Trafficking synaptic materials in complex morphologies involves tradeoffs in speed and precision

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ABSTRACT

Long lived changes in synaptic strength underlie persistent memories and in many cases require activity dependent transcription of plasticity-related genes. A well-known problem is to understand how appropriate gene products, such as synaptic proteins and mRNAs, get trafficked to the synapses that require them in spite of the complex dendritic arbors and thousands of postsynaptic sites typical of many kinds of neuron. Using a simple, biophysically-rooted model of microtubule transport we show that rapid delivery of plasticity-related cargo entails loss of specificity in the synapses that eventually receive it. The model predicts that neurons must balance the biochemical parameters that control trafficking to find a compromise between the precision and speed with which long-lived connectivity patterns are formed. Conversely, imbalances in transport parameters could lead to hypo-, or hyperconnectivity as well as loss of precision in associative memory, identifying a potential role for transport machinery in brain disorders that exhibit connectivity and plasticity phenotypes.

Keywords: Synaptic tagging hypothesis, LTP, Transport, Morphology

INTRODUCTION

Neurons often have complex dendritic trees, reflecting the dense and structured synaptic connectivity that supports nervous system function. In order to maintain appropriate connectivity, or to alter it as a means of encoding memories, numerous enzymes and macromolecules need to be transported along dendrites to the synapses that need them. How is this logistical problem solved at the subcellular level? And how might a biologically plausible solution constrain the behavior of synaptic plasticity 'rules' and learning in general?

Transport along dendrites is mediated by motor proteins (kinesins and dyneins) that walk along microtubules, carrying cargo at a rate that far exceeds passive diffusion [REFS]. The mechanisms that detach and sequester cargo at specific places in dendrites are not fully understood, but are known to depend on local biochemical signals that can be influenced by synaptic activity, such as transient elevations in cyclic AMP or $[Ca^{2+}]$ Mironov (2007); Wang and Schwarz (2009). It is therefore conceivable that synaptic cargo transport is controlled in a distributed fashion, in which synapses that require cargo trigger its release from the motor protein complexes as they pass. This idea is a variant of the well-known synaptic tagging hypothesis (Frey and Morris, 1997), which proposes that synapses produce a biochemical 'tag' that signals a requirement for synaptic building blocks as part of the plasticity process.

Not all aspects of synaptic plasticity require delivery of cargo from the soma. For example, some forms of long-term potentiation (LTP) depend on local protein synthesis and can function in isolated dendrites [REFS]. However, in the long run, the mRNAs and the machinery that supports such plasticity

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needs to be replenished. Moreover, many long-lasting forms of synaptic plasticity rely on anterograde transport of mRNAs (Kandel, 2001; Puthanveettil et al., 2008) and specific mRNAs are known to be selectively transported to regions of heightened synaptic activity (Steward et al., 1998; Steward and Worley, 2001; Moga et al., 2004) or to developing synaptic contacts (Lyles et al., 2006). The interplay between global distribution and local plasticity mechanisms therefore demands attention.

From simple assumptions about the dynamics of active dendritic transport we develop a model that can account for activity-dependent accumulation of molecular cargo at synaptic sites in realistic neuronal morphologies. Within this framework, we identify several ways in which local biochemical mechanisms can be orchestrated to produce arbitrarily complex distributions of molecular cargo. In addition to addressing the logistical challenges of dendritic transport, we identify several constraints the precision, flexibility, and speed of cargo delivery inherent to this model.

RESULTS

Model description

Transport along microtubules is mediated by kinesin and dynein motors, which mediate anterograde and retrograde transport, respectively (Hirokawa et al., 2010; Gagnon and Mowry, 2011). Cargo is often simultaneously bound to both forms of motor protein, resulting in stochastic back-and-forth movements with a net direction determined by the balance of opposing movements (Hancock, 2014; Buxbaum et al., 2014a) (Fig. 1A).

To obtain a general model that can accommodate variations in the biophysical details, we consider microtubule-based transport as a biased random walk along a one-dimensional cable (Bressloff, 2006; Newby and Bressloff, 2010; Bressloff and Newby, 2009). For each time step, the cargo moves left, right, or remains the in the same place. In the simplest model, the probabilities associated with each movement are fixed and independent across each time step (Fig. 1B, top panel). A potentially more realistic model incorporates *hysteresis* into the biased walk (see Methods). In this case, the cargo is more likely to continue in the same direction as it moved on the previous time step (Fig. 1B, bottom panel; see, e.g., Soundararajan and Bullock (2014)).

