

Behaviors of individual microtubules and microtubule populations relative to critical concentrations: dynamic instability occurs when critical concentrations are driven apart by nucleotide hydrolysis

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ABSTRACT The concept of critical concentration (CC) is central to understanding the behavior of microtubules (MTs) and other cytoskeletal polymers. Traditionally, these polymers are understood to have one CC, measured in multiple ways and assumed to be the subunit concentration necessary for polymer assembly. However, this framework does not incorporate dynamic instability (DI), and there is work indicating that MTs have two CCs. We use our previously established simulations to confirm that MTs have (at least) two experimentally relevant CCs and to clarify the behavior of individuals and populations relative to the CCs. At free subunit concentrations above the lower CC ($CC_{Elongation}$), growth phases of individual filaments can occur transiently; above the higher CC ($CC_{NetAssembly}$), the population's polymer mass will increase persistently. Our results demonstrate that most experimental CC measurements correspond to $CC_{NetAssembly}$, meaning that "typical" DI occurs below the concentration traditionally considered necessary for polymer assembly. We report that [free tubulin] at steady state does not equal $CC_{NetAssembly}$, but instead approaches $CC_{NetAssembly}$ asymptotically as [total tubulin] increases, and depends on the number of stable MT nucleation sites. We show that the degree of separation between $CC_{Elongation}$ and $CC_{NetAssembly}$ depends on the rate of nucleotide hydrolysis. This clarified framework helps explain and unify many experimental observations.

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Abbreviations used: CC, critical concentration; DI, dynamic instability; GDP, guanosine disphosphate; GTP, guanosine triphosphate; J, flux; MT, microtubule; PF, protofilament; SD, standard deviation.

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INTRODUCTION

The concept of critical concentration (CC) is fundamental to experimental studies of biological polymers, including microtubules (MTs) and actin, because CC is commonly understood to be the amount of subunit needed to obtain polymer. CC is used to characterize different polymers and to interpret the effects of polymer assembly regulators. In the standard framework for predicting the behavior of biological polymers, there is one CC, at which polymer assembly commences (e.g., Mirigian et al., 2013; Alberts et al., 2015). However, as indicated by other work (Hill and Chen, 1984; Walker et al., 1988), this framework fails to account for the dynamic instability (DI) displayed by MTs and other dynamically unstable polymers (e.g., PhuZ, ParM; Mitchison and Kirschner, 1984a; Garner et al., 2004; Erb et al., 2014). One purpose of the

work presented here is to examine the many experimental and theoretical definitions of CC in order to show how the definitions relate to each other. Another purpose is to clarify how the behaviors of individual dynamically unstable filaments and those of their populations relate to each other and to experimental measurements of CC. To address these problems, we simulated systems of dynamic MTs with one of the two ends of each MT fixed at a stable seed (also called a nucleation site or template), similar to MTs growing from centrosomes. We performed analyses that are directly comparable to those used in experiments. A significant advantage of computational modeling for this work is that it allows simultaneous examination of the behaviors of individual subunits, individual MTs, and the population's bulk polymer mass.

Traditional understanding of critical concentration based on equilibrium polymers

Traditionally, "the critical concentration" is understood to be the concentration of subunits needed for polymer assembly to occur ($CC_{PolAssem}$). Specifically, the textbook understanding is that polymers will grow at concentrations above $CC_{PolAssem}$, and will not grow below it. $CC_{PolAssem}$ can be measured by determining Q1 in a competing system (Figure 1, A and D). Equivalently, the CC has been defined as the concentration of free subunits left in solution once polymer assembly has reached a steady-state level in such a competing system ($CC_{SubSoln}$, measured by Q2 in Figure 1, A and D). This set of ideas is based on early empirical observations with actin (Oosawa *et al.*, 1959). These observations were initially given a theoretical framework by Oosawa and colleagues, who explained the behavior of actin by developing a theory for the equilibrium assembly behavior of helical polymers (Oosawa and Kasai, 1962; Oosawa, 1970). This equilibrium theory was extended to tubulin by Johnson and Borisy (1975).

The idea that polymer assembly commences at the CC is now used routinely to design and interpret experiments involving cytoskeletal polymers (e.g., Amayed *et al.*, 2002; Buey *et al.*, 2005; Wieczorek *et al.*, 2015; Concha-Marambio *et al.*, 2017; Díaz-Celis *et al.*, 2017; Schummel *et al.*, 2017), and it is a standard topic in cell biology textbooks (e.g., Alberts *et al.*, 2015; Lodish *et al.*, 2016). Over time, a set of experimental measurements and definitions of CC have emerged (Table 1 and Figure 1), all of which would be equivalent for an equilibrium polymer, that is, a polymer that reaches maximum assembly at thermodynamic equilibrium. In other words, for an equilibrium polymer, there is one CC that can be measured in multiple ways.

Nucleotide hydrolysis allows microtubules to exhibit dynamic instability

MTs (composed of subunits called tubulin dimers) are steady-state polymers, not equilibrium polymers, because they require a constant input of energy in the form of GTP (guanosine triphosphate) nucleotides to maintain a (highly) polymerized state. MTs exhibit a behavior known as DI, in which they stochastically switch between phases of growth and shortening via transitions known as catastrophe and rescue (Figure 1E) (Mitchison and Kirschner, 1984a; Walker *et al.*, 1988). The DI behavior of MTs is driven by GTP hydrolysis (conversion of GTP-tubulin to GDP-tubulin): tubulin subunits containing GTP assemble into MTs, while tubulin subunits containing GDP do not (this occurs because the k_{on} and k_{off} values for GTP-tubulin differ from those for GDP-tubulin). In contrast, tubulin subunits containing non- or slowly hydrolyzable

GTP analogues (e.g., GMPCPP) assemble into stable MTs that do not display DI (Hyman *et al.*, 1992). Though some details about the mechanism of DI remain unclear, the consensus explanation for DI behavior is that growing MTs have a cap of GTP-tubulin subunits (the "GTP cap") that stabilizes the underlying GDP-tubulin lattice. The MTs switch to rapid disassembly (i.e., undergo catastrophe) when they lose their stabilizing caps, exposing the unstable GDP-tubulin lattice below. When MTs regain their caps, they undergo rescue (transition from shortening to growth) (reviewed in Goodson and Jonasson, 2018). In contrast to equilibrium polymers, where individual filaments behave similarly to their populations, individual MTs and their populations can behave differently; for example, individual MTs can have shortening phases even when the overall polymer mass of the population is increasing.

Problems with applying equilibrium polymer theory to dynamically unstable polymers

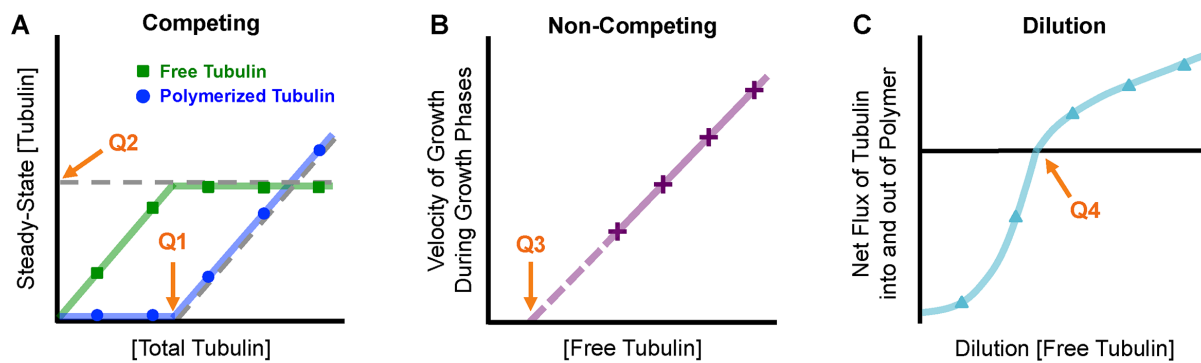
On the surface, it may seem reasonable to apply the traditional CC framework as outlined above (see also Table 1) to understanding DI polymers such as MTs, because this framework is founded on theory (albeit equilibrium polymer theory) and appears to be consistent with many experimental results (Howard, 2001). However, further consideration reveals problems with this approach.

For a theoretical equilibrium polymer, the CC has been defined as $k_{off}/k_{on} = K_D$ (CC_{KD} , Table 1), where k_{on} and k_{off} are the rate constants for attachment/detachment of a subunit to/from a filament tip and K_D is the equilibrium dissociation constant for addition of subunits to polymer (e.g., Oosawa and Asakura, 1975; Howard, 2001; see also Pollard, 2010, for general information about K_D for bimolecular reactions). Thus, polymer will undergo net assembly when [free subunit] is greater than CC_{KD} , because the rate of attachment ($k_{on} \times [\text{free subunit}]$) will be greater than the rate of detachment (k_{off}). Though this is frequently stated in textbooks, it is well recognized that the idea that "the CC" is the K_D is a serious oversimplification when applied to MTs, or to steady-state polymers more generally (Alberts *et al.*, 2015). More specifically, experimentally observed CCs for systems of dynamic MTs (however measured) cannot be equated to simple $k_{off}/k_{on} = K_D$ values because the GTP and GDP forms of tubulin have significantly different values of k_{off}/k_{on} . For example, the CC for GMPCPP (GTP-like) tubulin has been reported to be less than 1 μM (Hyman *et al.*, 1992), while that for GDP-tubulin is very high, perhaps immeasurably so (Howard, 2001).

Exactly how the measured CC value(s) for a system of dynamic MTs relate to the K_D values for GTP- and GDP-tubulin has not been established. However, intuition suggests that any CCs must lie between the respective K_D values for GTP- and GDP-tubulin (Howard, 2001). Consistent with this idea, experimentally reported values for mammalian brain tubulin CC typically lie between ~ 1 and $\sim 20 \mu\text{M}$ (e.g., Verdier-Pinard *et al.*, 2000; Bonfils *et al.*, 2007; Mirigian *et al.*, 2013; Wieczorek *et al.*, 2015).

Note that although the idea that $CC = K_D$ cannot apply in a simple way to a system of dynamic MTs, it can apply to tubulin polymers in the absence of hydrolysis, where assembly is an equilibrium phenomenon. Examples include systems containing only GDP-tubulin (when polymerized with certain drugs) or tubulin bound to non-/slowly hydrolyzable GTP analogues (e.g., GTP- γS , GMPCPP) (Hyman *et al.*, 1992; Díaz *et al.*, 1993; Buey *et al.*, 2005). However, even for an equilibrium polymer, the K_D might not

Traditional Measurements of Critical Concentration



D	Experimentally Measureable Quantity	Traditional Measurement Method (square brackets [] represent concentration)	Theoretical Critical Concentration corresponding to Q
Q1	Determined by measuring steady-state [polymerized tubulin] at different [total tubulin] in a competing (closed) system and extrapolating back to [polymerized tubulin] = 0.		CC _{PolAssem}
Q2	Plateau reached by steady-state [free tubulin] as [total tubulin] is increased in a competing system.		CC _{SubSoln}
Q3	[Free tubulin] at which $V_g = 0$ (where V_g is the growth velocity during growth phases of individual MTs), measured by plotting V_g as a function of [free tubulin] and extrapolating back to $V_g = 0$.		CC _{Elongation}
Q4	[Free tubulin] at which the flux (measured as the rate of change in [polymerized tubulin]) equals zero in a dilution experiment.		CC _{Flux}

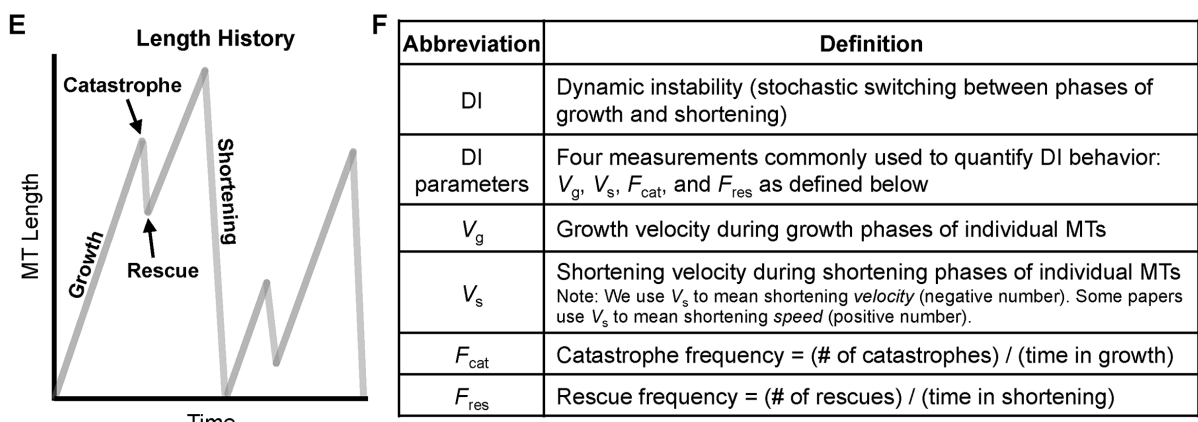


FIGURE 1: Classical understanding of MT polymer assembly behavior. See Table 1 for additional description of the CC measurements depicted here. [Free tubulin] is the concentration of tubulin dimers in solution, [polymerized tubulin] is the concentration of tubulin dimers in polymerized form, and [total tubulin] = [free tubulin] + [polymerized tubulin]. (A) In a competing (closed) system, [total tubulin] is held constant over time and MTs compete for tubulin. As typically presented in textbooks, the CC can be measured in a competing system by observing either the concentration of total tubulin at which MT polymer appears (Q1) or the concentration of free tubulin left in solution once the amount of polymer has reached steady state (Q2). (B) In a noncompeting (open) system, [free tubulin] is held constant over time. In such a system, CC is considered to be the minimum concentration of tubulin necessary for MT polymers to grow, which is estimated by measuring the growth rate of individual filaments (V_g) and extrapolating back to $V_g = 0$ (Q3). (C) In dilution experiments, MTs are grown under competing conditions until the system reaches polymer-mass steady state and then diluted into various [free tubulin]. The initial rate of change in [polymerized tubulin] is measured. Here, CC is the concentration of dilution [free tubulin] at which the rate of change in [polymerized tubulin] is zero (i.e., the dilution [free tubulin] at which the net flux of tubulin into and out of MT polymer is zero) (Q4). (D) Summary table of the definitions of the experimentally measurable quantities Q1–Q4 depicted in panels A–C. (E) Individual MTs exhibit a behavior called dynamic instability (DI), in which the individuals undergo phases of growth and shortening separated by approximately random transitions termed catastrophe and rescue. (F) Table of definitions of DI parameters (four measurements commonly used to quantify DI behavior).

Classical CC definition	Abbreviation	Experimental measurement of CC as applied to MT systems
Minimal concentration of <i>total</i> subunits (e.g., tubulin dimers) necessary for polymer assembly (Oosawa, 1970; Johnson and Borisy, 1975)	CC _{PolAssem}	CC _{PolAssem} is determined by measuring steady-state [polymerized tubulin] at different [total tubulin] in a competing system and extrapolating back to [polymerized tubulin] = 0. See Q1 in Figure 1A (also Figures 3, A and B, and 4).
Concentration of <i>free</i> subunits left in solution once equilibrium or steady-state assembly has been achieved (Oosawa, 1970; Johnson and Borisy, 1975) ^a	CC _{SubSoln}	CC _{SubSoln} is determined by measuring [free tubulin] left in solution at steady state for different [total tubulin] in a competing system and determining the position of the plateau reached by [free tubulin]. See Q2 in Figure 1A (also Figures 3, A and B, and 4).
Dissociation equilibrium constant for the binding of subunit to polymer, i.e., $CC = K_D = k_{off}/k_{on}$ (Oosawa and Asakura, 1975) ^b	CC _{KD}	CC _{KD} can be determined by separate experimental measurement of k_{on} and k_{off} for addition/loss of tubulin subunits to/from MT polymer, respectively, and then calculating the ratio k_{off}/k_{on} .
Concentration of <i>free</i> subunit at which the rate of association equals the rate of dissociation during the elongation phase (called S_c^e in Walker <i>et al.</i> , 1988; similar to c_1 in Hill and Chen, 1984) ^c	CC _{Elongation}	CC _{Elongation} is determined by measuring the growth rate during the growth state (V_g) at various values of [free tubulin] and extrapolating back to the [free tubulin] at which $V_g = 0$. See Q3 in Figure 1B (also Figure 7, A and B).
Concentration of <i>free</i> subunit at which the fluxes of subunits into and out of polymer are balanced, that is, where the net flux is zero (called c_0 in Hill and Chen, 1984)	CC _{Flux}	CC _{Flux} is determined by growing MTs to steady state at very high [total tubulin], then rapidly diluting to a new [free tubulin] and measuring the initial rate of change in [polymerized tubulin] (i.e., measuring [polymerized tubulin] flux). CC _{Flux} is the value of [free tubulin] where [polymerized tubulin] flux = 0. See Q4 in Figure 1C (also Figure 6).
Concentration of <i>free</i> subunit at which polymers transition from "bounded growth" to "unbounded growth" (called c_{cr} in Dogterom and Leibler, 1993)	CC _{Unbounded}	CC _{Unbounded} is the [free tubulin] at which the rate of change in average MT length transitions from equaling zero to being positive (Q5 in Figure 5). CC _{Unbounded} can be identified by measuring DI parameters from MT length histories (Figure 1, E and F) across a range of different [free tubulin] and determining the [free tubulin] at which $V_g F_{res} = IV_s F_{cat}$.

These definitions of CC are interchangeable for equilibrium polymers, but have not all been compared in a single analysis for DI polymers. For each CC definition, we have assigned a specific abbreviation and provided an example of an early publication where that definition was used. The terms CC_{PolAssem}, CC_{SubSoln}, etc., refer to theoretical values (concepts), and Q1, Q2, etc., refer to experimentally measurable quantities (i.e., values obtained through experimental approaches as indicated in the figures). All definitions except CC_{KD} can be applied to both equilibrium and steady-state polymers (CC_{KD} assumes that the system is at equilibrium and therefore can be applied to only equilibrium polymers). The traditional framework outlined here will be revised in *Results*, where we will show that these definitions are not all equivalent for DI polymers (see Tables 3 and 4 for a summary).

^aAssuming that assembly starts from a state with no polymer, maximal polymer assembly will occur at equilibrium for equilibrium polymers, and at polymer-mass steady state for steady-state polymers. Steady-state polymers will be (mostly) disassembled at thermodynamic equilibrium because the nucleotides in the system will be (effectively) entirely hydrolyzed.

^bThe idea that $CC = K_D$ for simple equilibrium polymers is derived as follows. The net rate of polymer length change at a single filament tip = rate of addition – rate of loss. The rate of addition is assumed to be k_{on} [free subunit], and the rate of loss is assumed to be k_{off} . Therefore, the rate at which new subunits add to a population of n polymers is $n \times k_{on}$ [free subunit], and the rate at which subunits detach from a population of n polymers is $n \times k_{off}$. At equilibrium, rate of polymerization = rate of depolymerization, so $n \times k_{on}$ [free subunit] = $n \times k_{off}$. Therefore, at equilibrium, [free subunit] = $k_{off}/k_{on} = K_D = CC_{KD}$.

^cCC_{Elongation} has been interpreted as the minimal concentration of free subunit needed to elongate from a growing polymer. The derivation of CC_{Elongation} is similar to that for CC_{KD}, but considers the behavior of a single filament, not a population, and can apply to steady-state polymers because it does not require equilibrium. For polymers displaying DI, measurements of CC_{Elongation} are performed during the growth state of DI. The derivation of CC_{Elongation} assumes that V_g is a linear function of [free subunit], i.e., $V_g = k_{on}^{growth}$ [free subunit] – k_{off}^{growth} , where k_{on}^{growth} and k_{off}^{growth} are observed rate constants during growth. Thus, the [free subunit] at which $V_g = 0$ is $k_{off}^{growth}/k_{on}^{growth} = CC_{Elongation}$.

TABLE 1: Traditional critical concentration definitions used in the literature.

equate to a simple k_{off}/k_{on} value (e.g., if the off rate is concentration-dependent [Gardner *et al.*, 2011]).

A related problem with applying equilibrium polymer theory to DI polymers is that it leaves open questions regarding how DI and energy utilization fit into the traditional CC framework. For example, how does the DI behavior of an individual filament in Figure 1, B and E, relate to the population-level behavior in Figure 1, A and C? Is there one experimentally relevant CC (as assumed from equilibrium polymer theory) or more than one? More broadly, why do some steady-state polymers (e.g., MTs) display DI, while others (e.g., actin) do not?

Previous efforts to clarify critical concentration as it applies to steady-state polymers

As one might imagine, some of these questions have been studied previously. However, ambiguity in understanding CC(s) of steady-state polymers still exists, in part because these earlier studies did not clearly relate their results to the classical definitions of CC, and few compared results between competing and noncompeting systems (Tables 1 and 2). A brief summary of some key previous efforts on CC for MTs is as follows (note that these focused on noncompeting systems):

Type of experiment/ simulation	Description
Competing	Closed system where [total tubulin] is held constant for the duration of the experiment and MTs compete for tubulin (e.g., in a test tube)
Noncompeting	Open system where [free tubulin] is held constant for the duration of the experiment (e.g., in a flow cell)
Dilution	System where MTs are grown to polymer-mass steady state under competing conditions at very high [total tubulin] and then moved into noncompeting conditions at various values of [free tubulin]

TABLE 2: Types of experiments/simulations.

