Compact Modeling of Allosteric Multisite Proteins: Application to a Cell Size Checkpoint

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Abstract

We explore a framework to model the dose response of allosteric multisite phosphorylation proteins using a single auxiliary variable. This reduction can closely replicate the steady state behavior of detailed multisite systems such as the Monod-Wyman-Changeux allosteric model or rule-based models. Optimal ultrasensitivity is obtained when the activation of an allosteric protein by its individual sites is concerted and redundant. The reduction makes this framework useful for modeling and analyzing biochemical systems in practical applications, where several multisite proteins may interact simultaneously.

As an application we analyze a newly discovered checkpoint signaling pathway in budding yeast, which has been proposed to measure cell growth by monitoring signals generated at sites of plasma membrane growth. We show that the known components of this pathway can form a robust hysteretic switch. In particular, this system incorporates a signal proportional to bud growth or size, a mechanism to read the signal, and an all-or-none response triggered only when the signal reaches a threshold indicating that sufficient growth has occurred.

Introduction

Protein phosphorylation is a common form of post-translational modification frequently used in nature to alter protein activity, for instance by changing the electrostatic properties of the protein or its spatial structure. The phosphorylation of the same protein at multiple different aminoacid residues is also very common, and it is found in proteins such as p34cdc25 [1], Sic1 [2], EGFR [3], Wee1 [4], Ste5 [5] and many others [6].

The differences in the function of single-site vs. multisite phosphorylation are not completely understood. Many multisite proteins are involved in regulatory processes that can benefit from the presence of bistability, hysteresis, or limit cycles, which require sufficiently nonlinear interactions in addition to the right type of feedback [7,8]. A reasonable hypothesis is that multisite phosphorylation can give rise to ultrasensitive dose responses, in a way that would not be possible in a comparable single-site system [9–13]. Many detailed mechanisms have also been proposed to explain the role of multisite systems in the emergence of bistability (for examples, see [14,15]).

On the other hand, such detailed multisite mechanisms are normally not used as part of actual mathematical models of biochemical interactions. This is because explicitly modeling multiple sites usually involves the introduction of numerous variables, one or more for each phosphorylation state, and realistic models are often too complex already to justify this additional effort. Systems that attempt to model biochemical reactions explicitly often use the assumption that the protein has two states, one active and one inactive, with a simple reaction to transform one into the other, effectively assuming that the protein only has one site. Other models are more phenomenological in nature and include, for example, Hill function terms in the equation that are less clearly tied to the actual biochemical reactions [16,17].

In this paper we describe a simple mechanistic approach for modeling multisite allosteric proteins. This approach, named modified fraction (MF) modeling, is capable of describing ultrasensitive dynamics without introducing a large number of additional variables. Under this framework one keeps track of the fraction of modified sites in the protein, and the concentration of active protein over time is estimated from this information. The protein is activated in a way that requires the phosphorylation of several but not all of the sites. The approximation becomes increasingly precise as the number of sites increases, with good estimates already for around four or more sites. In a sense, this mechanism can be considered a one-variable, quasi-steady state reduction of a model similar to the Monod-Wyman-Changeux allosteric system [18], although uses of MF modeling outside of MWC are also possible. The MF framework can also be extended to other types of multisite modification such as ligand binding, multisite transcription factor regulation, multisite methylation or acetylation, ubiquitination, etc [19,20].

Perhaps the best way to test the versatility of a modeling tool such as the one proposed is to implement it in an actual biochemical system. In the current work we describe a detailed mechanistic model of a cell size checkpoint. Cell size checkpoints halt the cell cycle at specific points until sufficient cell growth has...
Compact Multisite Modeling of Cell Size Checkpoint

Author Summary

A large number of proteins in the cell are modified post-translationally by phosphorylation at multiple specific locations. This can help bring about interesting dynamical behaviors such as bistability or all-or-none responses to stimuli. Such behaviors are in turn important for cellular decision-making, differentiation, or the regulation of cellular processes. In this paper we propose the use of a specific technique for modeling allosteric multisite proteins, which can be thought of as the reduced version of a more detailed set of reactions. After validating this technique computationally by comparison to more detailed systems, we apply it to a new model of a signal transduction pathway with several multisite proteins. The model is concerned with a mechanism in budding yeast that is thought to measure the extent of daughter bud growth and send a signal to initiate mitosis when sufficient growth has occurred. Using the given framework we derive an analytically tractable model that creates the desired all-or-none signal. Overall, we give quantitative support to a newly discovered biochemical pathway, and we show the utility of the new modeling framework in the context of a realistic biological problem.