While the position of individual cargoes can be highly stochastic, the net movement of a population of cargoes (Fig. 1C) is predictable. Figure 1D shows the distribution of 1000 molecules over time with (top panel) and without (bottom panel) hysteresis. In both cases, these simulations can be well-fit by an convection-diffusion partial differential equation (Fig. 1E-F), which is easier to analyze because it is

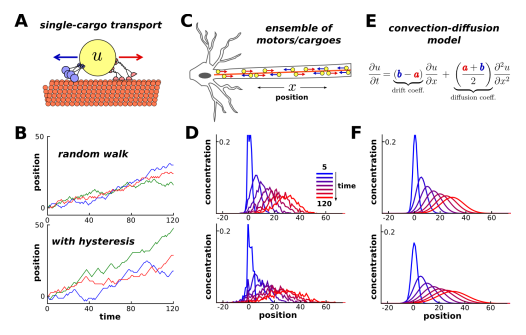


Figure 1. A simple mass-action model of dendritic transport

deterministic and contains fewer variables (Smith and Simmons, 2001).

The convection-diffusion equation can be numerically simulated by discretizing a dendritic tree into small compartments, and considering the movement of cargo between neighboring compartments as a chemical reaction with first order kinetics. In an unbranched cable with *N* compartments, the mass-action model is:

$$u_1 \stackrel{a_1}{\overleftarrow{b_1}} u_2 \stackrel{a_2}{\overleftarrow{b_2}} u_3 \stackrel{a_3}{\overleftarrow{b_3}} \dots \stackrel{a_{N-1}}{\overleftarrow{b_{N-1}}} u_N \tag{1}$$

where u_i is the amount of cargo in each compartment, and a_i and b_i respectively denote local rate constants for anterograde and retrograde transport; these parameters determine the local diffusion and drift coefficients as shown in figure 1E (see Methods). Similar mass-action schemes can be constructed to model transport in branched morphologies (Fig. 2A).

MOVE THIS PARAGRAPH TO METHODS/DISCUSSION? A central assumption of this model is that the movement of molecules between neighboring compartments is a stochastic and memoryless process. This assumption is reasonable if the run length of molecular motors is comparable to the size of the compartments; this appears to be the case under many (Muller et al., 2008; Verbrugge et al., 2009), thought not necessarily all (Dynes and Steward, 2007) circumstances (see Discussion). In simulations, we observed that this model was not very sensitive to violations of this assumption; the population dynamics of cargoes with substantial run lengths could still be well-fit by this class of models (Fig. 1F, lower panel; Supplementary Fig?).

A simple rule achieves desired spatial distributions of cargo

The exchange of u between neighboring compartments approaches an equilibrium/steady-state (ss) distribution over time. In an unbranched cable, this occurs precisely when:

$$\frac{u_i}{u_{i+1}}\bigg|_{SS} = \frac{b_i}{a_i} \tag{2}$$

(see Methods). Thus, the local trafficking rates, a_i and b_i , can be tuned to produce a desired spatial distribution of cargo. Let \tilde{u} denote the desired steady-state level of u in each compartment. For example, \tilde{u}_i might refer to the local concentration of biochemical signal that "tags" synapses for plasticity (Frey and Morris, 1997); such a tag might be positively correlated to the desired level of delivered cargo. To match this desired distribution, the ratio of local trafficking rates should satisfy:

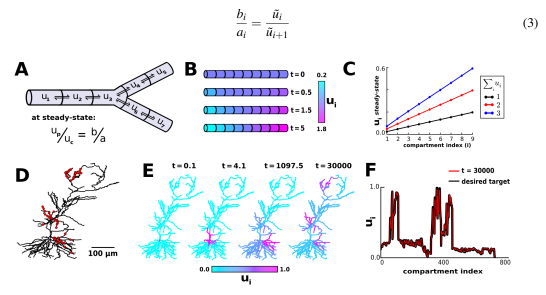


Figure 2. Delivery of cargo to active synapses in a complex dendritic tree

For example, a linear expression pattern emerges when $\tilde{u}_i \propto i$, resulting in $b_i/a_i = i/i + 1$ (Fig. 2B). This rule produces the expected linear expression gradient with the slope controlled by tuning the total amount of cargo in the cable (Fig. 2C).