1. In the 1980s, Hill and colleagues investigated some of the questions outlined above and worked to develop a theory of steady-state polymer assembly. Their conclusions included the idea that growth of MTs is governed by two distinct critical concentrations: a lower CC, which is the [free subunit] where “the mean subunit flux per polymer” during “phase 1” (the growth phase) equals zero, and an upper CC, which is the [free subunit] where “the mean net subunit flux per polymer” is zero (similar to Figure 1C; e.g., Hill and Chen, 1984, elaborated on in Hill, 1987). However, the published work did not clarify for readers the biological significance of these two CCs nor how they relate to the behaviors of individual filaments and their populations.
2. Later in the 1980s, Walker *et al.* used video microscopy to analyze in detail the behavior of individual MTs undergoing DI. They demonstrated that MTs observed *in vitro* have a “critical concentration for elongation” ($CC_{Elongation}$), which they described as the [free subunit] at which the rate of tubulin association (k_{on}^{growth} [free tubulin]) is equal to the rate of dissociation (k_{off}^{growth}) during the elongation phase (Walker *et al.*, 1988; Figure 1B; Table 1 and its footnotes). Consequently, at tubulin concentrations below $CC_{Elongation}$, there is no elongation. Later in this same paper, the authors discussed the existence of a higher CC above which a population of polymers will undergo “net assembly” (we refer to this as $CC_{NetAssembly}$). Thus, the analysis in this article clearly indicates that MTs have two CCs. However, this conclusion is not stated explicitly, and the article does not address the question of how either of the two Walker *et al.* CCs relates to the two CCs predicted by Hill.
3. In the 1990s, Dogterom *et al.* and Fygenson *et al.* used a combination of modeling (Dogterom and Leibler, 1993) and experiments (Fygenson *et al.*, 1994) to show that there is a “critical value of monomer density, $c = c_{cr}$,” above which MT growth is “unbounded” (i.e., the average length increases indefinitely and does not level off with time; Dogterom and Leibler, 1993; Dogterom *et al.*, 1995; Fygenson *et al.*, 1994). Hereafter, we refer to this c_{cr} as $CC_{Unbounded}$. Dogterom *et al.* also provided equations (similar to those proposed initially by Hill and Chen, 1984, and Walker *et al.*, 1988) that can be used to relate $CC_{Unbounded}$, which is a population-level characteristic, to the DI parameters (Figure 1, E and F), which describe individual-level behavior. One of the many significant outcomes of these papers was that they encouraged readers to think about how small changes to DI parameters (e.g., as caused by regulatory changes to MT binding proteins) could change the behavior of a system of MTs, especially in a cellular context. However, the implications of these articles for understanding CCs more broadly remained poorly appreciated because they did not explicitly relate $CC_{Unbounded}$ to the more classical CC definitions and measurements in Table 1 or to those discussed by Hill and Chen (1984) and Walker *et al.* (1988).

Thus, although DI has been studied for more than 30 years, confusion remains about how the traditionally equivalent definitions of CC and the interpretation of CC measurements should be adjusted to account for DI. Remarkably, the literature as yet still lacks a clear discussion of how the $CC_{Elongation}$ and $CC_{Unbounded}$ mentioned above relate to each other, to the CCs predicted by Hill, or to the classical experimental measurements of CC depicted in Figure 1A. To address this problem, we will investigate the following questions:

1. How many distinct CCs are produced by the different experimentally measurable quantities (Q values, Figure 1 and Table 1), which measurements yield which CC, and what is the practical significance of each?
2. How do these values relate to behaviors at the scales of subunits, individual MTs (e.g., Figure 1, B and E), and the bulk polymer mass of populations of MTs (e.g., Figure 1, A and C)?
3. How does the separation between distinct CCs relate to DI, and can the separation help to explain differences between steady-state polymers that display DI (e.g., tubulin) and those that do not (e.g., actin)?

Undoubtedly, many researchers have an intuitive understanding of the answers to at least some of these questions. However, the observation that even recent literature contains many references to “the” CC for MT assembly (e.g., Alfaro-Aco and Petry, 2015; Wieczorek *et al.*, 2015; Hussmann *et al.*, 2016; Schummel *et al.*, 2017) indicates that this problem deserves attention.

Biological implications of critical concentrations and practical significance for experimental design and interpretation

While these issues are interesting from a basic science perspective, they also have significant practical relevance: proper design and interpretation of experiments that involve perturbing MT dynamics (e.g., characterization of MT-directed drugs or proteins) require an unambiguous understanding of CCs and how they are measured (e.g., Verdier-Pinard *et al.*, 2000; Bonfils *et al.*, 2007; Hussmann *et al.*, 2016; Cytoskeleton Inc., n.d.).

For example, measuring the values of the CCs provides a way to characterize tubulin from different cell types and effects of polymer-binding proteins. The measured CCs serve as reference points separating concentration ranges where a system’s qualitative behaviors differ, analogous to a K_M for an enzyme–substrate system or a K_D for a binding reaction. Correspondingly, the relationship between the subunit concentration set by an experimenter or a cell and the CCs affects various properties of the resulting polymer population, such as the total amount of polymer present, the length distribution of filaments in the population, and the DI behavior of the filaments.

Examples of practical implications include the following:

- 1) Measuring the effect of a polymer-binding protein on the CCs

can be used not only to characterize the protein, but also to provide an intuitive understanding of its effects on polymer assembly over a range of subunit concentrations. 2) As we will return to in the *Discussion*, CCs are of value for understanding how MT behaviors studied *in vitro* pertain to *in vivo* behaviors, and relatedly, for understanding how cells can manipulate CCs for regulatory purposes. 3) Clarifying which CC is obtained from which measurement is necessary for correct interpretation of experiments (e.g., as our results below show, measurements commonly expected to yield the minimum subunit concentration needed for polymer assembly actually yield $CC_{NetAssembly}$). 4) Finally, we propose that the separation between different CCs can be used to understand behavioral differences between MTs and actin.

Additionally, even if measuring CCs is not the goal of an experiment, understanding the CCs is still relevant to experimental design and interpretation, because the relationship between the starting subunit concentration and the CCs affects the resulting polymer behavior and amount of polymer. For example, if the subunit concentration in an experiment is close to a CC, then a small variation in experimental conditions could lead to a drastic change in polymer mass and behavior. More specifically, if the control conditions and test conditions in an experiment with a MT binding protein have slightly different [free tubulin], with one just below $CC_{NetAssembly}$ and the other just above, then the predictable dramatic difference in results could be incorrectly attributed to the MT binding protein. Indeed, we suggest that incomplete understanding of CCs and their impact on designing and interpreting experiments is one reason for variability in the reported effects of MT binding proteins.

Summary of conclusions

Using systems of simulated MTs, we show that classical interpretations of experiments such as those in Figure 1 can be misleading in terms of understanding the behavior of individual MTs. In particular, we use the simulations to illustrate the fact that dynamically unstable polymers such as MTs do have (at least) two major experimentally distinguishable CCs, as originally proposed by Hill and colleagues (summarized in Hill, 1987). We clarify how the CCs relate to behaviors of individual MTs and populations of MTs. At [free tubulin] above the lower CC, growth phases of *individual* filaments can occur transiently, though experimentally observable growth phases may not occur until well above this CC. At [free tubulin] above the higher CC, the polymer mass of a large *population* will increase steadily; individual filaments in the population potentially still exhibit DI, but with net growth over sufficient time. We show that the lower CC corresponds to $CC_{Elongation}$ (Table 1) as measured by Walker *et al.* (1988), which can be described as the free tubulin concentration above which individual MTs can elongate during the growth phase. This CC can be measured by experimental quantity Q3 in Figure 1B. The higher CC corresponds to $CC_{Unbounded}$ (Table 1) as identified by Dogterom *et al.*, that is, the concentration of free tubulin above which “unbounded growth” occurs (Dogterom and Leibler, 1993; Dogterom *et al.*, 1995; Fygenson *et al.*, 1994). This upper CC can be measured by Q1, Q2, and Q4 in Figure 1, A and C. To clearly distinguish these two CCs and avoid confusing either with a situation where a physical boundary is involved, we suggest calling them $CC_{Elongation}$ and $CC_{NetAssembly}$, based on the terminology in Walker *et al.* (1988).

We show that most experiments intended to measure “the CC” actually measure $CC_{NetAssembly}$ (i.e., the higher CC). This conclusion means that “typical” MT DI (where MTs grow and depolymerize back to the seed) is limited to concentrations below what has tradi-

tionally been considered “the” CC needed for polymer assembly (i.e., Q1, Q2 in Figure 1A; Table 1). Related to the discussion above, this provides one example of why clarifying which CC is obtained from which measurement is relevant to experimental biology: if one thinks that an experiment is measuring the minimum concentration needed for polymer assembly, but it is actually measuring the CC for net assembly, then the biological implications of the experiment could be seriously misinterpreted.

Furthermore, we show that in competing systems (i.e., closed systems where MTs compete for a limited total number of tubulin subunits), the concentration of free tubulin at steady state ($[free\ tubulin]_{SteadyState}$) does not equal $CC_{NetAssembly}$, as would be expected from traditional interpretations of classic CC experiments (Figure 1A). Instead, $[free\ tubulin]_{SteadyState}$ asymptotically approaches $CC_{NetAssembly}$ as [total tubulin] increases.

In addition to the experimentally accessible $CC_{Elongation}$ and $CC_{NetAssembly}$, there are two more CCs (perhaps not experimentally accessible) that correspond to the K_D for the GTP and GDP forms of tubulin subunits. We suggest calling these CC_{KD_GTP} and CC_{KD_GDP} , respectively. For an equilibrium polymer system (e.g., one composed of only the GTP form), $CC_{Elongation}$ and $CC_{NetAssembly}$ would be equal to each other and to the relevant CC_{KD} . In addition, we demonstrate that the degree of separation between $CC_{Elongation}$ and $CC_{NetAssembly}$ depends on the GTP hydrolysis rate constant (k_H). We also show that $CC_{Elongation}$ can differ from CC_{KD_GTP} , contrary to previous assumptions that growing MTs always have GTP-tubulin at their tips (topmost subunits; e.g., Bowne-Anderson *et al.*, 2015).

Finally, we demonstrate that DI itself, in combination with detection limitations, can produce results (e.g., lack of detectable growth phases at concentrations near $CC_{Elongation}$, and sigmoidal seed occupancy plots) previously interpreted as evidence that growth from stable seeds requires a nucleation step.

This article focuses on systems composed of a predefined number of MTs with one end free and the other end anchored at a stable nucleation site, such as would exist for MTs growing from centrosomes. In other cases, MTs can have two free ends (plus and minus). For each of $CC_{Elongation}$ and $CC_{NetAssembly}$, the numerical value at the plus end could differ from the value at the minus end (Walker *et al.*, 1988). While our studies focus on MTs, we suggest that these CC definitions and interpretations can apply to steady-state polymers more generally, but are especially significant for those that exhibit DI.

RESULTS

Computational models: simplified model and detailed model

To investigate and clarify the concept of CC as it applies to dynamically unstable polymers, we used computational modeling. Computational models are ideal for addressing this type of problem because the biochemistry of the reactions can be explicitly controlled, and *in silico* experiments can be performed quickly and easily. Furthermore, it is possible to follow the behavior of the system at all relevant scales simultaneously: addition/loss of individual subunits to/from the free end of each filament, DI of individual filaments, and any changes in polymer mass of the population of filaments. In comparison, it is challenging to address these questions using physical systems because experiments have thus far been limited technically to measurements of one (or at most two) of these scales at a time.

In this work, we used both a “simplified” model of MT dynamics, in which MTs are modeled as simple linear polymers (Gregoretti *et al.*, 2006), and a “detailed” model, where MTs are

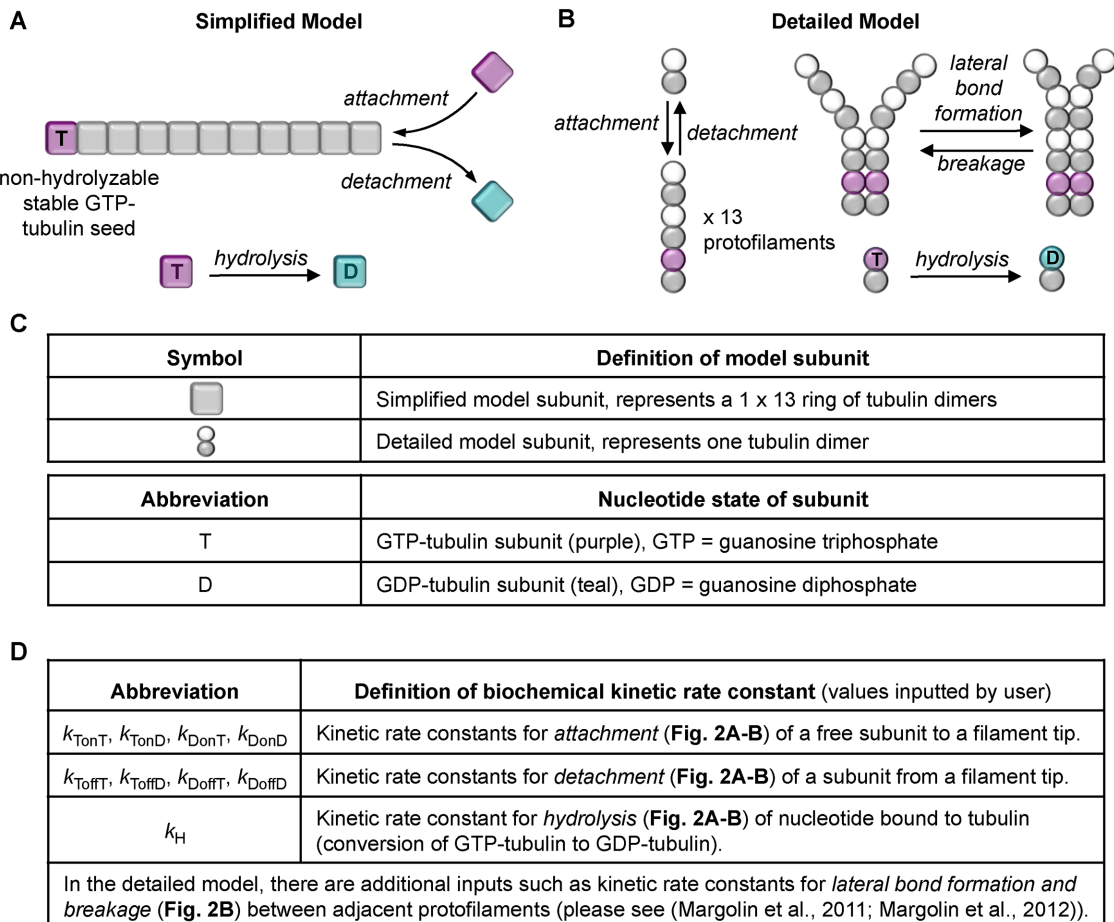


FIGURE 2: Processes that occur in the computational models. (A) In the *simplified model*, MTs are approximated as simple linear filaments that can undergo three processes: subunit addition, loss, and hydrolysis. Addition and loss can occur only at the tip. Hydrolysis can occur anywhere in the filament where there is a GTP subunit. (B) In the *detailed model*, there are 13 protofilaments, each of which undergoes the same processes as in the simplified model but also undergoes lateral bonding and breaking between adjacent protofilaments. (C) Information about the subunits in the models. In both models, the kinetic rate constants (panel D) controlling these processes are input by the user, and the MTs grow off of a user-defined constant number of stable MT seeds (composed of nonhydrolyzable GTP-tubulin). The standard DI parameters (V_g , V_s , F_{cat} , F_{res} ; see Figure 1, E and F) are emergent properties of the input rate constants, [free tubulin], and other aspects of the environment, such as the number of stable seeds. For more information about the models and their parameter sets, see Box 1, *Materials and Methods*, Gregoretti et al. (2006), and Margolin et al. (2011, 2012).

composed of 13 protofilaments (PFs), with lateral and longitudinal bonds between subunits (tubulin dimers) modeled explicitly (Margolin et al., 2011, 2012) (Figure 2). The simulations were designed to be intuitively understandable to researchers familiar with biochemical aspects of cytoskeletal polymers. Consequently, the rules governing the simulations correspond directly to biochemical reaction kinetics. Key elements of these models are described in Box 1.

We utilized both the simplified and detailed computational models because each has particular strengths for addressing problems related to MT dynamics. The simplified model has fewer kinetic parameters, all of which are directly comparable to parameters in typical analytical models (i.e., mathematical equations), and it is similar to single-protofilament models that have been used by other authors (e.g., Padinhateeri et al., 2012; Li and Kolomeisky, 2014; Aparna et al., 2017). Thus, the simplified model is useful for testing

analytical model predictions relating biochemical properties to individual filament level and bulk population level behaviors. In contrast, the increased resolution of the detailed model is important for testing the generality and relevance of conclusions derived from the simplified model.

In addition, the input kinetic rate constants in the two models were tuned to produce dynamic instability behavior that is quantitatively different between the two models (measured DI parameter values are in the Supplemental Excel file). It follows that the specific numerical values for CCs extracted from these two models will be different. However, as demonstrated by the results that follow, the behavioral changes that occur at each CC are qualitatively similar in the two models. Thus, these two models enable us to determine which conclusions are general and to avoid making conclusions that are specific to particular parameter sets or polymer types.

BOX 1: Key elements of the two computational models (simplified and detailed) used in this study.

1. The behaviors of the evolving systems of dynamic MTs in the simulations can be followed at the scales of subunits, individual filaments, or populations of filaments. For both models, the simulations spontaneously undergo the full range of DI behaviors (including rescue), and they can simulate systems of dynamic MTs for hours of simulated time (Gregoretti *et al.*, 2006; Margolin *et al.*, 2012).
2. Subunit addition/loss and GTP hydrolysis (both models) and lateral bond formation/breaking (detailed model only) are modeled as stochastic events that occur according to kinetic rate equations based on the biochemistry of these processes (Figure 2). In the detailed model, longitudinal bonds form first (i.e., subunits attach to a PF), and lateral bonds between a pair of subunits in two neighboring PFs can form only if the pair of subunits immediately below is already laterally bonded. Similarly, lateral bonds between PFs can break only at the interface between bonded and unbonded regions. In other words, lateral bonds between PFs zip open or closed, enabling dynamic cracks to exist between PFs at the MT tip, as was recently observed experimentally (McIntosh *et al.*, 2018). For more information about the effect of the lateral bonds in the simulations, see Margolin *et al.*, 2011, 2012, and Li *et al.*, 2014.
3. The user-defined (adjustable) parameters correspond to the following: the biochemistry of the proteins being studied (i.e., kinetic rate constants for the reactions listed above) and attributes of the environment that would be set by either the experimenter or the cell (e.g., the concentration of tubulin in the system, whether the system is competing (closed) or noncompeting (open), the number of stable seeds, and the system volume).
4. Because MTs in cells and in many *in vitro* experiments grow from stable seeds (nucleation sites such as centrosomes, axonemes, or GMPCPP seeds), our simulations assume that one end of each MT is fixed (as would be the case for growth from centrosomes) and that all addition and loss occur at the free end. In our simulations, the seeds are composed of nonhydrolyzable GTP–tubulin. Except where otherwise noted, the number of stable seeds was set to 100 in the simplified model and 40 in the detailed model.
5. As in physical experiments, emergent properties of the simulated systems include the DI parameters (V_g , V_s , F_{cat} , F_{res} ; see Figure 1, E and F) and the concentrations of free and polymerized tubulin at steady state. In particular, transitions between growth and shortening (catastrophe and rescue) are spontaneous processes that occur when the stabilizing GTP cap happens to be lost or regained as a result of the biochemical reactions described.
6. The kinetic rate constants used as input parameters for the detailed model were previously tuned to approximate the DI parameters of mammalian brain MTs *in vitro* (Margolin *et al.*, 2012). The simplified model parameters used here are modified from those of Gregoretti *et al.* (2006) and were chosen for use here because they produce DI behavior that is quantitatively different from that of the detailed model.

These attributes make these simulations ideal for studying the relationships between the concentration of tubulin, the behaviors of individual MTs, and the behaviors of populations of dynamic MTs. See *Materials and Methods*, Gregoretti *et al.* (2006), and Margolin *et al.* (2012) for additional details including input parameters.

Approach to understanding the relationship between microtubule behaviors and critical concentrations

The term “critical concentration” can have a specific thermodynamic meaning as the solute concentration at which a phase change occurs. Here we use the term operationally, as the concentration at which a behavioral change occurs. To clarify the concept of CC as it applies to MTs, we examined which of the commonly used CC definitions (outlined in Table 1) are meaningful in studying MTs, and for the set that are meaningful, which are equivalent. We determined how the various CC definitions in Table 1 relate to each other and to DI.

To perform this work, we used the simulations to examine simultaneously the behaviors of individual MTs and populations of MTs. More specifically, we ran sets of simulations for both the simplified and detailed models at various tubulin concentrations in both competing systems (closed systems with constant [total tubulin], as might happen in a test tube) and noncompeting systems (open systems with constant [free tubulin], similar to what might happen in a microscope flow cell). This approach mimics various experiments (Table 2) that are classically used to measure MT CC (Table 1). We then assessed and compared the behaviors of the individual MTs (e.g., DI parameters), population-level properties (e.g., [free tubulin] at steady state), and CCs as determined by the traditional definitions (Table 1).

For the work presented here, it is important to recognize that the relevant observations are the *behaviors* of the systems at different scales and the concurrence (or disagreement) between the values of CC that result from various definitions or measurement

approaches; the specific numerical CC values observed are simply outcomes of the particular input kinetic rate constants used and so are not by themselves significant. This situation is analogous to physical MTs, where DI parameters and CC values depend on the protein sequences, temperatures, and buffer conditions used (e.g., Williams *et al.*, 1985; Gildersleeve *et al.*, 1992; Fygenson *et al.*, 1994; Hussmann *et al.*, 2016; Schummel *et al.*, 2017).

We use the terms Q1, Q2, etc., to refer to specific experimentally measurable quantities (i.e., values obtained through experimental approaches, as indicated in the figures), and the terms CC_{KD}, CC_{PolAssem}, CC_{SubSoln}, etc., to refer to theoretical values (concepts) that may or may not correspond to particular experimentally measurable quantities and may or may not be equivalent. Table 1 summarizes traditional CC definitions and measurements used in the literature. Tables 3 and 4 summarize our clarifications of CC definitions and additional Q value measurements based on the results that will be presented in this work.

The results of our *in silico* experiments are presented in the following order:

1. We first present experiments with the computational simulations under competing conditions to assess the classical CC measurements.
2. Next, we present experiments with the simulations under noncompeting conditions to assess DI behavior relative to the CCs and to compare the results with those of the competing experiments.

- Once we have established which CC is provided by each measurement and the behavioral significance of the CCs, we then present simulations in which we alter k_H to investigate the relationship between k_H , the CCs, and DI behavior.
- At the end, we examine how our improved understanding of the relationship between CCs and DI can help explain experimental observations that growth of MTs from stable seeds appears to require a nucleation step.

