of soluble tubulin.

(A) To measure the diffusion of soluble GFP-tubulin, we used FLIP. Cells were exposed repeatedly to a focused laser beam (~1 μm) near one cell tip and GFP fluorescence intensity was measured (Methods).

(B) Fluorescence decay of cells expressing GFP-tubulin in a representative FLIP experiment at the indicated sorbitol conditions. Interphase cells in which MTs were depolymerized with 25 μg/ml methyl benzimidazol-2-yl-carbamate (MBC) were photobleached using a focused laser (blue stars). Scale bars, 4 μm.

(C) Loss of fluorescence intensity in cells from three osmotic conditions and the corresponding tubulin diffusion rate. Values (AVG +/- standard deviation) were normalized to initial intensity and expressed as concentrations (%). Data are n = 46 / 29 / 29 cells, for YE alone, YE + 1 M sorbitol, and YE + 1.5 M sorbitol, respectively, from three independent experiments. Dashed lines denote predictions from simulations of a 1D model (Methods) for various values of diffusion; these predictions were used to estimated diffusion rates from our experimental data. Simulation values are AVG +/- standard deviation for 5 simulations.

(D) Rates of MT polymerization and depolymerization in yeast cells as a function of tubulin diffusion rate. Data come from Figures 1 and 4. Correlations between diffusion and polymerization rate (P = 0.008) and depolymerization rate (P = 0.001) are significant according to Pearson's correlation test.
Figure 5: Increasing viscosity dampens rates of MT polymerization and depolymerization.
(A) Schematic of the in vitro reconstituted system for MT dynamics measurement.
(B) MT polymerization rates (green circles) and (C) MT depolymerization rates (red squares) were measured in MTs in solutions of varying viscosity. Data represent 3 repetitions with n > 70 MTs per condition.
(D) Rates of microtubule polymerization and depolymerization in vitro as a function of tubulin diffusion rate. Correlation is significant, p-value = 0.002 and 0.02 for polymerization and depolymerization rates respectively (Pearson correlation test).
(E) Schematic of the parameters used to model MT dynamic instability. Changing the association rate constant k+ was used to model the effect of viscosity. For more details see Methods.
(F) Apparent on-rate as a function of the association rate constant in the model.
(G) Apparent off-rate as a function of the association rate constant in the model.
(H) Scatter plot of the observed depolymerization rates versus the observed polymerization rates for the experiments in this study (moss (green circles), yeast cells (grey circles), yeast protoplasts (blue circles), in vitro (orange circles)). Each model presents a certain ratio (the slope of the regression) of depolymerization rate to polymerization rate but this ratio (the slope of the regression) is conserved when viscosity is increased in vitro or when cell volume is reduced by hyper-osmotic shocks (yeast, protoplasts, and moss). AVG +/- STD.
(I) Cytoplasm and viscosity have similar effects of MT polymerization and depolarization rates. Graph shows the relationship between normalized depolymerization rates and normalized polymerization rates for all the experimental conditions and the model prediction. AVG +/- propagated error.