These results extend directly to branched morphologies (see Methods). For any pair of connected compartments:

$$\frac{b}{a} = \frac{\tilde{u}_p}{\tilde{u}_c} \tag{4}$$

where u_p is the level in the "parent" compartment (closer to soma) and u_c is the level in the "child" compartment (closer to periphery); b and a refer to the local transport rate constants between this pair of compartments.

MOVE THIS PARAGRAPH TO DISCUSSION OR DELETE? These rules could be locally implemented by simple biochemical reaction networks. Consider the cytoplasmic calcium concentration, $[ca]_i$, as a candidate second messenger. Increases in $[ca]_i$ simultaneously arrest anterograde and retrograde microtubular transport (Wang and Schwarz, 2009). That is, for any pair of compartments, the anterograde rate constant is determined by calcium in the parent compartment, $a = f([ca]_p)$, and the retrograde rate constant is determined by calcium in the child compartment, $b = f([ca]_c)$, where f is a function that describes how calcium alters the transport rates. This results in:

$$\frac{b}{a} = \frac{f([ca]_c)}{f([ca]_p)} = \frac{\tilde{u}_p}{\tilde{u}_c} \tag{5}$$

where $\tilde{u}_i = 1/f([ca]_i)$.

To test the plausibility of this model in a complex morphology, we extended an existing multicompartment model of a CA1 pyramidal neuron (Migliore and Migliore, 2012). Excitatory synaptic input was delivered to 120 locations within three dendritic regions (red dots, Fig. 2D), and the average membrane potential in each electrical compartment determined the target level \tilde{u}_i in each compartment (Methods). This models how molecular cargo could be selectively trafficked to active synaptic sites. Figures 2E and 2F confirm that the spatial distribution of u approaches the desired steady-state.

Convergence rate

Biochemical processes are time-sensitive. For example, newly synthesized proteins must be delivered to synapses within \sim 1 hour to support long-term potentiation in CA1 pyramidal cells (Frey and Morris, 1997, 1998). It is therefore of substantial interest to examine factors that influence the speed of transport in the model.

In equations (4) and (5), we implicitly required each $\tilde{u}_i > 0$ in order to avoid division by zero. Intuitively, if $\tilde{u}_i = 0$ for some compartment, then no cargo can flow through that compartment, cutting of more peripheral compartments from the transport system. Similarly, if certain \tilde{u}_i are nearly equal to zero, then transport through these compartments will act as a bottleneck for transport, and convergence to the desired distribution will be slow.

Figure 3 illustrates and analyzes this intuition in a simple three compartment model. The two compartments on the end of the cable have the same desired level, $\tilde{u}_1 = \tilde{u}_3$; the middle compartment acts as a bottleneck when \tilde{u}_2 is very small (Fig. 3A). This results in the mass-action model:

$$u_1 = \frac{\varepsilon}{\frac{\varepsilon}{x}} u_2 = \frac{x}{\varepsilon} u_3 \tag{6}$$

We assume that u begins at one end of the cable, and examine the how taking ε to zero affects the convergence rate. Figure 3B shows that the convergence rate slows dramatically as ε decreases. This can be can be analyzed using basic linear systems theory; the convergence rate is given by the non-zero eigenvalue with smallest magnitude, which can be thought of as a rate-limiting step or process for the system (Methods, Supp. Fig. 1). Simulations confirmed this analysis and showed that the convergence rate diverges to infinity as ε approaches zero (Fig. 3C).

Qualitatively similar results were obtained in the multi-compartment CA1 neuron model. The model converged to a uniform target distribution more quickly than to a "bottleneck" target distribution, in which the middle third of the apical dendrite had low steady-state levels of cargo (Fig. 3D). Each pair of anterograde and retrograde rate constants were normalized to sum to one; thus, differences in convergence were not due to the scale of the trafficking rate constants.