Dynamically unstable polymers grow at concentrations below standard experimental quantities commonly thought to measure the critical concentration for polymer assembly

A typical way to measure “the CC” for MT assembly is to determine the [total tubulin] at which polymer assembles in a competing (closed) experiment such as that portrayed in Figure 1A, where Q1 measures what is traditionally considered to be the CC for polymer assembly ($CC_{PolAssem}$; e.g., Mirigian et al., 2013). An alternative approach treated as equivalent is to measure the concentration of free tubulin left in solution once steady-state polymer assembly has occurred (Figure 1A, Q2), traditionally considered to yield $CC_{SubSoln}$ (Mirigian et al., 2013). In other words, the expectation is that $Q1 \approx Q2$, and that these experimentally obtained quantities provide equivalent ways to measure the CC for polymer assembly, where $CC_{PolAssem} = CC_{SubSoln}$ (Table 1).

We tested these predictions by performing simulations of competing systems where individual MTs growing from stable seeds compete for a limited pool of tubulin (i.e., [total tubulin] is constant). This situation is analogous to a test-tube experiment in which MTs grow from preformed MT seeds, and both [polymerized tubulin] and [free tubulin] are measured after the system has reached polymer-mass steady state (Supplemental Figure S1, A–D).¹ Initial inspection of Figure 3, A and B shows that the [polymerized tubulin] and [free tubulin] curves are roughly similar to the expectations from the common understanding depicted in Figure 1A, and $Q1 \approx Q2$ (Figure 3, A and B).

However, closer examination of these data shows a key difference from the common understanding (Figure 1A): there is no sharp transition at either Q1 or Q2 (Figure 3, A and B). Significantly, small but nonzero amounts of polymer exist at [total tubulin] below reasonable estimates for Q1 (Figure 3, A and B; Supplemental Figure S1, E and F). In addition, the steady-state concentration of free tubulin ($[free\ tubulin]_{SteadyState}$) is not constant with respect to [total tubulin] for $[total\ tubulin] > Q1$. Instead, $[free\ tubulin]_{SteadyState}$ approaches an asymptote represented by Q2 (Figure 3, A and B). Nonetheless, Q1 is still approximately equal to Q2.² Consistent with these observations, examination of individual MTs in these simulations shows MTs growing and exhibiting DI at [total tubulin] below $Q1 \approx Q2$ (Figure 3, C and D; compare with Figure 3, A and B).

¹Polymer-mass steady state describes a situation where the polymer mass has reached a plateau and no longer changes with time (other than small fluctuations around the steady-state value; Supplemental Figure S1, A–D). Systems of dynamic MTs can also have other steady states (e.g., polymer-length steady state; see also Mourão et al., 2011).

²Because the transitions are not sharp, it can be difficult to determine the exact values of Q1 and Q2. Depending on how the measurements are performed, the values of Q1 and Q2 might appear different from each other. However, $Q1 = Q2$ does hold if the measurements are performed as follows: Q2 is the value of the horizontal asymptote that $[free\ tubulin]_{SteadyState}$ approaches as [total tubulin] increases; Q1 is the $[polymerized\ tubulin] = 0$ intercept of the line with slope 1 that [polymerized tubulin] approaches as [total tubulin] increases (Figure 3, A and B). Note that Q1 would be exactly equal to Q2 in a system with no measurement error, no noise, and no nonfunctional tubulin, but for a physical experiment these factors can interfere with the measurements.

These data (Figure 3) suggest that one of the most commonly accepted predictions of traditional CC understanding is invalid when applied to systems of dynamic MTs: instead of both Q1 and Q2 providing an experimental measure of the minimum concentration of tubulin needed for polymer assembly ($CC_{PolAssem}$), neither does, since MTs exhibiting DI appear at concentrations below $Q1 \approx Q2$. Correspondingly, the results in Figure 3, A and B, indicate that the CC called $CC_{SubSoln}$ would be more accurately defined as the asymptote approached by the $[free\ tubulin]_{SteadyState}$ as [total tubulin] increases, not the value of $[free\ tubulin]_{SteadyState}$ itself (Figure 1A).

The number of stable MT seeds impacts the sharpness of the transition at Q1 and Q2. Why is the transition at Q1 and Q2 in Figure 3, A and B, more gradual than the theoretical transition as depicted in Figure 1A? Previous results of our simplified model (Gregoretti et al., 2006) and other models (e.g., Vorobjev and Maly, 2008; Mourão et al., 2011) indicate that $[free\ tubulin]_{SteadyState}$ depends on the number of stable MT seeds. Therefore, we investigated how changing the number of stable MT seeds affects the shape of the curves in classical CC plots. Examination of the results (Figure 4, A and B; zoom-ins in Figure 4, C and D) shows that changing the number of MT seeds does change the sharpness of the transitions at Q1 and Q2. More specifically, when the number of MT seeds is small, a relatively sharp transition is seen at both Q1 and Q2 in graphs of steady-state [free tubulin] and [polymerized tubulin]; little if any bulk polymer is observed at [total tubulin] below Q1 (Figure 4, fewer seeds, darkest curves, similar to Figure 1A). In contrast, when the number of MT seeds is large, measurable amounts of polymer appear at concentrations well below Q1, and consequently $[free\ tubulin]_{SteadyState}$ approaches the Q2 asymptote more gradually (Figure 4, more seeds, lightest curves). Moreover, the data for various numbers of seeds all approach the same asymptotes (gray dashed lines, Figure 4). These observations indicate that the number of MT seeds does not impact the value of $Q1 \approx Q2$, but does affect how sharply steady-state [free tubulin] approaches the Q2 asymptote.

The observations thus far raise a question: because $CC_{SubSoln}$ is not the minimum tubulin concentration needed for polymer assembly ($CC_{PolAssem}$), what is the significance of $Q1 \approx Q2 \approx CC_{SubSoln}$ for MT behavior?

A critical concentration for net growth of individual microtubules and persistent growth of a population’s polymer mass ($CC_{NetAssembly}$)

To investigate the significance of Q2 (i.e., the asymptote approached by $[free\ tubulin]_{SteadyState}$ as [total tubulin] is increased; Figures 3, A and B, and 4), we examined the dependence of MT behavior on the concentration of free tubulin in noncompeting simulations. For these studies, we fixed [free tubulin] at various values instead of allowing polymer growth to deplete the free tubulin over time. This set of conditions is analogous to a laboratory experiment involving MTs polymerizing from stable seeds in a constantly replenishing pool of free tubulin at a known concentration, such as might exist in a flow cell.

As described above, Q1 and Q2 from competing systems do not yield the CC for polymer assembly ($CC_{PolAssem}$) as expected from traditional understanding. Instead, comparison with the noncompeting simulations (Figure 5) shows that Q1 and Q2 correspond to a different CC, which can be described as the [free tubulin] above which individual MTs will exhibit net growth over long periods of time (Figure 5, A and B). Equivalently, this CC can be described as the [free tubulin] above which the polymer mass of a large population of MTs will grow persistently (Figure 5, C and D); we use this

Competing Simulations

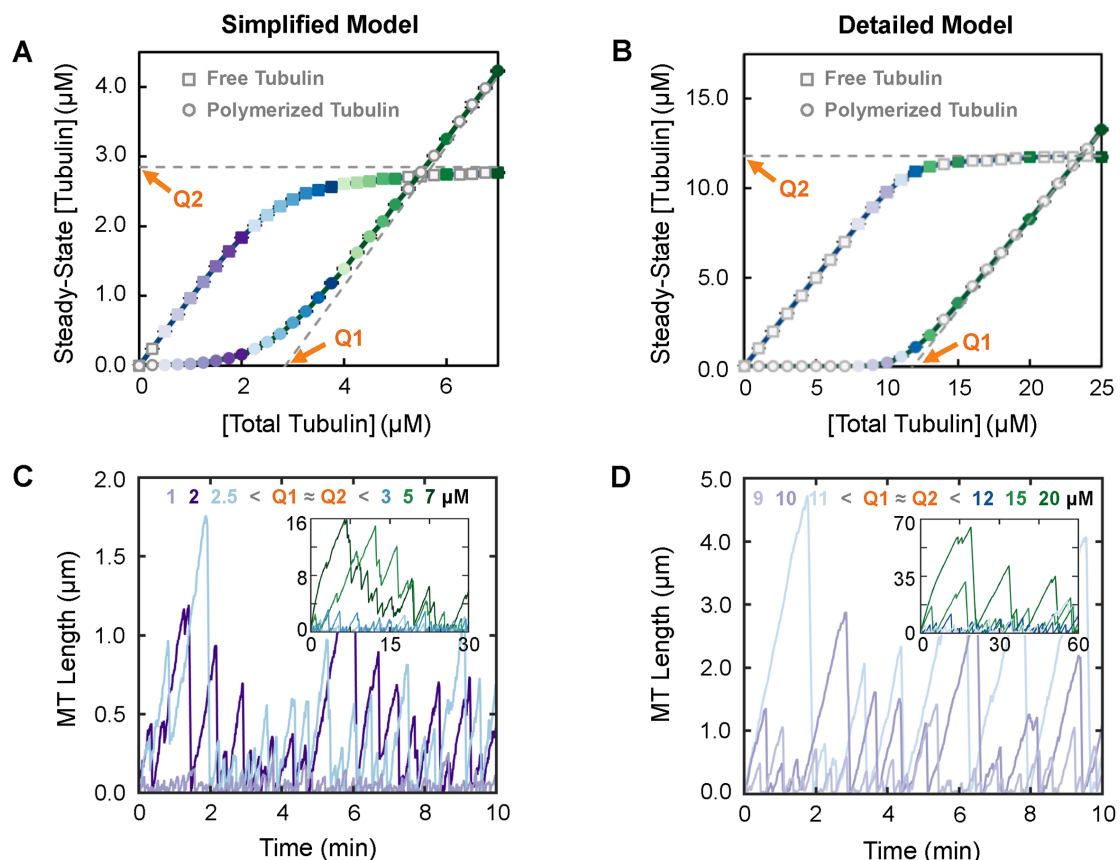


FIGURE 3: Behavior of MTs (populations and individuals) under conditions of constant total tubulin. Left panels: simplified model; right panels: detailed model. Colors of data points reflect the concentrations of total tubulin. (A, B) Classical CC measurements (compare with Figure 1A). Systems of competing MTs at total tubulin concentrations as indicated on the horizontal axes were each allowed to reach polymer-mass steady state (shown in Supplemental Figure S1, A–D). Then the steady-state concentrations of free (squares) and polymerized (circles) tubulin were plotted as functions of [total tubulin]. (C, D) Representative length history plots for individual MTs from the simulation runs used in panels A and B. The value of [total tubulin] for each length history is indicated in the color keys at the top of panels C and D. **Interpretation:** Classically, Q1 estimates $CC_{PolAssem}$, and Q2 estimates $CC_{SubSoln}$. However, as can be seen in panels C and D, MTs grow in both models at [total tubulin] below $Q1 \approx Q2$ ($\sim 2.85 \mu M$ in the simplified model and $\sim 11.8 \mu M$ in the detailed model). Consistent with this observation, the main text provides justification for the idea that CC as estimated by $Q1 \approx Q2$ instead measures $CC_{NetAssembly}$, the CC for persistent growth of a population's polymer mass and net growth of individual MTs over time. Note that the difference in the values of $Q1 \approx Q2$ between the two models is expected from the fact that the input kinetic parameters for the models were chosen to produce quantitatively different DI measurements in order to provide a test of the generality of conclusions about qualitative behaviors; the results show that the behaviors are indeed qualitatively similar between the two models. For additional data related to the competing simulations (e.g., plots of [free tubulin] and [polymerized tubulin] as functions of time), see Supplemental Figure S1. **Methods:** Data points in panels A and B represent the mean \pm one SD of the values obtained in three independent runs of the simulations. The values from each of three runs are averages over 15–30 min for the simplified model (panel 3A) and over 30–60 min for the detailed model (panel 3B). These time periods were chosen so that [free tubulin] and [polymerized tubulin] had reached their steady-state values (Supplemental Figure S1, A–D).

terminology based on the experimentally observed “persistent growth” in Komarova *et al.* (2002).

As discussed more below, this CC is the same as that previously identified by Dogterom *et al.* as the CC at which the transition from “bounded growth” to “unbounded growth” occurs (Dogterom and Leibler, 1993; Dogterom *et al.*, 1995; Fygenson *et al.*, 1994), by Walker *et al.* as the CC for “net assembly” (Walker *et al.*, 1988), and by Hill and Chen (1984) as the CC where net subunit flux equals zero. Note that a “bounded” system refers to one that has a constant steady-state polymer mass or average MT length; “unbounded” refers to a system where the polymer mass or average MT length exhibits net growth over time (Dogterom and Leibler,

1993; Dogterom *et al.*, 1995). This situation should not be confused with one in which the system of MTs experiences a physical boundary (e.g., MTs in cells). To avoid implying that a physical boundary is involved, we suggest using Walker's terminology: CC for net assembly, which we abbreviate $CC_{NetAssembly}$.

$CC_{NetAssembly}$ can be measured by Q5a, the [free tubulin] at which the steady-state net rate of change in average MT length (Figure 5, C and D, left axes) or in polymer mass (Figure 5, C and D, right axes) transitions from zero to positive. Additional approaches to measuring $CC_{NetAssembly}$ are discussed later.

The rate of change in [polymerized tubulin] is also described as the flux of tubulin into and out of polymer, typically abbreviated as

Competing Simulations

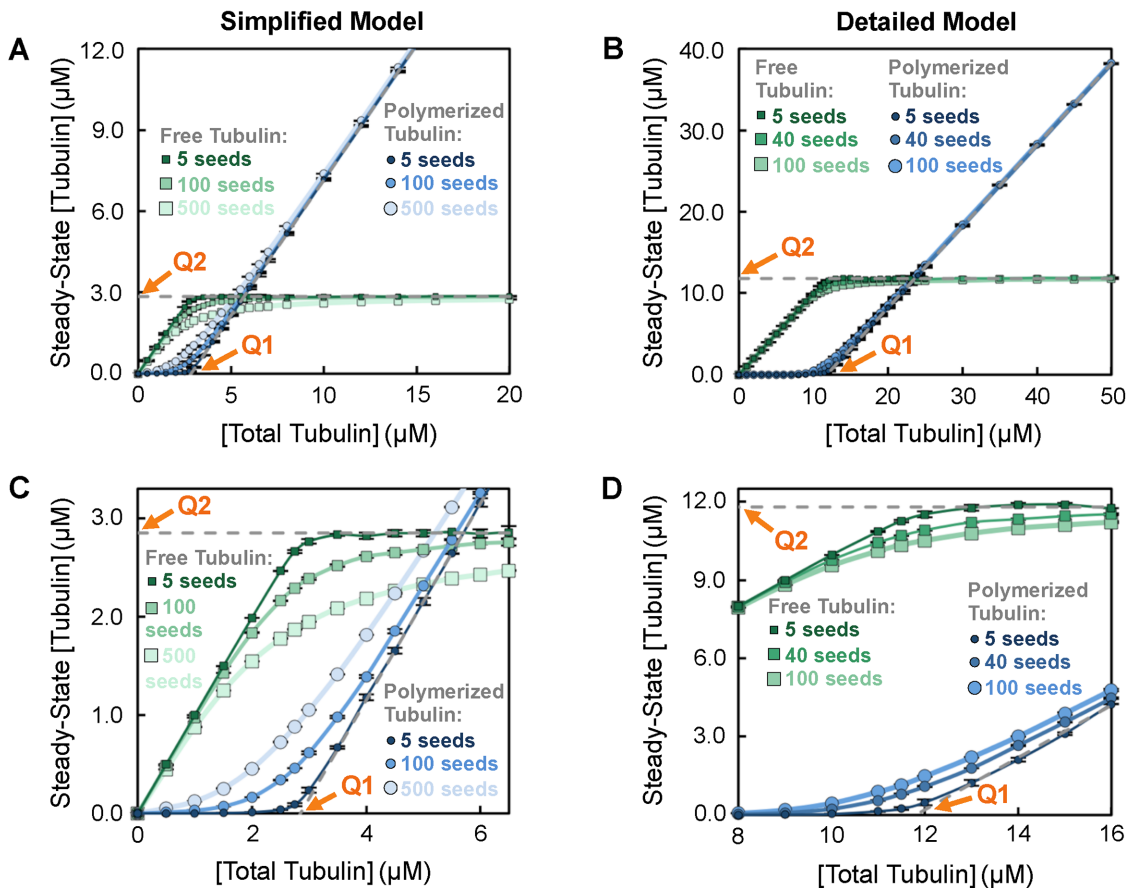


FIGURE 4: Impact of changing the number of MT seeds in competing systems. Steady-state concentrations of free (squares) and polymerized (circles) tubulin in competing systems as in Figure 3, A and B. (A, C) Simplified model with MTs growing from 5, 100, or 500 stable MT seeds (data for 100 seeds replotted from Figure 3A). (B, D) Detailed model with MTs growing from 5, 40, or 100 stable MT seeds (data for 40 seeds replotted from Figure 3B). Panels C and D show zoom-ins on the data plotted in panels A and B, respectively. The darker curves with smaller symbols correspond to fewer seeds and the lighter curves with larger symbols correspond to more seeds. **Interpretation:** These data show that changing the number of stable MT seeds alters the approach to the asymptotes determining Q1 and Q2 (dashed gray lines replotted here from Figure 3, A and B), but does not change the value of $Q1 \approx Q2$. **Methods:** Data points represent the mean \pm one SD of the values obtained in three independent runs of the simulations. Similar to Figure 3, [free tubulin] and [polymerized tubulin] from each run were averaged over a period of time after polymer-mass steady state was reached. The time to reach this steady state depends on the number of stable MT seeds (see Supplemental Figure S2). For the simplified model, the averages of [free tubulin] and [polymerized tubulin] were taken from 120 to 150 min for five MT seeds and from 15 to 30 min for 100 and 500 MT seeds. For the detailed model, the averages were taken from 100 to 150 min for five MT seeds and from 30 to 60 min for 40 and 100 MT seeds. We were able to use a larger number of seeds in the simplified model than in the detailed model because it is more computationally efficient.

J. We will use the abbreviation J_{Constant} for J as measured in constant [free tubulin] experiments, to distinguish it from J as obtained from other experiments discussed below and summarized in Supplemental Table S1. Using this terminology, Q5a is the [free tubulin] at which J_{Constant} transitions from equaling zero to being positive. Note that when the number of individual MTs is constant, the rate of change in the population's [polymerized tubulin] (Figure 5, C and D, right axes) is equivalent to the rate of change in average MT length (Figure 5, C and D, left axes) after converting units and dividing by the number of individual MTs (see also Mauro et al., 2019).

How microtubule behaviors relate to $CC_{\text{NetAssembly}}$. Examination of Figure 5 shows that MT polymerization behavior under noncompeting conditions (i.e., where [free tubulin] is constant) can be divided into two regimes:

Polymer-mass steady state: At concentrations of free tubulin below $CC_{\text{NetAssembly}}$ (measured by Q5a), both average MT length and [polymerized tubulin] within a population reach steady-state values that increase with [free tubulin] but are constant with time ($J_{\text{Constant}} = 0$, Figure 5, C and D; Supplemental Figure S3, A and B). Individual MTs in these systems exhibit what might be called “typical” DI: they undergo periods of growth and shortening, but they eventually and repeatedly depolymerize back to the stable MT seed (Figure 5, A and B).