occurred [21,22]. The mechanisms by which cell size checkpoints operate are poorly understood, and it is unclear whether they monitor actual cell size or parameters more closely related to the extent or rate of growth. In budding yeast, growth of a new cell is initiated when a daughter bud is formed on the surface of the cell [23]. The daughter bud initially grows in a polar manner, with all growth directed to the bud tip. Growth of the bud eventually switches to isotropic growth, in which the bud grows over its entire surface (see Figure 1A) [24]. The timing of the switch determines the duration of polar growth, which influences cell size and shape. It has been proposed in recent work by one of the authors that a cell size checkpoint controls the timing of this switch [25]. This checkpoint is the subject of our model. The variables are illustrated in Figure 1B and described in more detail below. See Tyson and Novak [26] for an accessible introduction to the systems-wide modeling of cell cycle checkpoints.

The MF approximation equation was first developed in [27], in the context of multisite systems with independent modification sites, with an emphasis on the estimation of the Hill exponents of sequential and nonsequential systems and on the comparison of their qualitative behavior. A major advance of the current paper, beyond the application to the cell cycle checkpoint, is to extend their qualitative behavior. A major advance of the current paper, beyond the application to the cell cycle checkpoint, is to extend this work to cooperative and allosteric systems. Such systems are by definition non-independent, since the modification of one site accelerates the rate of modification of its neighbors. Cooperative systems are also more common and much better characterized than independent ones. The validation of the approximation in cooperative systems is ultimately based on a computational comparison of the MF reduction with detailed cooperative models having n or 2^n variables.

In the first two Results sections we carry out a description of the modified fraction method to model multisite systems, and we compare simulations of the reduced model with those of a detailed mechanistic model. In the remaining two Results sections we carry out a mathematical analysis of the proposed checkpoint signaling pathway. We hypothesize that this interaction pathway has the capacity to produce a bistable signal responsible for a sudden switch from polar to isotropic growth, once the bud has undergone sufficient polar growth. The model presents several desirable qualities for a checkpoint, in particular a clear downstream signal when a sufficient polar bud growth has occurred.

Results

The MF framework describes a compact multisite mechanism

We start by describing the assumptions on our model in the context of multisite phosphorylation (although it could also be applied to other irreversible covalent modifications as well as noncovalent ligand binding). Suppose that a protein substrate \( P \) is phosphorylated by a kinase \( E \) at \( n \) possible sites and dephosphorylated by a phosphatase \( F \). The system is assumed to be nonsequential, so that there are \( 2^n \) different phosphoforms of \( P \), and the number of sites is thought to be relatively large e.g. \( n \geq 5 \). The system is cooperative in the sense that site phosphorylation accelerates the phosphorylation of neighboring sites. Since the number of sites is relatively large, the activation is thought to be cumulative and the effect of any individual site is assumed to be small. The sites are assumed to be equivalent to each other, in the
sense that the rate of phosphorylation and dephosphorylation is similar across all sites and that no site has a stronger effect on substrate activation than other sites. The activation of the substrate may be due to binding to another molecule or body, such as the cell membrane. It could also be due to an internal structural change that allows the substrate to interact differently with other proteins. Thus the active protein concentration can be defined as the concentration of the protein bound to a particular molecule or in a particular molecular state.

Suppose that the phosphorylations lead to the concerted, redundant activation of the protein. That is, multiple phosphorylations are necessary for activation (concerted), and not all sites need to be phosphorylated for full activation (redundant). We define the activity function \( h : [0,1] \rightarrow [0,1] \) such that the fraction of active protein with \( i \) phosphorylated sites is given by \( h(i/n) \).

There are many systems that likely fall within this general framework. For instance, Ste5 is a scaffold protein in budding yeast with \( n = 8 \) phosphorylation sites, which relays a pheromone response only when it is bound to the membrane [28,29]. When phosphorylated by Cdk1, it tends to unbind from the membrane, shutting down its activity. The sites are predicted to lie on an unstructured region of the protein and appear to act by changing the protein’s bulk electrostatic properties. In the paper [29], it was shown through site-directed mutagenesis that around five or more phosphorylations are necessary and sufficient for deactivation. Another recent example is the multisite phosphorylation of Cdc25 by Cdk1 in fission yeast, which was similarly studied in detail by mutating individual sites [30].