In addition to this global view of convergence (Fig. 3D), we considered how the transport bottleneck affected transport to individual dendritic compartments. Consider a scenario where transported cargo produces a local chemical reaction after a critical concentration threshold is reached; for example, a

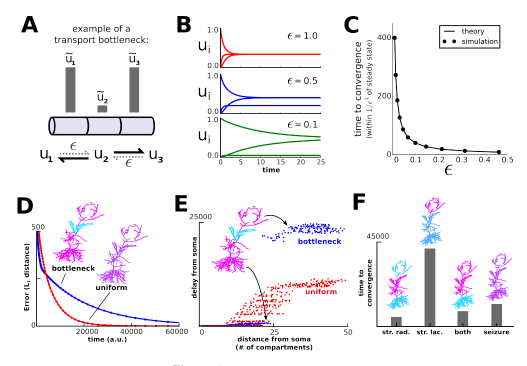


Figure 3. Bottlenecks in transport

recently potentiated synapse might be stabilized after enough plasticity-related factors are delivered from the soma. Figure 3E plots the transport delay, or duration of time it took for u_i to reach a pre-defined threshold for each compartment. We observed qualitatively similar results for different threshold values (data not shown).

As expected, introducing a bottleneck caused much longer delays to compartments distal to that bottleneck (Fig. 3E, upper right portion of plot). Interestingly, the presence of a bottleneck actually *shortened* the transport delay to proximal compartments, compared to the uniform target distribution (Fig. 3E, lower left portion of plot). This occurs because molecules that were delayed from entering the distal compartments spread throughout the proximal compartments, reaching higher levels earlier in the simulation.

One non-obvious prediction of this analysis is that transport to distal compartments should be quickened by allowing higher levels of u into the proximal compartments at steady-state (Fig. 3E). This might be tested experimentally by characterizing the convergence time of a retrogradely transported molecule that aggregates at recently activated synapses, such as Arc mRNA (Steward et al. (1998), see discussion). To illustrate this in the model CA1 cell, we characterized the time course of transport to the distal apical dendrite (stimulating stratum lacunosum/moleculare), proximal apical dendrite (stimulating stratum radiatum), entire apical dendrite (stimulating both layers), and entire cell (seizure). Interestingly, the model converged more slowly to distal input alone, than to paired distal and proximal input, or to an entirely uniform distribution (Fig. 3F).

Microtubular trafficking, detachment and degradation on separated time scales

The models above present a simple hypothesis for controlling molecular transport along an intricate system of microtubules. While certain types of molecular cargo stay on the microtubule network (e.g. mitocondria), others detach from the microtubules after a reaching a desired location. For example, dendritic mRNAs are transported within densely-packed granules, and can be locally released following granule disassembly (Krichevsky and Kosik, 2001; Buxbaum et al., 2014b). We model the detachment from the microtubule network as an irreversible process, and include a compartment-specific degradation

reaction. This results in the following mass-action model:

As before, a molecule u is transported along a network of microtubules (top row, in equation 6). In each compartment, molecules can irreversibly detach from the microtubules in a reaction $u_i \stackrel{c_i}{\longrightarrow} u_i^*$, where u^* denotes the biochemically active form of u, which is assumed to be inactive during transport. The final reaction, $u_i^* \stackrel{d_i}{\longrightarrow} \varnothing$, models degradation in each compartment. Note that only u^* is subject to degradation; the molecule is assumed to be protected from degradation during transport. This model can be extended to branched morphologies (Fig. 3A).

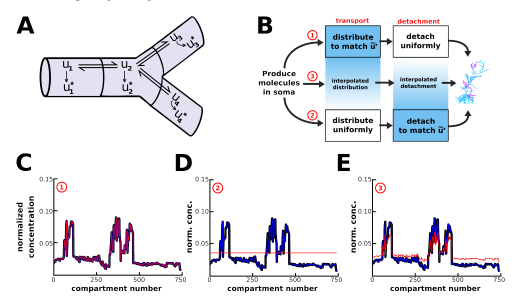


Figure 4. Multiple strategies for transport in a model including nonuniform microtubular detachment/activation

One way to analyze this system is to assume that these three processes — trafficking, detachment, and degradation — occur on separate timescales. If trafficking is sufficiently faster than detachment $(a, b \gg c)$, then u approaches a quasi-steady state distribution defined by our previous analysis (equation 5). We then choose detachment rate constants that transform the microtubular distribution into our desired distribution for u^* :

$$c_i \propto \frac{\tilde{u}_i^*}{\tilde{u}_i}$$
 (8)

Here, \tilde{u} and \tilde{u}^* respectively denote the quasi-steady state distributions for u and u^* , respectively. Due to the degradation reaction, the system no longer approaches a steady-state; however, as long as degradation is sufficiently slow $(c \gg d)$ the desired distribution is transiently achieved.