Polymer-growth steady state: At $CC_{\text{NetAssembly}}$, the populations of dynamic MTs undergo a major change in behavior: they begin to persistently grow in polymer mass. More specifically, when [free tubulin] is above label Q5a in Figure 5, C and D, there is no polymer-mass steady state where [polymerized tubulin] is constant over time (Supplemental Figure S3, A and B). Instead, the system of MTs

Non-Competing Simulations

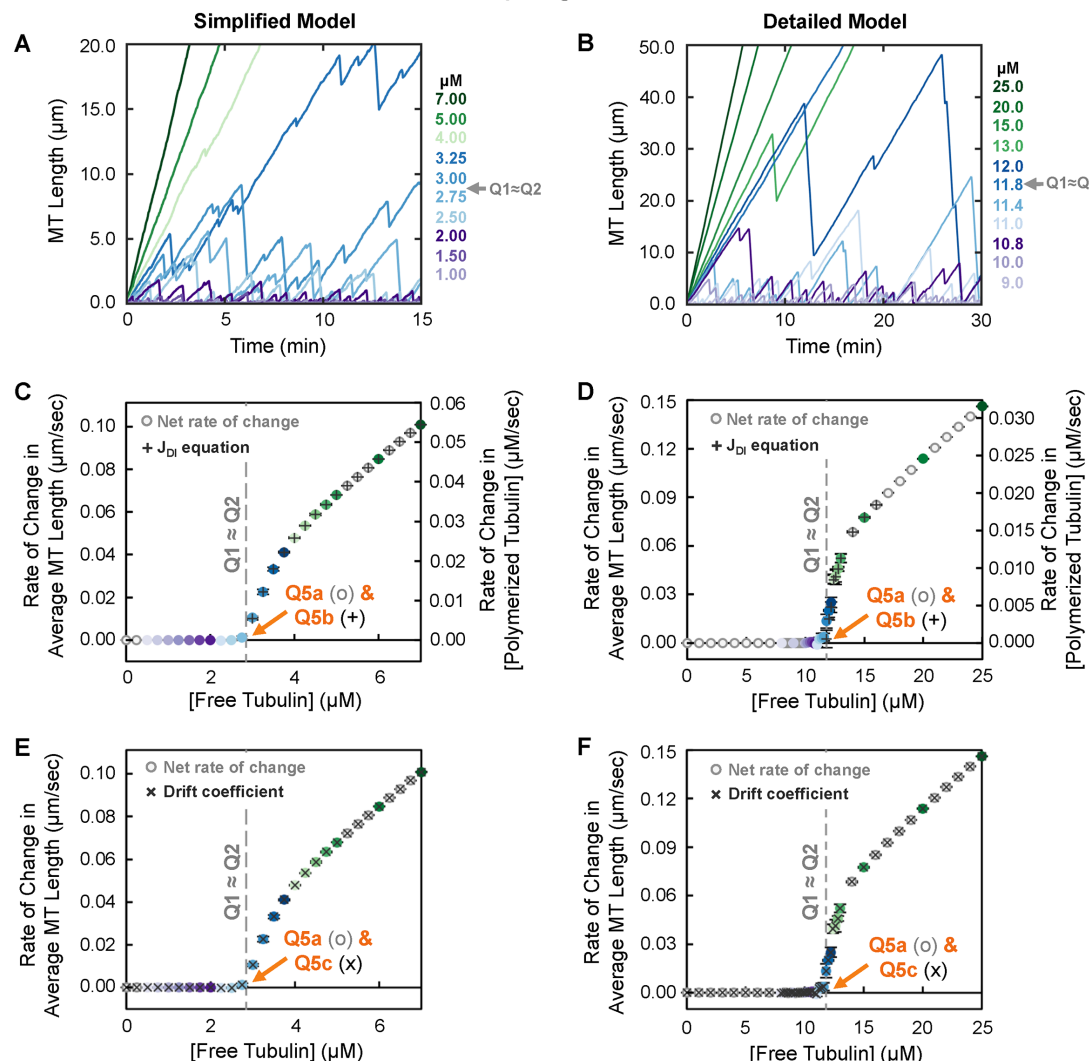


FIGURE 5: Behavior of MTs (individuals and populations) under conditions of constant free tubulin. Left panels: simplified model; right panels: detailed model. Colors of data points reflect the concentrations of free tubulin. (A, B) Representative length history plots for one individual MT at each indicated constant free tubulin concentration. (C, D) Steady-state net rate of change (o symbols) in average MT length (left axes) or in concentration of polymerized tubulin (right axes) for the free tubulin concentrations shown. Q5a indicates the concentration at which this rate becomes positive. This panel also shows the theoretical rate of change in average MT length (+ symbols) as calculated from the extracted DI measurements (Supplemental Excel file) using the equation $J_{DI} = (V_g F_{res} - IV_s F_{cat}) / (F_{cat} + F_{res})$ in the [free tubulin] range where $J_{DI} > 0$ (Eq. 1 in the “unbounded growth” regime) (Hill and Chen, 1984; Walker et al., 1988; Verde et al., 1992; Dogterom and Leibler, 1993). Q5b is the concentration at which J_{DI} becomes positive. (E, F) Drift coefficient (Komarova et al., 2002) of MT populations as a function of [free tubulin] (x symbols). Q5c is the concentration above which drift is positive. For ease of comparison, the rate of change in average MT length (o symbols) from panels C and D is replotted in panels E and F, respectively. For additional data related to these simulations, see Supplemental Figure S3. **Interpretation:** The results show that $Q5a \approx Q5b \approx Q5c$, hereafter referred to as Q5. At concentrations below Q5, populations of MTs reach a polymer-mass steady state where the average MT length is constant over time (the rate of change in average MT length or polymer mass is approximately zero; panels C and D), and the system of MTs exhibits zero drift (panels E and F). At free tubulin concentrations above Q5, populations of MTs reach a polymer-growth steady state where the average MT length and polymer mass increase over time at constant average rates that depend on [free tubulin] (panels C and D), and the system of MTs exhibits positive drift (panels E and F). Plots of average MT length vs. time are shown in Supplemental Figure S3, A and B. Note that the concentration range below Q5 corresponds to the “bounded” regime as discussed by Dogterom et al., while that above Q5 corresponds to the “unbounded” regime (Dogterom and Leibler, 1993). The overall conclusions of the data in this figure are that 1) MTs exhibit net growth (as averaged over time or over individuals in a population) at [free tubulin] above the value Q5 ($Q5a \approx Q5b \approx Q5c$) and 2) Q5 is similar to the value $Q1 \approx Q2$ (gray dashed line) as determined in Figure 3, A and B. Thus, Q1, Q2, and Q5 all provide measurements of the same CC, defined as $CC_{NetAssembly}$ in the main text. **Methods:** All population data points (panels C-F) represent the mean \pm one SD of the values obtained in three independent runs of the simulations. In panels C and D, the net rate of change was calculated from 15 to 30 min. In panels E and F, the drift coefficient was calculated using a method based on Komarova et al. (2002) (Supplemental Methods). See also Supplemental Table S1B for a summary of the measurement methods used in panels C-F.

arrives at a different type of steady state where [polymerized tubulin] increases at a constant rate ($J_{\text{Constant}} > 0$; Figure 5, C and D; Supplemental Figure S3, A and B). Individual MTs within these populations still exhibit DI (except perhaps at very high [free tubulin]), but they exhibit net assembly (Walker et al., 1988) if their behavior is assessed over sufficient time (Figure 5, A and B). This type of behavior is also described as unbounded growth by Dogterom and Leibler (1993).

Significantly, for both models, Q5a (Figure 5, C and D) lies at approximately the value of $Q1 \approx Q2$ (Figure 3, A and B). This observation indicates that $[\text{free tubulin}]_{\text{SteadyState}}$ in competing systems asymptotically approaches the same [free tubulin] at which MTs begin to exhibit net growth (i.e., unbounded growth) in noncompeting systems. In other words, these data show that $CC_{\text{SubSol}} \approx CC_{\text{NetAssembly}}$. This conclusion means that classical methods for measuring "the CC for polymer assembly" do not yield the CC at which individual DI polymers appear, but instead yield the CC above which the polymer mass of a population increases persistently and individual filaments exhibit net growth over sufficient time.

Other experimental methods for measuring $CC_{\text{NetAssembly}}$. As noted above, Dogterom and colleagues previously predicted the existence of a $CC_{\text{Unbounded}}$, the [free tubulin] at which MTs will transition from exhibiting "bounded growth" to exhibiting "unbounded growth" (Dogterom and Leibler, 1993; Dogterom et al., 1995). These growth regimes are characterized by the average MT length reaching a steady-state value (bounded) or increasing indefinitely (unbounded). The predicted existence of a $CC_{\text{Unbounded}}$ was experimentally verified by Fygenson et al. (1994).

An equation for the rate of change in average MT length as a function of the DI parameters had been presented previously (Hill and Chen, 1984; Walker et al., 1988):

$$J_{\text{DI}} = \frac{V_g F_{\text{res}} - |V_s| F_{\text{cat}}}{F_{\text{res}} + F_{\text{cat}}} \quad (1a)$$

Dogterom and colleagues (Verde et al., 1992; Dogterom and Leibler, 1993) then applied this equation to characterize bounded and unbounded growth in constant [free tubulin] systems:

$J_{\text{DI_piecewise}}$ = steady-state rate of change in average MT length

$$= \begin{cases} 0 & \text{during bounded growth} \\ J_{\text{DI}} > 0 & \text{during unbounded growth} \end{cases} \quad (1b)$$

Dogterom et al. identified $CC_{\text{Unbounded}}$ as the [free tubulin] at which $V_g F_{\text{res}} = |V_s| F_{\text{cat}}$ (indicated by the label Q5b in Figure 5, C and D). Significantly, $CC_{\text{Unbounded}}$ as predicted by Q5b from this equation evaluated with our DI parameter measurements matches Q5a (compare + symbols to o symbols in Figure 5, C and D; DI measurements in the Supplemental Excel file). Hence, $CC_{\text{NetAssembly}}$ corresponds to $CC_{\text{Unbounded}}$, and polymer-mass steady state and polymer-growth steady state correspond to "bounded growth" and "unbounded growth," respectively.

Determination of Q5b may not be an experimentally practical way to identify $CC_{\text{NetAssembly}}$, because measuring DI parameters across a range of concentrations requires extended (e.g., >tens of minutes) analysis of many individual MTs, which is laborious and time-consuming. An alternative approach to measuring $CC_{\text{NetAssembly}}$ that may be more tractable experimentally is to use video microscopy to simultaneously analyze the behavior of many individual MTs within a population according to the drift paradigm of Borisy and

colleagues (Vorobjev et al., 1997, 1999; Komarova et al., 2002). The drift coefficient is the mean rate of change in the position of the MT ends (for plus or minus ends separately), also described as the mean velocity of displacement of the MT ends. In cases where one end is fixed, as in our simulations, the drift coefficient is equivalent to the rate of change in average MT length. For a mathematical explanation of how MT behavior can be approximated by a drift-diffusion process, see Maly, 2002; Vorobjev and Maly, 2008; and Mirny and Needleman, 2010.

Here we used a method based on Komarova et al. (2002), which calculates the drift coefficient from the displacements of MT ends over small time steps, for example, between consecutive frames of a movie (see the Supplemental Methods for additional information). As shown in Figure 5, E and F (x symbols), and Supplemental Figure S3, G and H (all symbols), Q5c is the [free tubulin] below which a population of MTs at steady state exhibits zero drift and above which the population exhibits positive drift. Furthermore, zero drift corresponds to the state where the average length of MTs in the population is constant with time, and positive drift corresponds to the average MT length increasing persistently. As one might intuitively predict, $Q5a \approx Q5b \approx Q5c$ (Figure 5, C–F).

The evident similarity between the different measurements in Figure 5, C–F, suggests that the J_{DI} equation using DI parameters (Eq. 1; + symbols in Figure 5, C and D) (Hill and Chen, 1984; Walker et al., 1988; Verde et al., 1992; Dogterom and Leibler, 1993) and the drift equation using short-term displacements (Eq. S1 in the Supplemental Methods; x symbols in Figure 5, E and F) (Komarova et al., 2002) are simply two different representations of the same relationship. Indeed, both yield the rate of change in average MT length as functions of experimentally observed growth and depolymerization behaviors, which we examined in more detail in Mauro et al. (2019).

Measuring $CC_{\text{NetAssembly}}$ using population dilution experiments. Next we tested whether $CC_{\text{NetAssembly}}$ is the same as the CC obtained from the population dilution experiments in early studies of steady-state polymers (e.g., Carlier et al., 1984a,b; see Q4 in Table 1 and Figure 1C). These experiments measure the rate of change in [polymerized tubulin]. As noted above, this rate of change is also described the flux (J) of tubulin into or out of polymer. We refer to this flux measured in dilution experiments as J_{Dilution} to distinguish it from J_{Constant} discussed above, which is determined from constant [free tubulin] experiments. The measurement of J_{Dilution} is performed after a population of MTs at steady state is diluted into a large pool of free tubulin at a new concentration. The measured data from the dilution experiments are then used to produce $J(c)$ plots, where J_{Dilution} is plotted as a function of subunit concentration "c" (Figure 6, A and B). In these plots, "the CC" is identified as the dilution [free tubulin] at which $J_{\text{Dilution}} = 0$ (i.e., where the plotted curve crosses the horizontal axis, Q4). At this concentration, individual MTs undergo periods of growth and shortening, but the population-level fluxes into and out of polymer are balanced (i.e., net growth is zero). We refer to the CC measured via $J(c)$ plots as CC_{Flux} (Table 1). CC_{Flux} corresponds to one of the CCs that was identified by Hill and colleagues, variously named c_o in Hill and Chen (1984) and Chen and Hill (1985b) and a_α in Hill (1987).

Significantly, the value of CC_{Flux} as measured by Q4 in the dilution simulations corresponds to $CC_{\text{NetAssembly}}$ (gray dashed line, Figure 6, A and B) as measured by $Q1 \approx Q2$ in the competing simulations (Figure 3, A and B) and by Q5abc in the noncompeting simulations (Figure 5, C–F). Note also that for (dilution) [free tubulin]

Dilution (and Non-Competing) Simulations

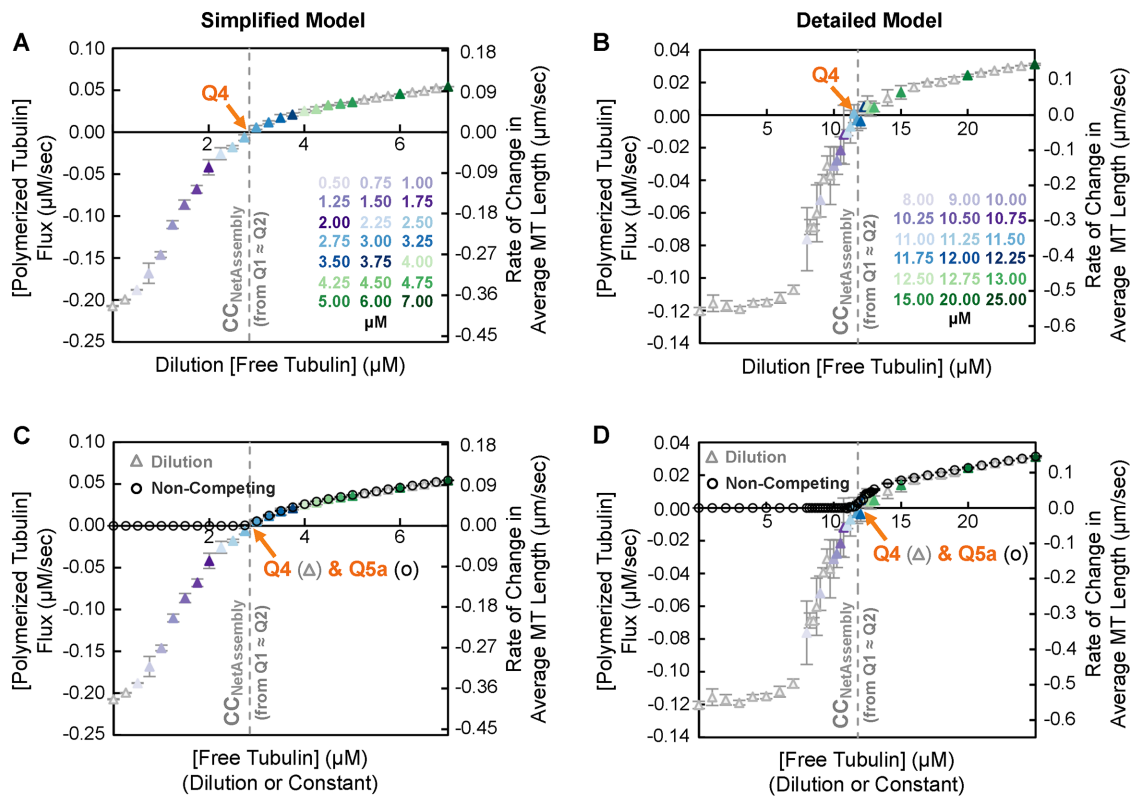


FIGURE 6: Flux of tubulin subunits into and out of MT polymer as a function of dilution [free tubulin] (i.e., a $J(c)$ plot as in Carlier et al., 1984a,b and Figure 1C). Left panels: simplified model; right panels: detailed model. (A, B) In the dilution simulations, competing systems of MTs at high [total tubulin] were allowed to polymerize until they reached polymer-mass steady state. The MTs were then transferred into (“diluted into”) the free tubulin concentrations shown on the horizontal axes. After a brief delay, the initial flux (rate of change in [polymerized tubulin] (left axes) or in average MT length (right axes) was measured. (C, D) Data replotted to show that the $J(c)$ curves from the dilution simulations (i.e., J_{Dilution}) in panels A and B (triangle symbols) and the net rate of change in average MT length from the constant [free tubulin] simulations (i.e., J_{Constant}) in Figure 5, C and D (circle symbols), overlay each other for [free tubulin] above $CC_{\text{NetAssembly}}$. **Interpretation:** These data show that CC as determined by Q4 from $J(c)$ plots is approximately the same value as $Q1 \approx Q2$ (gray dashed line), and thus Q4 also provides a measurement of $CC_{\text{NetAssembly}}$. **Methods:** Competing systems of MTs at 22 μM total tubulin were allowed to reach polymer-mass steady state. Then, at minute 10 of the simulation in the simplified model and at minute 20 of the simulation in the detailed model, the MTs were transferred into the free tubulin concentrations shown on the horizontal axes. After a 5-s delay, the flux was measured over a 10-s period (see Supplemental Figure S4 for plots of [free tubulin] and [polymerized tubulin] as functions of time). Note that the delay after dilution was necessary in the original experiments because of instrument dead time, but it is important for obtaining accurate $J(c)$ measurements because it allows the GTP cap size to respond to the new [free tubulin] (Duellberg et al., 2016; Bowne-Anderson et al., 2013; Mauro et al., 2019). For accurate measurements of J at low values of dilution [free tubulin], the predilution MTs should be sufficiently long so that none completely depolymerize during the 15-s period after dilution (Mauro et al., 2019). Data points for different concentrations of dilution [free tubulin] (see color key) represent the mean \pm one SD of the values obtained in three independent runs of the simulations.

above $CC_{\text{NetAssembly}}$, J_{Dilution} is superimposable with J_{Constant} (Figure 6, C and D). This observation might seem surprising, given the differences in the experimental approaches; however, it makes sense, because in each case the measurement is performed during a time period when [free tubulin] is constant and the rate of change, J , has reached its steady-state value for each [free tubulin] (Supplemental Figures S3, A and B, and S4, C–F). In contrast, J_{Dilution} and J_{Constant} differ from each other below $CC_{\text{NetAssembly}}$. This difference occurs because MTs in dilution experiments are sufficiently long so that they rarely depolymerize back to the MT seed during the measurement period, whereas MTs in constant [free tubulin] below $CC_{\text{NetAssembly}}$ repeatedly depolymerize to the seed. See Mauro

et al. (2019) for a discussion of how the shape of the $J(c)$ curve relates to individual growth and shortening behaviors (e.g., V_g , V_s , fractions of MTs that are growing or shortening).

Thus, all of the experimental approaches for measuring CC discussed thus far yield the CC for persistent growth of a population’s polymer mass and net growth of individual MTs over time ($CC_{\text{NetAssembly}} \approx Q1 \approx Q2 \approx Q4 \approx Q5abc$). This conclusion leaves us with an unresolved question: What is the significance of the remaining common experimental CC measurement, Q3, which is obtained from experiments measuring growth velocity during growth phases for individual MTs as a function of [free tubulin] (see Figure 1B and Table 1)?

A critical concentration for transient elongation phases (growth phases) of individual filaments ($CC_{Elongation}$)

$Q3$ (Figure 1B) has previously been used as a measure of the "critical concentration for elongation" ($CC_{Elongation}$; Walker *et al.*, 1988). According to standard models, $CC_{Elongation}$ is the free subunit concentration where the rate of subunit addition to an individual filament in the growth phase exactly matches the rate of subunit loss from that individual filament, meaning that individual filaments would be expected to grow at subunit concentrations above $Q3 \approx CC_{Elongation}$ (see Table 1 and its footnotes).

To determine the value of $Q3$ in our simulations, we used the standard approach for MTs as outlined in Table 1 (experiments in Walker *et al.*, 1988; see also the theory in Hill and Chen, 1984; Hill, 1987). We plotted the growth velocity (V_g) of individual filaments observed during the growth phase of DI as a function of [free tubulin] and extrapolated a linear fit back to the [free tubulin] at which V_g is zero.³ In addition to performing these measurements on the constant [free tubulin] simulations ($Q3$, Figure 7, A and B), we also used the growth phases that occurred in the dilution experiments to obtain a measurement of $CC_{Elongation}$ ($Q6$ in Figure 7, C and D). Comparing these measurements of $CC_{Elongation}$ in Figure 7, A–D, with the data in Figures 3–6 shows that in both models $CC_{Elongation}$ (as determined by $Q3 \approx Q6$) is well below $CC_{NetAssembly}$ as measured by any of the other approaches ($Q1 \approx Q2 \approx Q4 \approx Q5abc$).

This observation demonstrates that $Q3 \approx Q6$ provides information about MT behavior not provided by the other measurements. Specifically, because $Q3$ and $Q6$ are determined from measurements of the growth velocity of individual MTs during the growth phase of DI, $Q3$ and $Q6$ provide estimates of the [free tubulin] above which *individual filaments can grow transiently* (i.e., can extend during the transient growth phases of DI). Whether growth phases *will* occur also depends on other factors, including the rescue frequency and the frequency of initiating growth from seeds. Additionally, whether growth phases that occur are *observed* depends on length detection thresholds (further discussed below).

Comparison of V_g and J illustrates the relationship between $CC_{Elongation}$, $CC_{NetAssembly}$, and dynamic instability. To relate the observations thus far, note that V_g and $CC_{Elongation}$ come from measurements on individual MTs during only the growth state of DI, whereas J and $CC_{NetAssembly}$ are based on population-level measurements that encompass both growth and shortening phases. At any concentration at which both growth and shortening phases occur, V_g and J will necessarily differ (Figure 8, A and B, shows V_g and J replotted on the same axes). Hence, the difference between V_g and J , and correspondingly the difference between $CC_{Elongation}$ and $CC_{NetAssembly}$, are observable features of polymers that display DI (see also Mauro *et al.*, 2019).

As discussed earlier (Figure 5), at [free tubulin] above $CC_{NetAssembly}$, individual MTs experience net growth over sufficient time. Note that there is a range of [free tubulin] that is above $CC_{NetAssembly}$ but below the point where V_g and J become indistinguishable (Figure 8, A and B). This is the [free tubulin] range where net growth with appreciable DI occurs; in other words, the shortening phases that occur in this range are sufficient to create a

noticeable separation between V_g and J . If shortening were not occurring above $CC_{NetAssembly}$, then V_g and J would be equal for all values of [free tubulin] above $CC_{NetAssembly}$.

Farther above $CC_{NetAssembly}$, V_g and J converge on each other at very high [free tubulin] (Figure 8, A and B). This convergence occurs when [free tubulin] is sufficiently high so that catastrophe is rare and almost all MTs are growing (see the Supplemental Excel file for measured values of F_{cat} , time in growth, and time in shortening). Thus, in this range, measurements of individuals and populations give approximately the same results, leading to additional conclusions with practical significance for measuring the CCs. Specifically, linear extrapolation from $J_{Constant}$ at high [free tubulin] to obtain $Q7$ as shown in Figure 8, C and D, yields approximately the same value for $CC_{Elongation}$ as $Q3 \approx Q6$. Additionally, because $J_{Constant}$ and $J_{Dilution}$ match each other at high [free tubulin] (Figure 6, C and D), the $Q7$ extrapolation can also be performed on the $J_{Dilution}$ data to approximate $CC_{Elongation}$. Thus, both constant [free tubulin] experiments and dilution experiments can be used to obtain not only $CC_{NetAssembly}$ (via $Q4 \approx Q5abc$) but also $CC_{Elongation}$ (via $Q3 \approx Q6 \approx Q7$).

$CC_{Elongation}$ is not $CC_{PolAssem}$. The information above leads to the conclusion that $CC_{Elongation}$ represents a lower limit for individual MTs to exhibit the transient growth phases of DI. One might be tempted to use this idea to predict that $CC_{Elongation}$ is the concentration of free tubulin at which polymer appears (i.e., that $CC_{Elongation} = CC_{PolAssem}$). However, this prediction fails. Contrary to traditional expectations, there is no total or free tubulin concentration at which polymer assembly commences abruptly. Instead, the amount of polymer initially increases in a slow and nonlinear way with respect to [free tubulin], increasing more rapidly only as [free tubulin] approaches $CC_{NetAssembly}$ (Supplemental Figure S3, A–F). The same conclusion is reached whether polymer mass (Supplemental Figure S3, A and B), average MT length (Supplemental Figure S3, A–F), or maximal MT length (Supplemental Figure S3, C–F) is examined.

Additionally, the appearance of detectable polymer mass in a population will depend not only on whether individual MTs are exhibiting growth phases, but also on the number of individual MTs in the population. The results in Figure 4 demonstrate that the number of MT seeds strongly affects the [total tubulin] at which a population's [polymerized tubulin] first becomes noticeably non-zero. If there are few seeds for individual MTs to grow from, the appearance of noticeable [polymerized tubulin] does not occur until [total tubulin] is near or at $CC_{NetAssembly}$. As the number of seeds is increased, [polymerized tubulin] becomes detectable at lower values of [total tubulin] (Figure 4; compare progression from darker to lighter curves).

These observations indicate that MTs (and DI polymers more broadly) do not have a CC for polymer appearance ($CC_{PolAssem}$) as traditionally understood. $CC_{Elongation}$ is the tubulin concentration above which DI growth phases *can* occur, but significant amounts of polymer generally do not accumulate in experiments with bulk polymer until [free tubulin] nears or exceeds $CC_{NetAssembly}$ (Figure 3; Supplemental Figures S1–S3).

These behaviors might seem counterintuitive, but can be explained by the following reasoning. First, when [free tubulin] is just above $CC_{Elongation}$, the growth velocity during the growth phase is low ($V_g = 0$ at $Q3$) and the frequency of catastrophe (F_{cat}) is high (measured DI parameter values in the Supplemental Excel file). Then, under these conditions, individual MTs will be both short (Figure 5, A and B; Supplemental Figure S3, A–F) and short-lived (Figure 5, A and B), and thus difficult to detect. As [free tubulin]

³This V_g versus [free tubulin] relationship is expected to be linear on the basis of the assumption that growth occurs according to the equation $V_g = k_{TonT}$ [free tubulin] – k_{ToffT} , where the first term corresponds to the rate at which GTP-tubulin attaches to a GTP tip, and the second term corresponds to the rate at which GTP-tubulin detaches from a GTP tip (Bowne-Anderson *et al.*, 2015). We return to this relationship later in the main text.

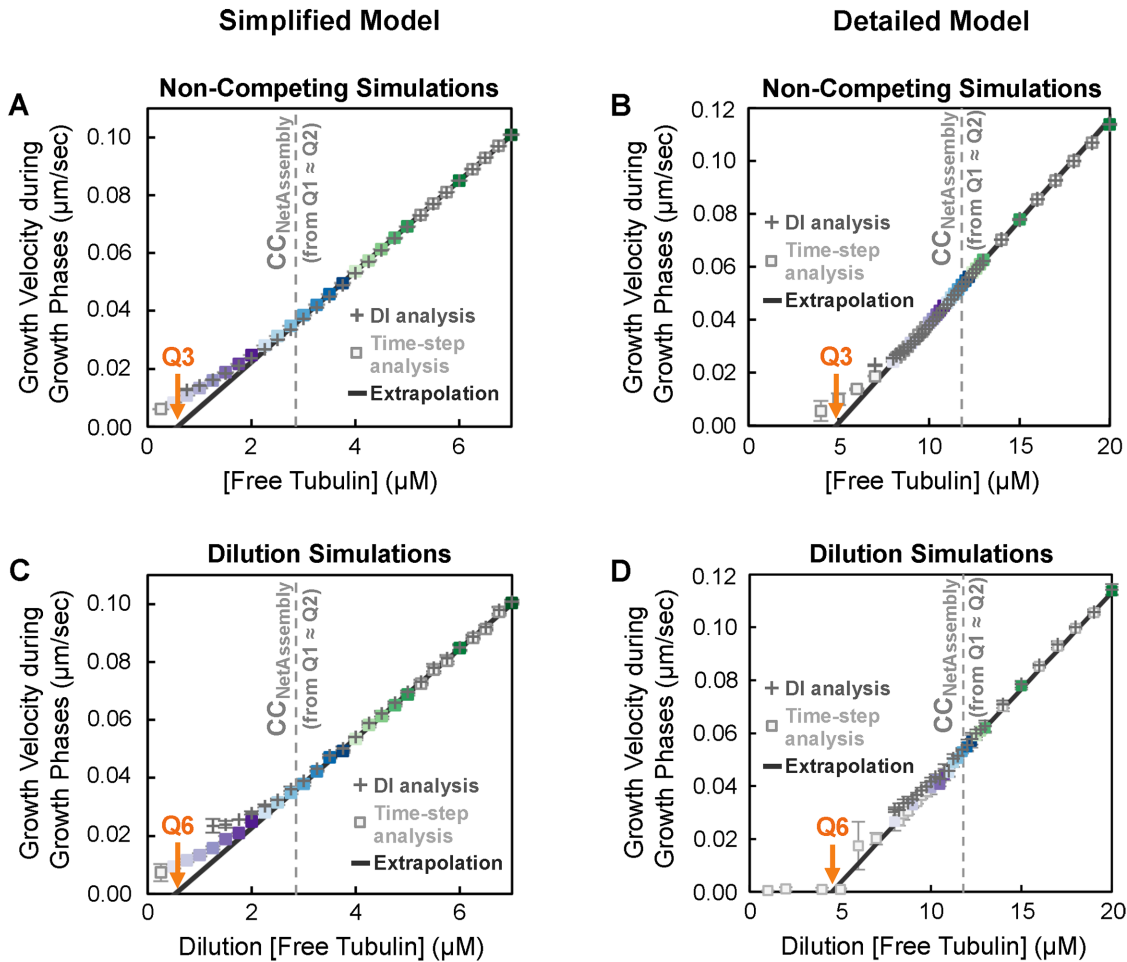


FIGURE 7: Growth velocity of individual MTs during the growth state as a function of [free tubulin]. Left panels: simplified model; right panels: detailed model. Colors of data points reflect the concentrations of free tubulin. (A–D) Growth velocity (V_g) measured using growth phases from either the constant [free tubulin] simulations of Figure 5 (panels 7A and 7B) or the dilution simulations of Figure 6, A and B (panels 7C and 7D). Each of panels A–D shows V_g as measured by a standard DI-based analysis method (+ symbols) and a time-step method (square symbols) inspired by Komarova et al. (2002). Regression lines (solid black) were fitted to the linear range of these data and extrapolated back to $V_g = 0$ to obtain Q3 for the constant [free tubulin] simulations (panels A and B) and Q6 for the dilution simulations (panels C and D). **Interpretation:** These data show that the CC as measured by Q3 is approximately equal to that measured by Q6 and is different from $CC_{NetAssembly}$ (gray dashed line) as measured by $Q1 \approx Q2$ ($\approx Q4 \approx Q5$) from Figures 3–6. The text provides justification for the idea that Q3 and Q6 estimate $CC_{Elongation}$, the CC above which individual filaments can exhibit the transient growth phases of DI. **Methods:** In panels A and B (constant [free tubulin]), the V_g measurements were taken during the time period from minute 15 to minute 30 of the simulations (chosen so that the system has reached either polymer-mass or polymer-growth steady state). In panels C and D (dilution simulations), the V_g data were acquired from 5 to 15 s after the dilution, that is, the J/c measurement period described in Figure 6. For the DI-based analyses (panels A–D, + symbols), we used a custom MATLAB program to identify and quantify growth phases by finding peaks in the length history data. The time-step method (panels A–D, square symbols) divides each length history into 2-s intervals and identifies intervals during which there is a positive change in the MT length. See the Supplemental Methods for more information about both methods. Regression lines were fitted to the time-step measurements of V_g for [free tubulin] in ranges where the V_g data are approximately linear as a function of [free tubulin]: from 3 to 7 μM for the simplified model (panels A and C) and from 7 to 15 μM for the detailed model (panels B and D). All data points represent the mean \pm one SD of the values obtained in three independent runs of the simulations. Data points are shown only at concentrations where at least one detected growth segment occurred in each of the three replicates.

rises, MTs will experience growth phases that last longer (because F_{cat} drops) and also have higher growth velocity (Figure 7). The combined impact of these two effects creates a nonlinear relationship between [free tubulin] and [polymerized tubulin] or equivalently the average MT length observed at steady state; it similarly creates a nonlinear relationship between [free tubulin] and maximal MT length as observed within a period of time (Supplemental Figure S3, C–F).

Measurement of $CC_{Elongation}$ by Q3, Q6, or Q7 is approximate. $CC_{Elongation}$ and $CC_{NetAssembly}$ are intrinsic properties of a system (i.e., a particular protein sequence in a particular buffer or cell type), whereas the experimental measurements (Q values) are subject to measurement error and are therefore approximate. The measurements of $CC_{Elongation}$ by Q3, Q6, or Q7 can be particularly sensitive to measurement error and noise because they are based on extrapolations.

Non-Competing Simulations

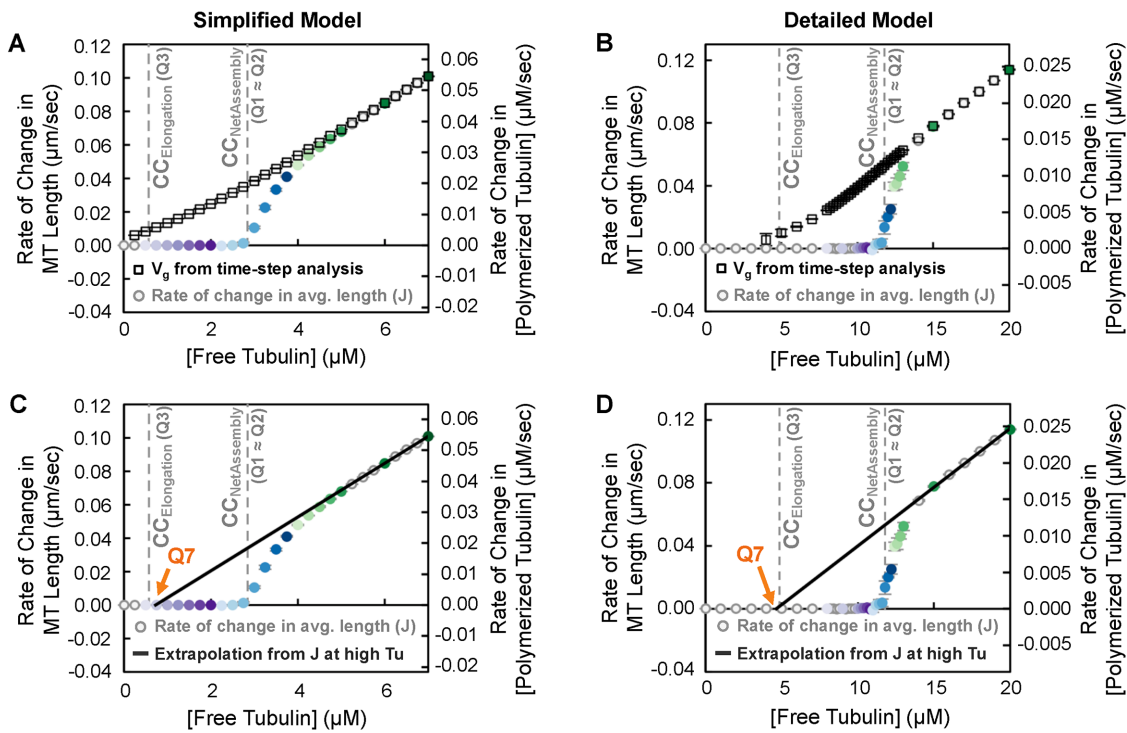


FIGURE 8: Comparison of V_g and J (panels A and B) and an alternative method for measuring $CC_{Elongation}$ (panels C and D). Left panels: simplified model; right panels: detailed model. In all panels, the gray dashed lines represent $CC_{Elongation}$ as measured by Q3 (Figure 7, A and B) and $CC_{NetAssembly}$ as measured by $Q1 \approx Q2$ (Figure 3, A and B). (A, B) Overlay of V_g measured from the time-step analysis of growing individual MTs (square symbols; replotted from Figure 7, A and B) and $J_{Constant}$ measured from the net rate of change in average MT length of the MT population (circle symbols; replotted from Figure 5, C and D), both from the constant [free tubulin] simulations. **Interpretation of panels A, B:** These data show that at high [free tubulin], $J_{Constant}$ approaches the V_g of individual MTs. These two data sets converge because at sufficiently high [free tubulin] individual MTs are growing (nearly) all the time, as seen in the length histories (Figure 5, A and B). Thus, $CC_{Elongation}$, which was obtained from V_g in Figure 7, should also be obtainable by extrapolating from the $J_{Constant}$ data at sufficiently high [free tubulin]. (C,D) Extrapolation to obtain $Q7$ from $J_{Constant}$. **Interpretation of panels C, D:** In each of the models, the value of $Q7$ is approximately equal to $Q3 \approx Q6$ (Figure 7). Thus, $Q7$ provides another way of estimating $CC_{Elongation}$. **Methods:** Regression lines were fitted to $J_{Constant}$ for [free tubulin] in ranges where the $J_{Constant}$ data are approximately linear as a function of [free tubulin]: from 6 to 7 μM for the simplified model (panel C) and from 14 to 20 μM for the detailed model (panel D). $Q7$ is the x-intercept of the regression line. Note that the [free tubulin] ranges used for determination of $Q7$ are higher than those used for $Q3$ and $Q6$ because the $Q7$ extrapolation requires conditions where (almost) all MTs in the population are growing (few if any depolymerization phases).

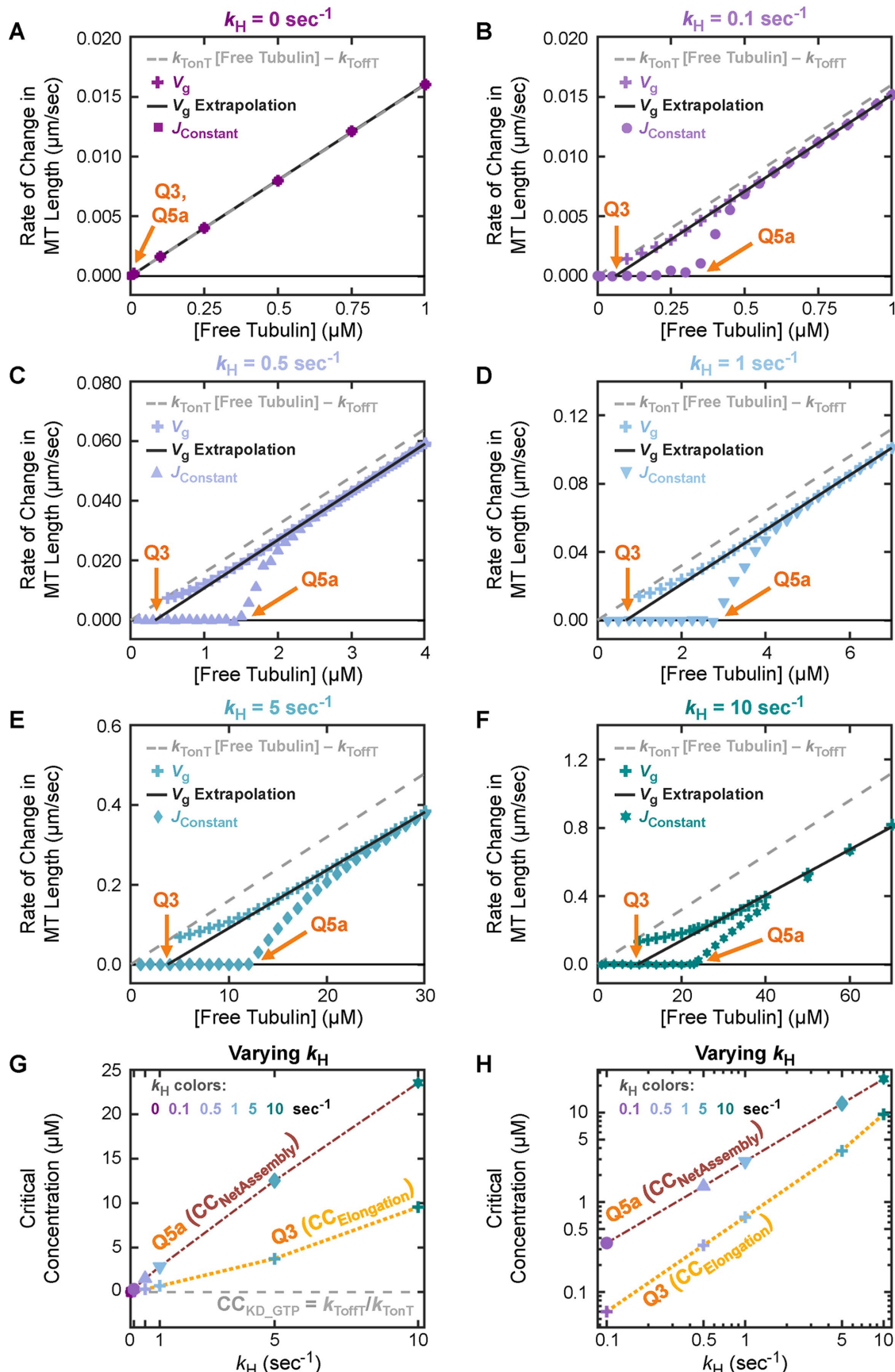
More specifically, because $Q3$ and $Q6$ are determined by extrapolations from regression lines fitted to plots of V_g versus [free tubulin], small changes in the V_g data (e.g., from noise) can be amplified in extrapolating to the $V_g = 0$ intercept. Additionally, in the simulation results, nonlinearities are observed in the V_g versus [free tubulin] plots in both models. In the presence of noise and/or nonlinearities, the values of $Q3$ and $Q6$ will depend on the [free tubulin] range where the regression lines are fitted to the V_g plots.

The deviations from linearity in the simulation plots are explained in part by measurement bias: at the lowest [free tubulin], there are few growing MTs, all of which are short (Figure 5, A and B; Supplemental Figure S3, C–F). The measured V_g data are biased toward the MTs that happened to grow fast enough and long enough to be detected. In particular, at low concentrations, there are very few detected growth phases, and the time in detected growth phases is a small fraction of the total time (see the DI measurements in the Supplemental Excel files). These observations indicate that the lowest concentrations should not be used in the linear extrapolation to

identify $Q3$ or $Q6$. To our knowledge, such deviations from linearity at low concentrations have not been detected experimentally. However, the simulations generate considerably more data and at smaller length thresholds than is possible with typical experiments. Because measurement bias could also be a problem in physical systems, we speculate that similar effects may eventually be seen experimentally.

Given the nonlinearities and the measurement bias described above, one might be concerned that detection thresholds would affect the measured value of $CC_{Elongation}$. We therefore compared two different analysis methods for determining V_g (Figure 7). Specifically, for the DI analysis method (Figure 7, + symbols), we set a threshold of 25 subunits (200 nm) of length change to detect growth or shortening phases (we set this threshold to be comparable to typical length detection limits in light microscopy experiments). In contrast, for the time-step method (Figure 7, square symbols), we did not impose a threshold on the length change during each time step (see the Supplemental Methods). The V_g results from the two

Simplified Model, Non-Competing Simulations



methods agree well with each other in the [free tubulin] range used to determine $CC_{Elongation}$ (i.e., the range where V_g is approximately linear). Thus, in implementing V_g analysis to estimate $CC_{Elongation}$, the regression lines should be fitted to the linear region to avoid the effect of detection thresholds. If the regression lines are not fitted in the tubulin range where V_g is linear, then Q3 and Q6 will be less accurate approximations of $CC_{Elongation}$.

Depending on the specific system, Q7 may be a less accurate approximation than Q3 or Q6. Q7 is obtained from $J_{Constant}$ at free tubulin concentrations that are sufficiently high so that (almost) all MTs are growing (i.e., where V_g and $J_{Constant}$ overlap, Figure 8). Because the Q7 extrapolation is performed from higher concentrations than the Q3 or Q6 extrapolations, measurement error or noise in the data can be further amplified. Moreover, V_g and $J_{Constant}$ may not overlap until tubulin concentrations are so high that experimental measurements may no longer be feasible (e.g., because of problems such as free nucleation).

Both in the detailed model and in physical MTs, an additional factor can cause V_g to have nonlinearities as a function of [free tubulin] and therefore likely interferes with the accuracy of identifying $CC_{Elongation}$ via Q3, Q6, or Q7. Previous work has provided experimental and theoretical evidence that the GTP–tubulin detachment rate depends on the tubulin concentration (Gardner *et al.*, 2011), which is contrary to the assumptions classically used to determine $CC_{Elongation}$ and would contribute to nonlinearity of V_g versus [free tubulin]. This observation has been explained by the occurrence of concentration-dependent changes in the MT tip structure (Coombes *et al.*, 2013).

If one wished to estimate $CC_{Elongation}$ from a competing system at steady state, V_g would need to be plotted as a function of the emergent [free tubulin]_{SteadyState}. However, measuring V_g and the emergent [free tubulin]_{SteadyState} in the same experiment might be impractical. Additionally, [free tubulin]_{SteadyState} in a competing system is below $CC_{NetAssembly}$, so this approach might not provide data over a range of [free tubulin] sufficient for the extrapolation to determine $CC_{Elongation}$. In other words, depending on the specific

system, most or all of the [free tubulin] range where the measured V_g is linear might be above $CC_{NetAssembly}$ and would therefore not be accessible in a steady-state competing system.