This model admits a spectrum of strategies for controlling the distribution of molecular cargo (Fig. 4B). One strategy (Fig. 4C) is to choose microtubular trafficking rates such that \tilde{u} exactly matches \tilde{u}^* , and choose the detachment rate constants to be spatially uniform (c_i = constant). Figure 4C shows that the desired spatial distribution is achieved both along the microtubules (red line) and for the detached/activated cargo (blue line).

Figure 4D shows the strategy on the other end of the spectrum. It begins by uniformly distributing cargo along the microtubules, and locally detaches cargo at a rate proportional to the desired level $(c_i \propto \tilde{u}_i^*)$. Unlike the first solution, this strategy avoids the transport bottlenecks examined in Figure 3, and can achieve target patterns with \tilde{u}^* equal to zero in certain compartments by setting $c_i = 0$.

Figure 4E shows the behavior of an intermediate model, whose parameters are a linear interpolation between the extreme strategies shown in Figure 4C and 4D. Thus, effective trafficking systems can be achieved by a spectrum of strategies, which may be suited to different situations and purposes (see Discussion).

Non-specific cargo delivery occurs when time scales aren't separated

We have thusfar considered models that produce a desired distribution of cargo to arbitrary precision. However, real neurons might tolerate some sloppiness in synaptic trafficking (or even utilize it as a substrate for complex plasticity rules). Thus, we examined the consequences of relaxing the separation of time scales between transport and detachment. In general, this improves the speed at which cargoes are delivered to synapses, but decreases the accuracy or specificity of delivery.

To illustrate this, we considered an experiment in which cargo is delivered to recently activated synaptic sites.

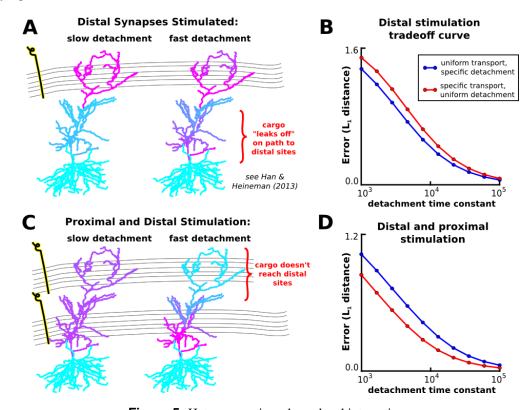


Figure 5. Heterosynaptic and non-local interactions.

Tradeoff between speed of consolidation and specificity of potentiation

We sought to understand the general relationship

DISCUSSION

A microscope image of a typical neuronal dendritic tree hints at the complex logistical task that the neuron must solve to maintain synapses and alter them, as appropriate, during development and learning. Synapses, such as the extensively-studied excitatory synapses of the mammalian nervous system, comprise hundreds of different proteins, including receptors, structural and anchoring proteins, and cytosolic

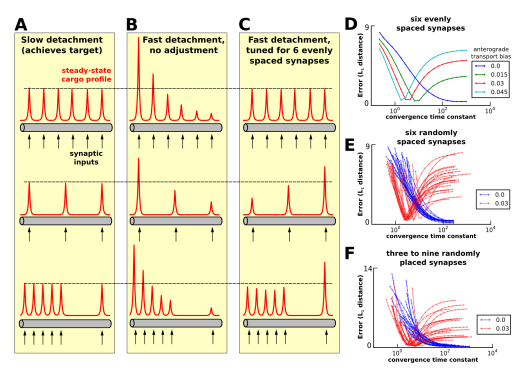


Figure 6. Tradeoff between speed of consolidation and specificity of potentiation.

signalling enzymes [REFs]. How do these components find their way to the synapse, and how do synapses that have varying demands for these components receive an appropriate supply?

Local protein synthesis near synaptic sites offers an elegant solution to this problem. Many mRNAs for synaptic proteins are distributed in dendrites, along with ribosomes and translational regulation machinery that can provide new proteins as they are needed [REFs]. However, these components need to be appropriately distributed to begin with and on a long timescale their distribution needs to be maintained. On the other hand, some forms of long-term synaptic plasticity have a transcription-dependent component, suggesting that somatic biosynthesis and dendritic trafficking are important for maintaining and controlling synaptic connectivity in addition to local mechanisms.

We examined simple models of microtubular transport to investigate how such a global network can be organized and controlled in complex neuronal morphologies. Previous models of microtubular-based transport have examined systems with spatially uniform rates of anterograde and retrograde transport (Smith and Simmons, 2001; Bressloff, 2006). We hypothesized that complex spatial patterns could emerge by relaxing this assumption. One might naïvely expect that elaborately branched morphologies obfuscate the relationship between these local mechanisms and the global distribution of molecular cargo. We showed that the relationship is actually quite simple: the steady-state ratio between neighboring compartments is the reciprocal ratio of the trafficking rate constants connecting those compartments.

Local activity signals provide a means to sequester dendritic cargo as it passes a synapse, providing a way for plasticity-inducing synaptic stimulation to 'tag' a synapse. Direct evidence of activity-dependent arrest (Soundararajan and Bullock, 2014) and accumulation (Krichevsky and Kosik, 2001; Buxbaum et al., 2014b) of mRNAs and other actively transported biomolecules suggests that the kind of mechanism we studied here is involved in regulating synapses in parallel with local synthesis mechanisms. We are agnostic about the precise identity of the activity signal that controls transport and sequestration, but note that experiments have shown that local calcium (Wang and Schwarz, 2009) and ADP (Mironov, 2007), alter anterograde and retrograde transport rates. Similarly, experimental data indicate that degradation of mRNA can be spatially heterogeneous and activity-dependent (Farris et al., 2014). Incorporating these additional processes into the model created new possibilities for molecular trafficking; the same expression pattern can be achieved by nonuniform microtubular transport followed by uniform release or from uniform transport followed by nonuniform release (Fig. 4).

There is experimental evidence for both strategies. Kim and Martin (2015) identified three mRNAs that were uniformly distributed throughout cultured *Aplysia* sensory neurons, but were targeted to synapses at the level of protein expression by localized translation. In contrast, the nonuniform expression of *Arc* mRNA is closely matched to the pattern of Arc protein (Farris et al., 2014; Steward et al., 2015).

Each of these transport strategies might be tailored for different constraints or purposes. For example, a uniform mRNA profile could facilitate shifts in the protein expression pattern, since the changing the pattern of local translation is presumably faster than re-distributing the mRNA. In other cases, it might be undesirable to rapidly shift protein expression based on transient changes to the neuron or circuit; controlling expression at the level of mRNA might prevent this. We showed that it is possible to interpolate between these two strategies (Fig. 4E), which might allow neurons to trade off between the relative advantages of each.

FRX - in this disease there is aberrantly high translation of mRNAs. This is like a high sequestration rate. Leads to higher connectivity but perhaps lower specificity.

Other assumptions of the mass-action model - Second, the model assumes are many copies of u in each compartment. This allows us to approximate the mass as a continuous variable, and the dynamics of the system as deterministic. Third, the number of molecular motors and capacity of the microtubules is not saturated by the level of u in any compartment. This means that the rate of transport from compartment A to compartment B linearly increases with the amount of u in compartment A.

METHODS

Microtubule Transport Model (Fig. 1)

Transport in an unbranched cable (equation 1) leads to the following system of differential equations:

$$\begin{array}{rcl} \dot{u}_1 & = & -a_1u_1 + b_1u_2 \\ \dot{u}_2 & = & a_1u_1 - (a_2 + b_1)u_2 + b_2u_3 \\ & \vdots \\ \dot{u}_i & = & a_{i-1}u_{i-1} - (a_i + b_{i-1})u_i + b_iu_{i+1} \\ & \vdots \\ \dot{u}_N & = & a_{N-1}u_{N-1} - b_{N-1}u_N \end{array}$$

Similar systems result from branched mass-action models. The steady-state relationship in equation 5 can be immediately verified at either end of the cable:

$$\dot{u}_1 = 0 \Rightarrow -a_1 u_1 + b_1 u_2 = 0 \Rightarrow \frac{u_1}{u_2} \bigg|_{ss} = \frac{b_1}{a_1}$$