In summary, both detection issues and actual structural features can potentially make observed V_g measurements nonlinear with respect to [free tubulin]. As a result, the value obtained for $CC_{Elongation}$ from $Q3 \approx Q6 \approx Q7$ may depend on what [free tubulin] range is used for the linear fit. These observations mean that these values (Q3, Q6, Q7) provide at best approximate measurements of $CC_{Elongation}$.

Effect of the hydrolysis rate constant (k_H) on $CC_{Elongation}$ and $CC_{NetAssembly}$

The results above show that $CC_{Elongation}$ is obtained from measurements of individual MTs that are in the growth phase, while $CC_{NetAssembly}$ is obtained from measurements performed on populations (or on individual MTs over sufficient time) that include both growth and shortening phases (see also Hill, 1987; Walker *et al.*, 1988). Thus, the coexistence of growth and shortening phases (i.e., DI itself) occurs in conjunction with the separation between $CC_{Elongation}$ and $CC_{NetAssembly}$. DI in turn depends on nucleotide hydrolysis, since GTP–tubulin is prone to polymerization and GDP–tubulin is prone to depolymerization. Therefore, to develop an improved understanding of the separation between $CC_{Elongation}$ and $CC_{NetAssembly}$ in DI polymers, we next examined the effect of the hydrolysis rate constant k_H on $CC_{Elongation}$ and $CC_{NetAssembly}$. To allow a straightforward comparison between the observed behaviors and the input kinetic parameters, we utilized the simplified model.

More specifically, we ran simulations in the simplified model across a range of k_H values, while holding the other biochemical kinetic parameters constant, under both constant [free tubulin] (Figure 9; Supplemental Figures S5 and S6) and competing (Supplemental Figure S7) conditions. From these data, we determined $CC_{Elongation}$ as measured by Q3 (Figure 9; Supplemental Figure S5A), and $CC_{NetAssembly}$ as measured by Q5a (Figure 9; Supplemental Figure S5B) and also by $Q1 \approx Q2$ (Supplemental Figure S7). As an

FIGURE 9: Effect of varying the rate constant for nucleotide hydrolysis (k_H) in the simplified model. Noncompeting simulations of the simplified model were performed for various values of k_H (all other input kinetic rate constants are the same as in the other figures). Each of panels A–F corresponds to a different value of k_H , ranging from 0 to 10 s^{-1} , as indicated in the panel titles. (A–F) The growth velocity during growth phases (V_g ; + symbols; color-coded by k_H value) and the rate of change in average MT length ($J_{Constant}$; color and symbol vary by k_H value) as functions of [free tubulin]. We also plot the theoretical equation for V_g , which assumes that growing ends have only GTP–tubulin at the tips (gray dashed line). Note that the scales of the axes vary among panels A–F; for data replotted at the same scale, see Supplemental Figure S5. (G, H) $CC_{Elongation}$ and $CC_{NetAssembly}$ as functions of k_H , with $CC_{Elongation}$ and $CC_{NetAssembly}$ measured respectively by Q3 and Q5a from panels A–F. The axes have linear scales in panel G and log scales in panel H. The vertical separation between $CC_{Elongation}$ and $CC_{NetAssembly}$ at each k_H in the log–log plot (panel H) represents their ratio $CC_{NetAssembly}/CC_{Elongation}$. **Interpretation:** When k_H is zero (panel G; see also panel A), $CC_{Elongation}$ and $CC_{NetAssembly}$ are equal to each other and to CC_{KD_GTP} . As k_H increases (panels G and H; see also panels B–F), the values of $CC_{Elongation}$ and $CC_{NetAssembly}$ increase; they also diverge from each other and from CC_{KD_GTP} . Thus, the separations between CC_{KD_GTP} , $CC_{Elongation}$, and $CC_{NetAssembly}$ depend on k_H . To see how DI behavior relates to the CCs, see Supplemental Figure S6 for representative length–history plots of individual MTs at each k_H value presented here. At any given [free tubulin], as k_H increases, V_g and J each decrease relative to the rate for an equilibrium polymer with only GTP–tubulin ($k_{\text{Toff}}[\text{free tubulin}] - k_{\text{Toff}}$). V_g and J also diverge from each other. As discussed above, V_g and J differ from each other in the [free tubulin] range where DI occurs. **Methods:** The simulations were performed using the simplified model with 50 stable MT seeds. V_g was measured using the DI analysis method (Supplemental Methods). The steady-state $J_{Constant}$ was measured from the net change method (Supplemental Table S1B; see also Q5a, Table 4). All measurements were taken from 40 to 60 min. V_g data points are plotted only at concentrations where detected growth phases constituted at least 2% of the total time analyzed ($20 \text{ min} \times 50 \text{ MTs} = 1000 \text{ min analyzed}$). Regression lines (black solid line) were fitted to the V_g data points in the [free tubulin] range above $CC_{NetAssembly}$ and then extrapolated back to $V_g = 0$.

additional test of the effect of varying k_H , we also performed competing and noncompeting simulations in a parameter set with a higher value of CC_{KD_GTP} (Supplemental Figure S8).

When the hydrolysis rate constant k_H equals zero, only GTP-tubulin subunits are present. As would be expected, the behavior is that of an equilibrium polymer: no DI occurs (see length histories in Supplemental Figure S6A), and all observed CC values correspond to the K_D for GTP-tubulin as defined by the input rate constants. In other words, when k_H is zero, $CC_{Elongation} = CC_{NetAssembly} = CC_{KD_GTP} = k_{ToffT}/k_{TonT}$ (Figure 9A). When k_H is greater than zero in these simulations, both GTP- and GDP-tubulin subunits contribute to polymer dynamics, concurrent with the appearance of DI (Supplemental Figure S6, B–F). As k_H increases, $CC_{Elongation}$ (Q3) and $CC_{NetAssembly}$ (Q5a) both increase and diverge from each other (Figure 9; Supplemental Figures S5 and S8), and DI occurs over a wider range of [free tubulin] (Supplemental Figure S6). In all cases, the value of $CC_{NetAssembly}$ as measured by Q1 and Q2 is similar to the value as measured by Q5a (Figure 9; Supplemental Figures S5, S7, and S8).

$CC_{Elongation}$ can differ from CC_{KD_GTP} . In addition to showing that nucleotide hydrolysis drives $CC_{Elongation}$ (Q3) and $CC_{NetAssembly}$ (Q5a) apart, the results in Figure 9 also show that hydrolysis drives both away from CC_{KD_GTP} (x-intercept of gray dashed line in Figure 9, A–F; gray dashed line in Figure 9G). In particular, while the relationship $CC_{KD_GTP} = k_{ToffT}/k_{TonT}$ is independent of k_H , we observe that $CC_{Elongation}$ changes with k_H . This could be viewed as surprising because one might expect $CC_{Elongation}$ to equal CC_{KD_GTP} even in the presence of DI. The reasoning behind this expectation is as follows.

First, the rate of growth of an individual MT in the growth state has been assumed to change linearly with [free tubulin] according to the following relationship (Walker et al., 1988):⁴

$$V_g = k_{on}^{growth} [\text{free tubulin}] - k_{off}^{growth} \quad (2)$$

where k_{off}^{growth} and k_{on}^{growth} (called k_{-1}^e and k_2^e in Walker et al., 1988) are effective (observed) rate constants for loss and addition of GTP-tubulin subunits on a growing tip. By "effective" we mean that they are emergent quantities extracted from the V_g data, as opposed to directly measured kinetic rate constants. More specifically, the values of k_{on}^{growth} and k_{off}^{growth} are measured from the slope and the y-intercept, respectively, of a regression line fitted to V_g data, given Eq. 2. Because $CC_{Elongation}$ is measured as the value of [free tubulin] at which V_g is zero, setting Eq. 2 equal to zero and solving for [free tubulin] leads to the conclusion that $CC_{Elongation} = k_{off}^{growth}/k_{on}^{growth}$. This ratio $k_{off}^{growth}/k_{on}^{growth}$ is measured as the x-intercept of the regression line (Eq. 2; Walker et al., 1988).

Second, it is commonly assumed that rapidly growing tips have only GTP-subunits at the end (e.g., Howard, 2001; Bowne-Anderson et al., 2015). Under this assumption, and also assuming that all unpolymerized tubulin is bound to GTP, Eq. 2 becomes

$$V_g = k_{TonT} [\text{free tubulin}] - k_{ToffT} \quad (3)$$

which leads to the prediction that $CC_{Elongation} = k_{ToffT}/k_{TonT} = CC_{KD_GTP}$

Instead, the results (Figure 9, A–F; Supplemental Figures S5A and S8) show that Eq. 3 fits the data well only when k_H is close to zero. As k_H increases, the V_g regression line and $CC_{Elongation}$ diverge from the values that would be predicted from Eq. 3. More specifically, when k_H is greater than zero, the effective k_{on}^{growth} and k_{off}^{growth} (slope and intercept of V_g in Eq. 2) in the simulations diverge from k_{TonT} and k_{ToffT} ; in this case, V_g does not satisfy Eq. 3, and $CC_{Elongation}$ diverges from CC_{KD_GTP} . In considering these observations, recall that the kinetic rate constants (e.g., k_{TonT} , k_H) in our simulations are input by the user. In contrast, the values of V_g and $CC_{Elongation}$ are emergent properties of the system. Taken together, the results indicate that GDP subunits can influence behavior during growth phases.

Possible mechanisms for exposure of GDP-tubulin at growing microtubule tips. There are strong reasons to expect that GDP subunits will influence growth phase behavior in physical MTs. The idea that growing MT tips could have GDP-tubulin subunits might seem surprising, but GDP-tubulin subunits could become exposed on the surface of a growing tip either by detachment of a surface GTP subunit from a GDP subunit below it or by direct hydrolysis. The first mechanism conflicts with earlier ideas that GTP subunits rarely detach, but is consistent with recent experimental data indicating rapid exchange (attachment and detachment) of GTP subunits on MT tips (Gardner et al., 2011; Coombes et al., 2013; see also Margolin et al., 2012).

The idea that GDP-tubulin cannot be exposed at MT tips during growth phases may be a remnant of vectorial hydrolysis models,⁵ where GDP-tubulin would become exposed only when the GTP cap is entirely lost (at least for single-protofilament models). However, various authors have shown that vectorial hydrolysis is neither sufficient (Flyvbjerg et al., 1994, 1996; Padinhateeri et al., 2012) nor necessary (Margolin et al., 2012; Padinhateeri et al., 2012) to explain MT DI behavior.

Additionally, Hill and colleagues examined both vectorial and random hydrolysis models. In the vectorial hydrolysis model, the growth velocity satisfied an equation equivalent to Eq. 2, which assumes only GTP tips during growth (Hill, 1987). In their random hydrolysis model, the observed (emergent) slope and intercept of V_g did not equal the input rate constants for addition and loss of GTP subunits, as explicitly pointed out in (Hill and Chen, 1984; Hill, 1987). This conclusion from Hill's random hydrolysis model is consistent with the results of our model, which also has random hydrolysis.

The conclusion that $CC_{Elongation} \neq CC_{KD}$ also helps explain the observation from earlier in the paper that there is no concentration at which polymer assembly abruptly commences (i.e., there is no $CC_{PolAssem}$). Instead, the amount of polymer increases slowly with increasing [free tubulin] (Supplemental Figure S3, A–F). More specifically, although the MTs typically reach experimentally detectable lengths (e.g., >200 nm, depending on the method used) at some concentration above $CC_{Elongation}$ (Supplemental Figure S3, A–F), polymerization of a few subunits can occur even below $CC_{Elongation}$ (Supplemental Figure S3, E and F; square symbols in Figure 7). When [free tubulin] is above CC_{KD_GTP} , attachment to a GTP subunit will be more favorable than detachment; thus, small flickers of growth can occur. In contrast, $CC_{Elongation}$ is the [free tubulin] above

⁴The symbol \approx may be more appropriate than $=$ because this equation assumes 1) that V_g increases linearly with [free tubulin] and 2) that the detachment rate is independent of [free tubulin]. Our V_g results presented above indicate that assumption 1) may be inaccurate. See Gardner et al. (2011) for evidence against assumption 2).

⁵In vectorial hydrolysis models, hydrolysis occurs only at the interface between the GDP-tubulin lattice and the GTP-tubulin cap (e.g., Carlier et al., 1987; Hill, 1987). In contrast, in random hydrolysis models, any internal GTP subunit can hydrolyze (terminal GTP subunits may also hydrolyze, depending on the model).

which a MT may exhibit the extended growth phases of DI. The dependence of $CC_{Elongation}$ on k_H indicates that attachment must in some sense outweigh both detachment and hydrolysis of GTP subunits in order for extended growth phases to occur.

Dynamic instability can produce relationships previously interpreted as evidence of a nucleation process for growth from stable seeds

Previously, two experimental observations have been interpreted as evidence that growth of MTs from stable templates (e.g., centrosomes, axonemes, GMPCPP seeds) involves a nucleation process (e.g., conformational maturation or sheet closure; Wieczorek et al., 2015; Roostalu and Surrey, 2017). First, MTs are generally not observed growing at [free tubulin] near $CC_{Elongation}$. Second, when the fraction of seeds occupied is plotted as a function of [free tubulin], the shape of the resulting curve is sigmoidal, suggesting a cooperative process and/or a thermodynamic barrier. In this section, we show that these two nucleation-associated behaviors are observed in our simulations, which is notable because neither model incorporates an explicit nucleation step (our seeds are composed of nonhydrolyzable GTP-tubulin, so the rates of attachment to and detachment from the seed are the same as those for a GTP-tubulin tip). We show that both experimentally observed relationships can result from DI in combination with length detection thresholds. The behavior of DI polymers relative to $CC_{Elongation}$ and $CC_{NetAssembly}$, as described above (e.g., Figure 5, A and B; Supplemental Figure S3, A–F), can therefore be helpful in understanding these relationships.

Failure to detect MT growth events in experiments at [free tubulin] near $CC_{Elongation}$ can result from physical detection limitations coupled with DI. As described above, when [free tubulin] is near $CC_{Elongation}$, V_g is small and F_{cat} is high, meaning that MTs are short (Supplemental Figure S3, A–F) and short-lived (Figure 5, A and B); the average MT length remains small until [free tubulin] is closer to $CC_{NetAssembly}$ (Supplemental Figure S3, A–F). This behavior, coupled with length detection thresholds (such as would be imposed by physical experiments), could make it difficult to detect MTs at [free tubulin] near $CC_{Elongation}$. To test this hypothesis, we used the simulations (which output the MT length without any detection threshold) to examine the effect of imposing length detection thresholds similar to those in physical experiments.

Indeed, when we imposed a 200-nm detection threshold (comparable to light microscopy) on the length change needed for a growth phase to be recognized (Figure 7, + symbols), we saw that MT growth that was detected in the absence of this threshold (Figure 7, square symbols) was no longer detected. These results indicate that failure to observe MTs growing from stable seeds at [free tubulin] near $CC_{Elongation}$ can result from using experimental methods that have length detection limitations, providing evidence that such behavior can result from processes other than nucleation.

A sigmoidal P_{occ} curve is predictable from detection thresholds and microtubule population length distributions resulting from dynamic instability. P_{occ} is the proportion of stable MT templates/seeds that are occupied by a (detectable) MT (Figure 10, A and B). Previous experimental work has shown that P_{occ} has a sigmoidal shape when plotted as a function of [free tubulin] (e.g., Mitchison and Kirschner, 1984b; Walker et al., 1988; Wieczorek et al., 2015). This shape has been interpreted as evidence that starting a new MT from a seed is harder than extending an existing MT and thus that growth from seeds involves a nucleation process (e.g., Wieczorek

et al., 2015; compare Figure 11, A and B). However, the V_g analysis described above led us to hypothesize that this sigmoidal P_{occ} shape can also result from the combination of length detection thresholds and DI.

To test this hypothesis, we examined P_{occ} as a function of [free tubulin] with varying detection thresholds (Figure 10, C and D; Supplemental Figure S9). The results show that at each [free tubulin] (below $CC_{NetAssembly}$), as the detection threshold is increased, the detected P_{occ} decreases (i.e., fewer MTs are longer than the threshold), resulting in a sigmoidal shape emerging. The steepness of the sigmoid increases as the length detection threshold is increased. These observations indicate that the sigmoidal shape can result simply from imposing a length detection threshold on a system (such as dynamic MTs) where some of the filaments are shorter than the detection threshold. In the presence of DI with complete depolymerizations back to the seeds (as occurs below $CC_{NetAssembly}$), MTs will necessarily be below any nonzero detection threshold for at least some amount of time (see also Fygenson et al., 1994). Indeed, Hill previously presented a formula for P_{occ} as a function of the DI parameters and length detection thresholds (Hill, 1984).

The P_{occ} curve reaches 1 at [free tubulin] near $CC_{NetAssembly}$. The results in Figure 10 and Supplemental Figure S9 provide another observation relevant to understanding CCs: in both models, P_{occ} approaches 1 as [free tubulin] approaches $CC_{NetAssembly}$ (except possibly at very small thresholds, where P_{occ} nears 1 at lower [free tubulin]). This result is predictable, with or without a nucleation process, because only at [free tubulin] above $CC_{NetAssembly}$ (where the population undergoes net growth) would all active seeds be occupied by MTs longer than an arbitrarily chosen length threshold. This full occupancy would occur if sufficient time were allowed, because at [free tubulin] above $CC_{NetAssembly}$, all MTs will eventually become long enough to escape depolymerizing back to the seed. Thus, the idea that $P_{occ} = 1$ at [free tubulin] above $CC_{NetAssembly}$ after sufficient time may provide a practical way to identify $CC_{NetAssembly}$ experimentally (see also Chen and Hill, 1985a; Fygenson et al., 1994; Dogterom et al., 1995).

Taking all this information together, we propose that a combination of DI itself and the existence of detection thresholds contributes to phenomena (failure to observe growing MTs at [free tubulin] near $CC_{Elongation}$, Figure 7; and sigmoidal P_{occ} plots, Figure 10, Supplemental Figure S9) that have previously been interpreted as evidence that growth of MTs from stable seeds involves a nucleation process (e.g., Wieczorek et al., 2015). In fact, any process that makes growth from a seed more difficult than extension of a growing tip (e.g., a nucleation process such as sheet closure) would make the P_{occ} curve more steplike, not less so (Figure 11: compare panels B and C). While we cannot exclude the existence of nucleation processes such as conformational maturation or sheet closure in physical MTs, our work suggests that neither sigmoidal P_{occ} curves nor absence of detectable MTs on seeds at [free tubulin] near $CC_{Elongation}$ is sufficient evidence to conclude that growth from templates (e.g., centrosomes, stable seeds) involves a physical nucleation process.

DISCUSSION

The behavior of microtubules is governed by two major critical concentrations

Using dynamic MTs in our computational simulations, we examined the relationships between subunit concentration and polymer assembly behaviors for DI polymers. Our results show that there is no $CC_{PolAssem}$ as traditionally defined, meaning that there is no

Critical concentration	Representative figures	Critical concentration description	Equivalent to (see Table 1) ^a	Measured by (see Table 4)
$CC_{NetAssembly}$	1, A and C, 3–6	CC above which the polymer mass of a population will increase persistently, and individual filaments will undergo net growth over time	$CC_{SubSoln}$ ^b CC_{Flux} ^c $CC_{Unbounded}$	Q1, Q2, Q4, Q5
$CC_{Elongation}$	1B, 7, 8	CC above which individual filaments can exhibit the transient growth phases of DI	$CC_{Elongation}$	Q3, Q6, Q7
CC_{KD_GTP}	9	Equilibrium dissociation constant for binding of a free GTP subunit to a GTP subunit at a polymer tip		Any of the Q values above, under conditions where GTP is not hydrolyzed
CC_{KD_GDP}		Equilibrium dissociation constant for binding of a free GDP subunit to a GDP subunit at a polymer tip		GDP–tubulin alone does not form MTs, so CC_{KD_GDP} is not straightforwardly measured

For steady-state polymers (including DI polymers), $CC_{KD_GTP} \leq CC_{Elongation} \leq CC_{NetAssembly} \leq CC_{KD_GDP}$, but for equilibrium polymers, $CC_{KD} = CC_{Elongation} = CC_{NetAssembly}$.

^a $CC_{PolAssem}$ is not listed here because there is no threshold concentration at which polymers abruptly appear. Instead, the measurement classically expected to yield $CC_{PolAssem}$ (see Q1 in Table 4) actually yields $CC_{NetAssembly}$.

^b $CC_{SubSoln}$ is classically defined as the value of [free tubulin]_{SteadyState} in a competing system whenever [total tubulin] is above " $CC_{PolAssem}$ " (Table 1; Figure 1A). However, $CC_{SubSoln}$ is more accurately defined as the asymptote approached by [free tubulin]_{SteadyState} as [total tubulin] is increased (Q2 in Figures 3, A and B, and 4).

^cIt should be stressed that CC_{Flux} is the [free tubulin] at which the population-level fluxes of tubulin into and out of polymer are balanced, while individual MTs may grow and shorten when [free tubulin] = CC_{Flux} .

TABLE 3: Revised understanding of critical concentration for dynamically unstable polymers.

concentration where MTs abruptly come into existence. Instead, there are at least two major CCs. There is a lower CC ($CC_{Elongation}$), above which individual filaments can grow transiently, and an upper CC ($CC_{NetAssembly}$), above which the polymer mass of a population of filaments will increase persistently (Figure 12, A–C). For [free tubulin] above $CC_{NetAssembly}$, individual MTs may still undergo DI (Figure 12A, length history with triangle symbol), but will exhibit net growth over time (Figure 12A, triangle and circle symbols). What might be considered “typical” or “bounded” DI (where individual MTs repeatedly depolymerize back to the seeds) occurs at [free tubulin] between $CC_{Elongation}$ and $CC_{NetAssembly}$ (Figure 12, A and D, diamond symbols).

$CC_{Elongation}$ is estimated by Q3, Q6, and Q7, and $CC_{NetAssembly}$ is estimated by Q1, Q2, Q4, and Q5abc (Figure 12, C and F; Tables 3 and 4). Classical CC measurements (Q1 and Q2 in Figure 1A) do not yield the traditionally expected $CC_{PolAssem}$, but instead yield $CC_{NetAssembly}$ (Q1 and Q2 in Figure 12F). Importantly, [free tubulin]_{SteadyState} in a competing system does not equal $CC_{NetAssembly}$, but approaches $CC_{NetAssembly}$ asymptotically as [total tubulin] increases and depends on the number of stable seeds (Figure 12F; compare dark and light green lines).

Bulk polymer experiments can create the illusion that $CC_{NetAssembly}$ corresponds to $CC_{PolAssem}$. The conclusion that MTs grow transiently at [free tubulin] between $CC_{Elongation}$ and $CC_{NetAssembly}$ might appear to conflict with experimental observations reporting that bulk polymer is detectable only above Q1 (Figure 1A; see, e.g., Johnson and Borisy, 1975; Mirjigan et al., 2013). As discussed, Q1 provides a measure of $CC_{NetAssembly}$, but is traditionally expected to provide the CC for polymer assembly, $CC_{PolAssem}$. This apparent conflict can be resolved by recognizing that the fraction of total subunits converted to polymer will be small until the total tubulin concentration nears $CC_{NetAssembly}$ and will depend on the number of individual MTs in the population. In particular, for [total tubulin] < $CC_{NetAssembly}$, [free tubulin] will be approximately equal to [total tubulin], and [polymerized tubulin] will be low (Figure 12F, dark lines), unless there are many stable seeds (Figure 12F, light curves). In

contrast, for [total tubulin] > $CC_{NetAssembly}$, all free tubulin in excess of $CC_{NetAssembly}$ will be converted from free to polymerized form if sufficient time is allowed (Supplemental Figure S1, A–D).⁶ This conversion will happen because the average MT filament will experience net growth until [free tubulin] falls below $CC_{NetAssembly}$ (Figure 12D; compare early in time to later in time). The outcome of these relationships is that in bulk polymer experiments with few individual MTs, little if any MT polymer mass will be detected⁷ until the total tubulin concentration is above $CC_{NetAssembly}$ (Figure 12F, dark blue line), even though dynamic individual MT filaments can exist transiently at tubulin concentrations below $CC_{NetAssembly}$ (Figure 3, C and D). Thus, the experimental quantities Q1 and Q2 may look like the traditionally expected minimum concentration for polymer assembly ($CC_{PolAssem}$), but they actually represent the CC for persistent growth of a population’s polymer mass and net growth of individual MTs over time ($CC_{NetAssembly}$).

P_{occ} plots can create the illusion that there is a [free tubulin] at which microtubule assembly commences abruptly, i.e., that $CC_{PolAssem}$ exists. P_{occ} plots with length detection thresholds (such as thresholds intrinsic to microscope-based experiments) (Figure 10, A and B) may have led to the conclusion that there is a $CC_{PolAssem}$ at which P_{occ} first becomes positive. However, at low [free tubulin], MTs are short and short-lived as a result of low V_g and high F_{cat} , as described above, and therefore can be undetectable by standard microscopy. By varying the length detection threshold imposed on simulation data (Figure 10, C and D), it can be seen that the [free tubulin] at which P_{occ} first becomes positive depends on the threshold. These results, together with the polymer mass, average

⁶More precisely, as indicated by the earlier discussion of Figure 3, A and B, all subunits in excess of [free tubulin]_{SteadyState} will be converted to polymer; [free tubulin]_{SteadyState} is necessarily below but perhaps close to $CC_{NetAssembly}$.

⁷The amount of polymer present depends on the kinetic rate constants of the particular system and the number of stable seeds (Figure 4). The amount of polymer detected depends on the amount of polymer actually present and on what the experimental setup can detect.

Q value	Representative figures	Description of experimentally measurable quantity	CC estimated by Q
Q1	1A, 3, A and B, 4	Q1 is the x-intercept of the line (with slope = 1) approached by steady-state [polymerized tubulin] as [total tubulin] is increased in a competing system.	CC _{NetAssembly}
Q2	1A, 3, A and B, 4	Q2 is the horizontal asymptote approached by [free tubulin] _{SteadyState} as [total tubulin] is increased in a competing system.	CC _{NetAssembly} (= CC _{SubSoln})
Q3	1B, 7, A and B	Q3 is the [free tubulin] at which $V_g = 0$. Q3 is estimated by plotting V_g as a function of [free tubulin], fitting a regression line to the approximately linear part of the V_g data, and extrapolating back to the [free tubulin] at which $V_g = 0$.	CC _{Elongation}
Q4	1C, 6	Q4 is the [free tubulin] at which J , the rate of change in [polymerized tubulin], equals zero in a dilution experiment ($J < 0$ when dilution [free tubulin] < Q4; $J > 0$ when dilution [free tubulin] > Q4). ^a Q4 is determined by growing MTs to polymer-mass steady state at high [total tubulin], then rapidly diluting to a new [free tubulin] and measuring the rate of change in [polymerized tubulin] after a short delay. ^b	CC _{NetAssembly} (= CC _{Flux})
Q5 (a, b, and c)	5, C–F	Q5 is the [free tubulin] above which the rate of change in average MT length is positive in an experiment where [free tubulin] is held constant and the population has reached polymer-mass or polymer-growth steady state ($J = 0$ when [free tubulin] < Q5; $J > 0$ when [free tubulin] > Q5). ^c Q5 can also be described as the concentration above which the population drift coefficient is positive. We use the name Q5a, Q5b, or Q5c depending on how J is measured.	CC _{NetAssembly} (= CC _{Unbounded})
Q5a	5, C–F, 6, C and D	Q5a is Q5 with J calculated from the net rate of change in a population's average MT length between two time points; that is, $J = (\text{average length at time B} - \text{average length at time A}) / (\text{time B} - \text{time A})$, called J_{Net} in Mauro <i>et al.</i> (2019).	
Q5b	5, C and D	Q5b is Q5 with J calculated from measured DI parameters using the $J_{\text{DI_piecewise}}$ equation (Eq. 1b). Q5b is the [free tubulin] at which $V_g F_{\text{res}} = V_s F_{\text{cat}}$.	
Q5c	5, E and F	Q5c is Q5 with J calculated by summing displacements measured over short time steps (see the Supplemental Methods subsection on measuring drift coefficient, called J_{TimeStep} in Mauro <i>et al.</i> , 2019).	
Q6	7, C and D	Q6 is measured in the same way as Q3, but using growth phases from a dilution experiment after the system has been diluted into constant [free tubulin] conditions (instead of [free tubulin] being constant for the entire experiment, as with Q3).	CC _{Elongation}
Q7	8, C and D	Q7 is the x-intercept of the line approached by J as [free tubulin] is increased (note that J approaches the line when [free tubulin] >> CC _{NetAssembly}).	CC _{Elongation}

See Table 3 for descriptions of the CCs.

^a J can be defined in terms of polymer mass or average MT length (Supplemental Table S1; see also Section 1.3.2 of Mauro *et al.*, 2019): J = rate of change in [polymerized tubulin] = flux of tubulin into and out of polymer (e.g., in $\mu\text{M/s}$) or J = rate of change in average MT length = drift coefficient (e.g., in $\mu\text{m/s}$).

^bThe delay allows the GTP cap size to adjust in response to the new [free tubulin] after the dilution.

^cThe closer [free tubulin] is to CC_{NetAssembly}, the longer it will take for the system to reach steady state. If J is measured before polymer-mass steady state has been reached for [free tubulin] < CC_{NetAssembly}, then J will appear to be positive for [free tubulin] near but below CC_{NetAssembly}; this would make it difficult to identify the precise value of Q5. The transition from $J = 0$ to $J > 0$ at Q5 will be sharper the longer the system is allowed to run (see also Mauro *et al.*, 2019).

TABLE 4: Summary of experimentally measurable quantities (Q values) used to estimate CCs.

length, and maximal length data (Supplemental Figures S1, C–F, and S3, A–F), indicate that there is no concentration at which assembly of DI polymers commences abruptly.

Two additional critical concentrations help define polymer behaviors. In addition to the major CCs (CC_{Elongation} and CC_{NetAssembly}), there are at least two additional CCs that impact MT assembly. The first of these is CC_{KD_GTP} = $k_{\text{OffT}}/k_{\text{OnT}}$, which

corresponds to the K_D for binding of a free GTP–tubulin subunit to a GTP–tubulin at a MT tip. The second additional CC is the K_D for binding of a free GDP–tubulin subunit to a GDP–tubulin at a MT tip, CC_{KD_GDP} = $k_{\text{OffD}}/k_{\text{OnD}}$. Because CC_{KD_GTP} and CC_{KD_GDP} provide biochemical limits on the behavior of GTP–tubulin and GDP–tubulin, any CCs must lie between these two nucleotide-specific CCs (CC_{KD_GTP} ≤ CC_{Elongation} ≤ CC_{NetAssembly} ≤ CC_{KD_GDP}). CC_{KD_GTP} is the [free tubulin] above which GTP–tubulin polymers

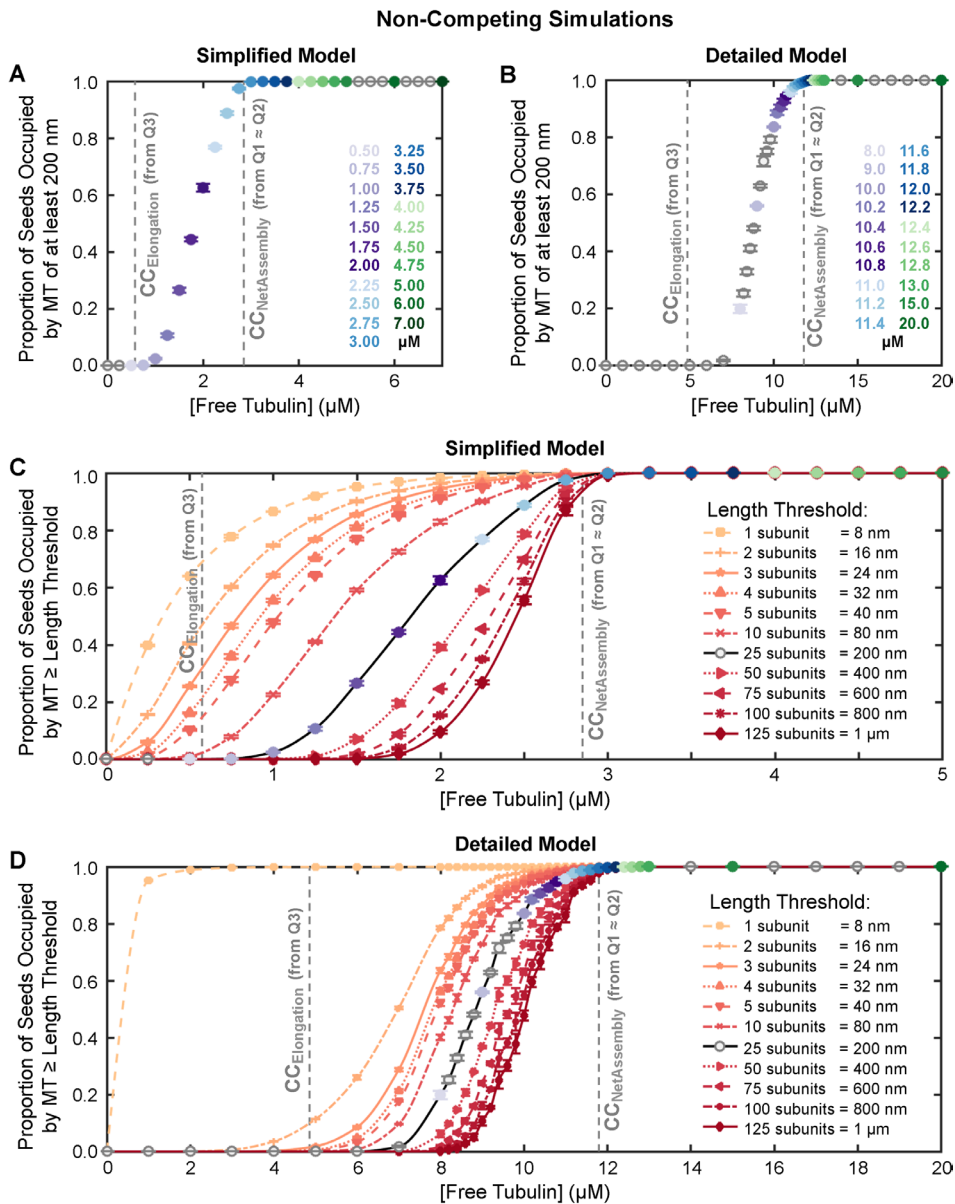


FIGURE 10: Relationship between P_{occ} (proportion of stable MT seeds that are occupied) and [free tubulin]. Simplified model in panels A, C; detailed model in panels B, D. The raw data analyzed in this figure are from the same noncompeting (constant [free tubulin]) simulations used in Figures 5, 6, C and D, 7, A and B, and 8. In all panels, the gray dashed lines represent $CC_{Elongation}$ (Q3 from Figure 7, A and B) and $CC_{NetAssembly}$ (Q1, Q2 from Figure 3, A and B).

(A, B) Proportion of stable seeds bearing “experimentally detectable” MTs (P_{occ}) as a function of [free tubulin]. Here detectable MTs are those with length ≥ 25 subunits = 200 nm (chosen because the Abbe diffraction limit for 540-nm (green) light in a 1.4 NA objective is ~ 200 nm). (C, D) P_{occ} with detection thresholds varied from 1 subunit (8 nm) to 125 subunits (1000 nm). The data with the 25-subunit threshold are replotted from panels A and B. **Interpretation:** The data in panels A and B show that with a detection threshold similar to that in typical fluorescence microscopy experiments, little polymer is observed growing off of the GTP-tubulin seeds in either model until [free tubulin] is well above $CC_{Elongation}$. More specifically, with this 200-nm threshold, P_{occ} does not reach 0.5 until [free tubulin] is more than halfway from $CC_{Elongation}$ to $CC_{NetAssembly}$. Note that the lowest value of [free tubulin] at which 100% of the seeds have a detectable MT corresponds to $\sim CC_{NetAssembly}$ (see also Fygenson et al., 1994; Dogterom et al., 1995). The data in panels C and D show that short MTs (with lengths below the 200-nm detection threshold from panels A and B) are present at free tubulin concentrations near $CC_{Elongation}$ (see also maximum MT length data in Supplemental Figure S3, C–F).

Additionally, we note that the P_{occ} curve of the detailed model is steeper than that of the simplified model when the same threshold is compared. We suggest that this results from the more cooperative nature of growth in the detailed (13-protofilament) model, which is an outcome of interactions between protofilaments. **Methods:** All data points represent the mean \pm one SD of the P_{occ} values obtained in three independent runs of the simulations. The values from each run are averages from 25 to 30 min, chosen so that P_{occ} has reached its steady-state value. MT length is measured as the number of subunit lengths above the seed. Note that in the detailed model, the MT length is the average of the 13 protofilament lengths and can therefore have noninteger values; see Supplemental Figure S9 for fractional thresholds below 2 subunits, which fill in the large gap between the thresholds of 1 and 2 subunits.

Schematics (not simulation data), Non-Competing Systems

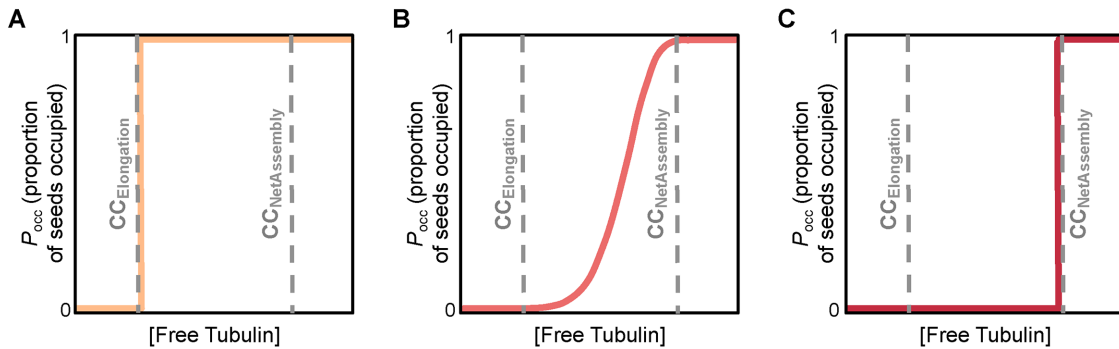


FIGURE 11: Hypothetical P_{occ} vs. [free tubulin] curves, where P_{occ} is the proportion of seeds that are occupied by MTs.

It might have been expected that GTP-like seeds should start growing once [free tubulin] is above $CC_{Elongation}$, and that P_{occ} would therefore increase abruptly from 0 to 1 for [free tubulin] at or just above $CC_{Elongation}$, similarly to the step function in panel A. In contrast, sigmoidal P_{occ} curves, similar to panel B, have been observed experimentally (Mitchison and Kirschner, 1984b; Walker et al., 1988; Dogterom et al., 1995; Wieczorek et al., 2015). Obtaining a sigmoidal shape (B) instead of a step function (A) has been interpreted as evidence of a nucleation process that makes growth of MTs from stable seeds more difficult than extension from a growing end (e.g., Wieczorek et al., 2015). However, as discussed in the text, this sigmoidal shape can be a consequence of DI in combination with experimental length-detection limitations, and therefore is not necessarily evidence of a nucleation process. Note that a nucleation process that makes growth from seeds more difficult would lead to a P_{occ} curve that increases more rapidly from 0 to 1 and does so at [free tubulin] near $CC_{NetAssembly}$, similarly to the step function in panel C. This behavior can be explained in the following way. When [free tubulin] is below $CC_{NetAssembly}$, MTs will repeatedly depolymerize back to the seed. When nucleation from seeds is difficult, it will take longer for a new growth phase to initiate after each complete depolymerization; seeds will therefore remain unoccupied for longer times and the proportion of seeds in the population that are occupied at any particular time will be smaller. Thus, as the difficulty of nucleation increases, the shape of the P_{occ} curve would change from a sigmoid (as in panel B) to a step function at $CC_{NetAssembly}$ (as in panel C).

will grow in the absence of hydrolysis and provides the lower limit for short-term assembly of polymers in the presence of hydrolysis. As the hydrolysis rate constant increases, $CC_{Elongation}$ (the [free tubulin] above which the growth phases of DI can occur) can diverge from CC_{KD_GTP} (Figure 12G; compare yellow $CC_{Elongation}$ line to gray CC_{KD_GTP} line). Unlike CC_{KD_GTP} , CC_{KD_GDP} is not straightforwardly measurable for MTs, because GDP-tubulin subunits alone do not polymerize into MTs (Howard, 2001), but could be relevant to other steady-state polymers (e.g., actin).

Separation between the critical concentrations is created by GTP hydrolysis. By running simulations in the simplified model at different k_H values, we show that increasing k_H causes $CC_{Elongation}$ and $CC_{NetAssembly}$ to diverge from each other and from CC_{KD_GTP} (Figure 9; Supplemental Figures S5 and S8). We expect that the magnitude of the separation between the various CCs will depend not on the value of k_H per se, nor on any individual rate constants, but rather on the relationships between the various rate constants. This is a topic of ongoing investigation. We speculate that the separation between the CCs has significance for understanding the difference between actin and tubulin, as discussed more below.

Relationship to previous work

As discussed above, the idea that MTs and other steady-state (energy-using) polymers have two major CCs was first investigated in depth by Hill and colleagues, who studied the behavior of these systems using a combination of theory, computational simulations, and experiments (Carlier et al., 1984a; Hill and Chen, 1984; Hill, 1987). Their c_1 corresponds to our $CC_{Elongation}$; their c_0 corresponds to our $CC_{NetAssembly}$ (Hill and Chen, 1984; note that these CCs were given other variable names elsewhere in Hill's works).

Moreover, Hill and Chen concluded that "rather long" MTs grow at concentrations "well below" what they referred to as the "real" CC (corresponding to our Q4 in Figure 1C), in contrast to equilibrium polymers, where the average length is very small until [free subunit] is "extremely close" to the CC (Hill and Chen, 1984). However, the significance of this work for MT DI behavior was not fully incorporated into the CC literature, perhaps because it was not clear how Hill's two CCs related to classical CC measurements (e.g., Q1 and Q2 in Figure 1A).

Walker et al.'s (1988) seminal article on DI parameters included measurements of two different CCs, which they termed the CC for elongation ($CC_{Elongation}$ in our notation) and the CC for net assembly ($CC_{NetAssembly}$ in our notation). They calculated their value of the CC for net assembly from their measured DI parameters using a version of the J_{DI} equation (see equation on p. 1445 of Walker et al., 1988). However, perhaps because the article focused on $CC_{Elongation}$ and did not directly relate either of these CCs to those predicted by Hill and colleagues, the idea that MTs have two CCs still did not become widely acknowledged. Soon thereafter, the articles of Dogterom et al. and Fygenson et al. were important in showing clearly and intuitively how the behavior of MTs changes at the CC for unbounded growth (equivalent to $CC_{NetAssembly}$), which they described using the J_{DI} equation shown in Eq. 1 (Dogterom and Leibler, 1993; Dogterom et al., 1995; Fygenson et al., 1994). However, these authors did not relate their CC for unbounded growth to the CCs discussed by Hill or Walker et al. or to more classical CCs (Table 1; Figure 1).