The same procedure can be iteratively applied down the cable to show that equation 5 holds globally. The same result can be shown in a branched tree morphology. Here, each compartment has exactly one parent, but can have multiple children at branch points. Consider a parent-child pair of compartments, u_p and u_c , with anterograde/retrograde transport rate constants a and b. At steady-state, there is no net flux between u_c and its children:

$$\dot{u}_c = au_p - bu_c + \underbrace{\sum_j (b_j u_j - a_j u_c)}_{=0, \text{ at ss}}$$

Thus, we have:

$$\dot{u}_c = au_p - bu_c = 0 \Rightarrow \left. \frac{u_p}{u_c} \right|_{ss} = \frac{b}{a}$$

To study how the system converges to steady-state, we can re-express the model as a matrix differential equation, $\dot{\mathbf{u}} = A\mathbf{u}$, where $\mathbf{u} = [u_1, u_2, ... u_N]^T$ is the state vector, and A is a matrix. For a general branched morphology, the matrix A will be nearly tridiagonal, with off-diagonal elements corresponding to branch points; matrices in this form are called Hines matrices (Hines, 1984). For the simpler case of an unbranched

cable, A is tridiagonal:

$$A = \begin{bmatrix} -a_1 & b_1 & 0 & \dots & 0 \\ a_1 & -b_1 - a_2 & b_2 & 0 & & & \\ 0 & a_2 & -b_2 - a_3 & b_3 & \ddots & \vdots \\ \vdots & 0 & a_3 & \ddots & & 0 \\ & & \ddots & & -b_{N-2} - a_{N-1} & b_{N-1} \\ 0 & & \dots & 0 & a_{N-1} & -b_{N-1} \end{bmatrix}$$

For both branched and unbranched morphologies, each column of A sums to zero, which reflects conservation of mass within the system. The rank of A is exactly N-1 (this can be seen by taking the sum of the first N-1 rows, which results in -1 times the final row). Thus, the nullspace of A is one-dimensional (red lines in Supp. Fig. 1).

The desired distribution, $\tilde{\mathbf{u}}$, is an eigenvector that spans the nullspace of A. All other eigenvalues A are negative, which can be shown using the Gershgorin circle theorem. The convergence rate is determined by the non-zero eigenvalue with smallest magnitude of A. We can increase the convergence rate (while still maintaining $\tilde{\mathbf{u}}$) by multiplying all rate constants by some common value. However, this trivial strategy ignores the fact that there are biophysical limits on how fast molecular motors can move.

Stochastic Model of Transport

We model the stochastic transport of a single molecule with a discrete-time Markov chain model. Let the vector $\mathbf{p}^{(k)} = [p_1^{(k)}, p_2^{(k)}, ..., p_N^{(k)}]^T$ denote the probability of finding the molecule in each of the N compartments on the kth time step. For example, $p_1^{(0)}$ is the initial probability that the molecule is in the first compartment (which is the soma in our implementation). The state-transition matrix S describes the probabilistic anterograde/retrograde transport of the molecule between neighboring compartments in each discrete time step:

$$\mathbf{p}^{(k+1)} = S\mathbf{p}^{(k)}$$

The state-transition matrix is related to the mass-action system matrix A by the following relation:

$$S = \Delta t A + I$$

where Δt is the length of a time step (assumed to be very small), and I is the identity matrix. The columns of S sum to one, and, since Δt is small, all elements of S lie on the interval [0,1]. This transformation shifts and scales the eigenvalues of A to lie on the interval [0,1], without modifying the eigenvectors of the system. The steady-state probability distribution is given by the eigenvector of S with eigenvalue 1 (CITE), which is identical to the vector spanning the nullspace of S and therefore the steady-state concentration distribution of the mass-action model.

0.1 Transport Model With Detachment and Degradation (Fig. 4)

We can also interpolate between these two strategies to achieve the target distribution. Let F be a scalar between 0 and 1, and let \tilde{u}^* be normalized to sum to one. We choose a_i and b_i to achieve:

$$\tilde{u}_i = F \ \tilde{u}_i^{\star} + (1 - F)/N$$

along the microtubular network and choose c_i to satisfy

$$c_i \propto \frac{\tilde{u}_i^{\star}}{F \ \tilde{u}_i^{\star} + (1 - F)/N}$$

Here, N is the number of compartments in the model. Setting F = 1 results in the simulation in figure 4C, while setting F = 0 results in the simulation in figure 4D. Setting F = 0.3 results in the simulation shown in figure 4E, which uses an interpolated strategy.

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