Some of the continued confusion about CC may have resulted from the fact that published experimental work typically involved either competing conditions or noncompeting conditions but not both. More specifically, classical experiments for determining "the critical concentration" (e.g., Figure 1A) involved competing

Summary of Conclusions

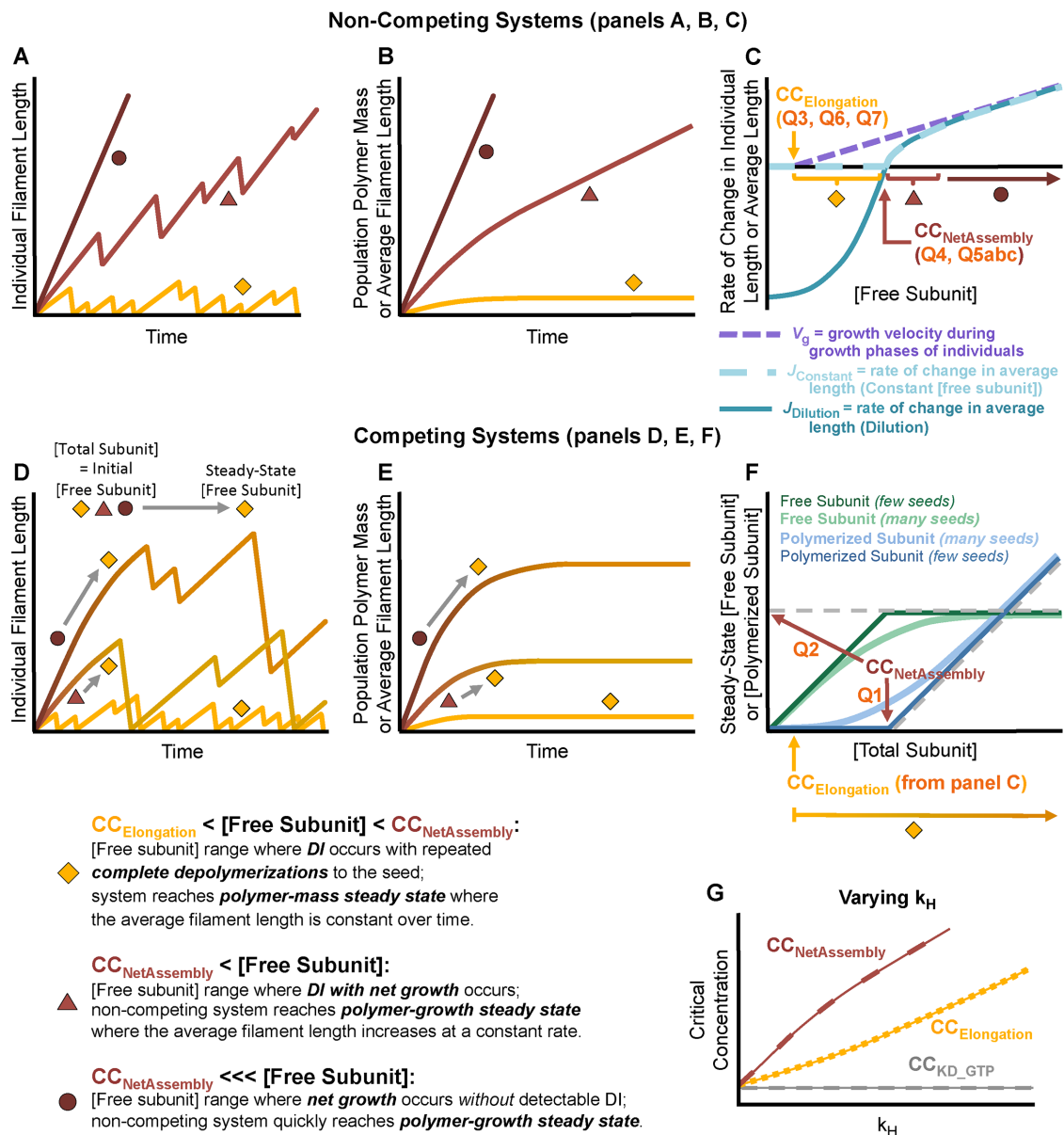


FIGURE 12: Schematic summary of the relationships between DI behavior and CCs for DI polymers. First row: noncompeting systems ([free subunit] is held constant over time). Second row: competing systems ([total subunit] is held constant over time). Symbol key at bottom left. CCs and Q values are summarized in Tables 3 and 4. (A, D) Length histories of individual filaments in systems with three different values of [free subunit] (panel A) or [total subunit] (panel D). (B, E) The polymer mass or average filament length of populations containing the individuals in panels A or D as indicated by the corresponding symbols (similar to Supplemental Figures S3, A and B, and S1, C and D, respectively). (C) Rate of polymerization/depolymerization vs. [free subunit] for individual filaments during growth phases (V_g) and populations of filaments (J_{Constant} , J_{Dilution}). More specifically, the panel shows 1) the growth velocity of individual filaments during the growth phase (V_g , purple dashed line; similar to Figure 7); 2) the net rate of change in average filament length of a population of filaments as assessed from experiments with [free subunit] held constant for the entire time of the experiment (J_{Constant} , light turquoise dashed curve; similar to Figure 5, C–F); and 3) the net rate of change in average filament length of a population of filaments as assessed from dilution experiments (J_{Dilution} , dark turquoise solid curve; similar to Figure 6). Notice that J_{Constant} and J_{Dilution} are superimposed for any [free subunit] $> CC_{\text{NetAssembly}}$, and that these two curves approach V_g for [free subunit] $>> CC_{\text{NetAssembly}}$. (F) Emergent [free subunit] (green) or [polymerized subunit] (blue) as functions of input [total subunit] for competing systems at polymer-mass steady state (similar to Figures 3, A and B, and 4). (G) Effect of changing k_H , the rate constant for nucleotide hydrolysis (similar to Figure 9G). **Interpretations:**

- Behavior of individual filaments for [free subunit] below and above $CC_{\text{NetAssembly}}$: In both competing and noncompeting systems, when [free subunit] is between $CC_{\text{Elongation}}$ and $CC_{\text{NetAssembly}}$, individual filaments display steady-state DI in which they eventually and repeatedly depolymerize back to the seed

conditions, but much of the previous work described above was performed under conditions of constant [free tubulin] (e.g., Figure 1, B and C). Walker *et al.* (1988) did note in their Discussion section that the concentration of free tubulin at steady state in their competing system was below their calculated CC for net assembly, contrary to the expectation that $[\text{free tubulin}]_{\text{SteadyState}}$ would equal the CC for net assembly. They attributed this difference to "uncertainties inherent in [their] assumptions and measurements" (Walker *et al.*, 1988). Instead, as shown above, the observation that $[\text{free tubulin}]_{\text{SteadyState}}$ approaches $\text{CC}_{\text{NetAssembly}}$ without actually reaching it is a predictable aspect of DI. More specifically, $[\text{free tubulin}]_{\text{SteadyState}}$ will be measurably below $\text{CC}_{\text{NetAssembly}}$ if [total tubulin] is not high enough relative to the value of $\text{CC}_{\text{NetAssembly}}$ and/or if the number of stable seeds is large (Figures 3, A and B, and 4).

More recently, Mourão *et al.* (2011) focused on systems of MTs growing under competing conditions. Using stochastic simulations and mathematical analysis to study MT growth from stable seeds, they examined a quantity that they called "a baseline steady state free subunit concentration (MD_{SS})," which is conceptually similar to our $\text{CC}_{\text{SubSoln}}$ (measured by Q2). They concluded that $[\text{free tubulin}]_{\text{SteadyState}}$ is not equal to MD_{SS} but below it; our results are consistent with this conclusion. In particular, they demonstrated how the separation between $[\text{free tubulin}]_{\text{SteadyState}}$ and MD_{SS} depends on various factors including the number of stable MT seeds. The dependence of MT behavior on subunit concentration was not their primary focus, so they did not explicitly show that $[\text{free tubulin}]_{\text{SteadyState}}$ asymptotically approaches $MD_{SS} = \text{CC}_{\text{NetAssembly}}$ as [total tubulin] increases (Figures 3, A and B, and 4); however, they did perform simulations at three different values of [total tubulin], and their results are consistent with our conclusions. Additionally, the criterion that they used to determine the value of MD_{SS} is that MD_{SS} is the free tubulin concentration at which $V_g/|V_s| = F_{\text{cat}}/F_{\text{res}}$. We note that this equation is algebraically equivalent to $|V_s|F_{\text{cat}} = V_g F_{\text{res}}$,

which was the criterion given by Dogterom and Leibler (1993) for identifying the CC for unbounded growth (equivalent to our $\text{CC}_{\text{NetAssembly}}$).

Thus, there has been a need for a unified understanding of how CCs relate to each other and to MT behaviors at different scales. Our work fills this gap by clearly showing how the behaviors of individual MTs and populations of MTs relate to each other, to [free subunit] and [total subunit], and to a range of different experimental measurements in both competing and noncompeting systems (conclusions summarized in Figure 12 and Tables 3 and 4). Taken together, our simulations and analyses should provide a more solid foundation for understanding the behavior of MTs and other DI polymers under varied concentrations and experimental conditions.

Implications for systems with free nucleation

As discussed above, all of our studies in this article are performed with a defined number of stable MT seeds and one free end. For systems with two free ends (plus and minus), each end would have its own values of $\text{CC}_{\text{Elongation}}$ and $\text{CC}_{\text{NetAssembly}}$, and behaviors of the individual ends would depend on the system's [free tubulin] relative to the CCs. Additional behaviors such as filament treadmilling could arise.

In systems with free nucleation and disappearance of MTs, the number of MTs in a population can change over time. If the number of MTs is increasing with time, then the population's polymer mass could increase while the average filament length stays the same or even decreases; this could occur, for example, with constant [free tubulin] below $\text{CC}_{\text{NetAssembly}}$ and a high rate of free nucleation, as could occur if there is autocatalytic nucleation (Ishihara *et al.*, 2016).

In a competing system with free nucleation at steady state, the sharpness of the transition at Q1 and Q2 would likely depend on the difficulty of nucleation; difficult nucleation would lead to a sharp transition, whereas the transition would become more gradual with an increasing rate of nucleation (Johnson and Borisy, 1975). For

(panels A and D, diamond symbols). When [free subunit] is above $\text{CC}_{\text{NetAssembly}}$, individual filaments display net growth over time, while still undergoing DI (panel A, triangle symbol) except perhaps at very high concentrations (panel A, circle symbol).

- **Behavior of individual filaments in competing systems over time:** If initial [free subunit] > $\text{CC}_{\text{NetAssembly}}$ in a competing system, then individual filaments initially experience net growth (panel D, triangle and circle symbols). Because [total subunit] is constant in competing systems, [free subunit] decreases as subunits polymerize. Net growth of individual filaments continues until [free subunit] drops below $\text{CC}_{\text{NetAssembly}}$.
- **Polymer-mass and polymer-growth steady states:** Given sufficient time, competing systems reach polymer-mass steady state, where the average filament length (panel E), the population's [polymerized subunit] (panels E and F), and [free subunit] (panel F) have finite steady-state values governed by the input [total subunit]. In noncompeting systems, polymer-mass steady state is reached only if [free subunit] is below $\text{CC}_{\text{NetAssembly}}$ (panel B, diamond symbol). In contrast, noncompeting systems with [free subunit] above $\text{CC}_{\text{NetAssembly}}$ reach polymer-growth steady state (panel B, triangle and circle symbols), where the average filament length increases at a constant rate J (panel C).
- **Effect of number of stable MT seeds in competing systems:** $[\text{Free subunit}]_{\text{SteadyState}}$ in competing systems does not equal $\text{CC}_{\text{NetAssembly}}$ but approaches $\text{CC}_{\text{NetAssembly}}$ as [total subunit] is increased, and the sharpness of the approach depends on the number of seeds (panel F; compare light and dark curves). In particular, for many seeds, $[\text{free subunit}]_{\text{SteadyState}}$ is noticeably below $\text{CC}_{\text{NetAssembly}}$ even at very high [total subunit].
- **Varying k_H :** When $k_H = 0$ (equilibrium polymer), there is one CC: $\text{CC}_{\text{KD_GTP}} = \text{CC}_{\text{Elongation}} = \text{CC}_{\text{NetAssembly}}$. When $k_H > 0$ (steady-state polymer), $\text{CC}_{\text{Elongation}}$ and $\text{CC}_{\text{NetAssembly}}$ are distinct from each other and from $\text{CC}_{\text{KD_GTP}}$. At sufficiently low k_H , $\text{CC}_{\text{Elongation}}$ and $\text{CC}_{\text{NetAssembly}}$ would be experimentally indistinguishable. As k_H increases, $\text{CC}_{\text{Elongation}}$ and $\text{CC}_{\text{NetAssembly}}$ both increase, but $\text{CC}_{\text{KD_GTP}}$ does not change with k_H . The [free subunit] range where DI with complete depolymerizations occurs, that is, the range between $\text{CC}_{\text{NetAssembly}}$ and $\text{CC}_{\text{Elongation}}$ (bracket with diamond symbol in panel C), increases with k_H .

biologically relevant free nucleation rates, the transition would likely be sharp, and one might mistakenly conclude that $CC_{NetAssembly}$ corresponds to a hypothetical CC for nucleation. This is because only when $[free\ tubulin] > CC_{NetAssembly}$ would newly nucleated MTs be likely to persist.

Studies of systems with two free ends and free nucleation are a topic for future work.

Concurrence between different approaches for measuring microtubule behavior has practical significance

As shown in Figure 5, C–F, there is concurrence between three seemingly disparate ways of analyzing MT behavior by measurements of J (flux of subunits into and out of polymer): 1) the net rate of change in $[polymerized\ tubulin]$ (Figure 5, C–F, o symbols), which is a bulk property obtained by assessing the change in mass of the population of polymers between two points in time; 2) the J_{DI} equation (Figure 5, C and D, + symbols), which uses DI parameters extracted from individual filament length history plots obtained over tens of minutes; 3) the drift coefficient (Figure 5, E and F, x symbols; Supplemental Figure S3, G and H, all symbols) as measured by observing individual MTs in a population of MTs for short periods of time (e.g., 2-s time steps across as little as 1 min). These approaches differ in attributes including physical scale, temporal scale, and experimental design. While the similarity of the data produced by these different approaches may initially be surprising, it can be shown that the equations underlying these measurements are algebraically equivalent if certain assumptions are met (Mauro *et al.*, 2019). The agreement between the results of these measurements indicates that the experimentally more tractable time-step approach (Komarova *et al.*, 2002) can be used to measure $CC_{NetAssembly}$ and should be used more frequently to quantitatively assess MT assembly behavior in the future (see Mauro *et al.*, 2019, for additional considerations relevant to implementing this approach).

Biological significance of having two major critical concentrations

The understanding of critical concentration as presented above should help resolve apparently contradictory results in the MT literature. In particular, our results indicate that reported measurements of “the” CC for MT polymerization vary at least in part because some experiments measure $CC_{Elongation}$ (e.g., Walker *et al.*, 1988; Wieczorek *et al.*, 2015), while others measure $CC_{NetAssembly}$ (e.g., Carlier *et al.*, 1984a; Dogterom *et al.*, 1995; Mirigian *et al.*, 2013). This clarification should help in design and interpretation of experiments involving CC, especially those investigating the effects of MT-binding proteins (e.g., Amayed *et al.*, 2002; Wieczorek *et al.*, 2015; Hussmann *et al.*, 2016), osmolytes (e.g., Schummel *et al.*, 2017), or drugs (e.g., Buey *et al.*, 2005; Verma *et al.*, 2016).

Furthermore, it is important to emphasize that $CC_{Elongation}$ and $CC_{NetAssembly}$ are fundamental attributes of a specific type of tubulin in a particular environment, similarly to the way a K_D characterizes a protein–protein interaction or a K_M characterizes an enzyme–substrate reaction. Thus, we suggest using $CC_{Elongation}$ (e.g., as measured by Q3) and $CC_{NetAssembly}$ (especially as measured by Q5c from the time-step drift coefficient approach) in addition to using DI parameters as a way to characterize tubulin (or other proteins that form polymers) and the activities of proteins that alter polymer assembly (see also the discussion in Komarova *et al.*, 2002).

CCs can also be helpful in connecting the behavior of MTs between *in vitro* and *in vivo* studies, including studies of how cells can

exploit CCs for regulatory purposes. For example, MTs in many interphase cell types grow persistently (perhaps with catastrophe and rescue, but with net positive drift) until they reach the cell edge, where they undergo repeated cycles of catastrophe and rescue with rare complete depolymerizations (Komarova *et al.*, 2002). We showed previously that this persistent growth is a predictable outcome of having enough tubulin in a confined space: if sufficient tubulin is present, the MTs grow long enough to contact the cell boundary, which causes catastrophe; this drives the $[free\ tubulin]$ above its natural steady-state value, which reduces catastrophe, enhances rescue, and induces the persistent growth behavior (Gregoretti *et al.*, 2006). In light of the current results, we can now phrase this previous work more succinctly: persistent growth of MTs in interphase cells occurs when catastrophes induced by the cell boundary drive $[free\ tubulin]$ above $CC_{NetAssembly}$. In contrast, at mitosis, when the MTs are more numerous and thus shorter, $[free\ tubulin]$ remains below $CC_{NetAssembly}$. Furthermore, a cell can potentially regulate the transition from interphase to mitosis through MT binding proteins that alter the value of $CC_{NetAssembly}$ and/or the degree of nucleation. See also Verde *et al.*, 1992; Dogterom and Leibler, 1993; Dogterom *et al.*, 1995; Gregoretti *et al.*, 2006; Vorobjev and Maly, 2008; and Mourão *et al.*, 2011, for relevant data and discussions.

Relevance to other steady-state polymers

Though our studies presented here were formulated specifically for MTs, we suggest that they can be applied to any nucleated steady-state polymers that display DI, and perhaps to steady-state polymers more broadly. In particular, we propose that the key characteristic that distinguishes dynamically unstable steady-state polymers (e.g., mammalian MTs) from other steady-state polymers (e.g., mammalian actin) is as follows: for DI polymers, $CC_{Elongation}$ and $CC_{NetAssembly}$ are separable values driven apart by hydrolysis, but for other polymers, they are either identical (as is true for equilibrium polymers) or so close as to be nearly superimposed (e.g., mammalian actin). Whether or not DI is physiologically relevant for a given polymer type in a specific cellular environment will depend on how the values of $CC_{Elongation}$ and $CC_{NetAssembly}$ relate to the cellular subunit concentration.

MATERIALS AND METHODS

Simulations

Simplified model (Figure 2A). The simplified model of stochastic MT dynamics was described previously (Gregoretti *et al.*, 2006), but the implementation used here was updated significantly. First, the code was rewritten in Java so that it could be implemented more easily on personal computers. Second, the time between events is now sampled using an exact version of the Gillespie algorithm (Gillespie, 1976), instead of an approximate version with a fixed time step. This change improves the accuracy with which the simulation carries out the underlying biochemical model with user-input rate constants. Third, the simulation was adjusted so that each simulated subunit now corresponds to an 8-nm MT ring (1×13 dimers) instead of a 20-nm MT brick (2.5×10 dimers) as in Gregoretti *et al.* (2006). Also, the simulations in Gregoretti *et al.* (2006) had a cell edge, which limited the MT lengths; the simulations presented here have no physical constraints on the MT lengths. The change in subunit size and the lack of a physical boundary in the present simulation mean that the numerical values of the DI parameters and Q measurements (Figures 3–8, left panels) are not directly comparable between this implementation and our earlier publication (Gregoretti *et al.*, 2006). However, the general behavior of the simulation is the same. The input parameters used here are as follows:

k_{TonT}	2.0 / $\mu\text{M s}$	kinetic rate constant for addition of GTP-tubulin onto GTP MT end
k_{TonD}	0.1 / $\mu\text{M s}$	kinetic rate constant for addition of GTP-tubulin onto GDP MT end
$k_{\text{ToffT}}, k_{\text{ToffD}}$	0.0 /s	kinetic rate constant for loss of GTP-tubulin from GTP or GDP MT end
$k_{\text{DoffT}}, k_{\text{DoffD}}$	48 /s	kinetic rate constant for loss of GDP-tubulin from GTP or GDP MT end
k_{H}	1 /s	kinetic rate constant for nucleotide hydrolysis (GTP-tubulin \rightarrow GDP-tubulin)
Vol	500 fL	volume of simulation

Unless otherwise indicated, each of the simplified model simulations was run with MTs growing from 100 stable seeds composed of nonhydrolyzable GTP-tubulin.

Detailed model (Figure 2B). The detailed model of stochastic MT dynamics was first developed in Margolin *et al.* (2011, 2012) and later utilized in Gupta *et al.*, 2013; Li *et al.*, 2014; Duan *et al.*, 2017, and Mauro *et al.*, 2019. The core simulation is the same as in these prior publications, but this version has minor modifications including the addition of a dilution function to enable production of $J(c)$ plots such as those in Figure 6. Please refer to Margolin *et al.* (2012) for detailed information on the model, its parameter set C, and how its behavior compares with that of *in vitro* DI. Briefly, parameter set C (used here) was tuned in Margolin *et al.* (2012) to approximate Walker's DI parameters for *in vitro* dynamics of mammalian brain MTs at [free tubulin] = 10 μM (Walker *et al.*, 1988). Unless otherwise indicated, each of the detailed model simulations was run with MTs growing from 40 stable seeds composed of nonhydrolyzable GTP-tubulin in a volume of 500 fL.

For both models, the Supplemental Excel file provides the numerical values of the DI parameters, the numbers of growth and shortening phases, total times in growth and shortening, and total length changes during growth and shortening, as measured by our automated DI analysis tool (described in the Supplemental Methods). The values of the DI parameters for the detailed model are similar to those that we published previously for this model (Margolin *et al.*, 2012; Duan *et al.*, 2017).

Analysis

$\text{CC}_{\text{NetAssembly}}$ is estimated by determining Q1, Q2, Q4, or Q5abc (Figures 3–6). $\text{CC}_{\text{Elongation}}$ is estimated (perhaps poorly) by determining Q3, Q6, or Q7 (Figures 7 and 8). See Table 4 for information on how to perform each of the Q measurements. The figure legends provide details about applying the measurements to the simulation data, including information about the time periods during which measurements were performed. The time periods were chosen to ensure that the variable being measured (e.g., rate of change in average length) has reached its steady-state value. For most of the measurements, this occurs when the simulated system has reached either polymer-mass steady state (noncompeting systems with [free tubulin] < $\text{CC}_{\text{NetAssembly}}$, Supplemental Figure S3, A and B; competing systems, Supplemental Figure S1, A–D) or polymer-growth steady state (noncompeting systems with [free tubulin] > $\text{CC}_{\text{NetAssembly}}$, Supplemental Figure S3, A and B). In the Supplemental Methods, we describe the time-step analysis method (based on Komarova *et al.*, 2002) used to measure drift and V_g , as well as our DI analysis method, used to measure V_g , V_s , F_{cat} , and F_{res} .

Code availability

The simulation codes (written in Java) and analysis codes (written in MATLAB) are available upon request.

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