According to the modified fraction framework, we estimate the concentration of a particular protein state from the overall fraction of modified sites. For instance, if the protein has \( n = 3 \) sites and the fraction of phosphorylated sites is \( p \), then the fraction of protein with only the first and last sites phosphorylated is roughly

\[
P_{101} \approx p^2 (1-p).
\]

Here \( P \) is the total protein concentration. This is not an equality since cooperative effects introduce correlations among the sites, i.e. the sites are not independent of each other, but it is an approximation assuming cooperative effects are sufficiently weak. Multiplying on both sides by \( P \) and adding over all possible phosphoforms with \( i \) phosphorylations, the concentration \( P_i \) of proteins with exactly \( i \) phosphorylated sites out of a total of \( n \) sites is

\[
P_i \approx P \binom{n}{i} p^i (1-p)^{n-i}.
\]

One can estimate the overall concentration \( P \) of active protein as

\[
P = \sum_{i=0}^{n} \binom{n}{i} P_i \approx \sum_{i=0}^{n} \binom{n}{i} \binom{n}{i} (1-p)^{n-i}.
\]

A key aspect of this formula is that if we denote the right hand side by \( f_a(p) \), it can be shown that \( f_a(p) \) converges to \( Ph(p) \) as \( n \) increases [31]. This gives the approximation

\[
P \approx Ph(p), \quad (1)
\]

which becomes increasingly precise for large \( n \). Notice that the different quantities in this formula can potentially be measured in the lab - the active protein concentration via an activity assay, the total protein concentration via Western blot, and the activity function \( h(x) \) through site-directed mutagenesis.

A timescale decomposition argument can be made to use this approximation away from steady state. If \( p(t) \) is the fraction of active sites over time, and the timescale of protein activation is much faster than the rate at which \( p(t) \) changes, then one can approximate \( P(t) \approx Ph(p(t)) \) at any given time using the same formula. This produces a convenient method for modeling multisite systems under the given assumptions by keeping track of the variable \( p(t) \), without creating \( n \) let alone \( 2^n \) variables. On the other hand, if \( p(t) \) or any other process affecting protein activation, is at least as fast as protein activation itself, then nontrivial dynamics might take place such as limit cycle oscillations, and the approximation can introduce errors.

It is necessary to calculate the fraction of phosphorylation \( p \) itself. Assuming linear rates of phosphorylation and dephosphorylation for \( p \), one obtains the system

\[
p_{in} \frac{zE}{P} \frac{dP}{dt} = \beta \frac{P}{P} - \delta P,
\]

That is, \( dp/dt = zEP_{in} - \beta P \), where \( p_{in} = 1 - p \) is the fraction of inactive sites. This is the default for the model, although any other rate equation for \( dp/dt \) can be used, including Michaelis-Menten complex formation at the level of the individual sites.

As for the activity function \( h(x) \), any sigmoidal function can be used, including functions measured directly by experiments. By default we assume the following form, which we will derive in the next section:

\[
h(x) = \frac{e^x}{\delta + e^x}.
\]

See also [5,9,27] for other uses and derivations of this formula in the literature. The function \( h(x) \) can actually be highly switch-like for large \( \gamma \), which illustrates how small increases in the kinase \( E \) can result in large activity changes in the protein \( P \), unlike linear rate models with only one site. Other forms for the activity function \( h(x) \) have effectively been considered by Kapuy et al [32], and also by Wang et al [10].

In Figure 2A we show the relationship between the fraction of phosphorylated sites \( p \) and the active protein concentration \( P \) for a particular choice of the MF parameters \( x, \beta, \gamma, \delta \), using the approximation formula \( P \approx Ph(p) \) (and chosen to fit the detailed model described in the next section for \( n = 6 \)). Notice that the activation is concerted and redundant, in that a minimal threshold of phosphorylation is required for activation, and activation is reached for less than full phosphorylation.

The MF model reproduces the dynamics of detailed allosteric systems.

A validation of the performance of this model for \( n = 6 \) is now shown in Figure 2 in the context of a system similar to the classical and widely used Monod-Wyman-Changeux model of an allosteric multisite protein [18]. The original MWC model describes the binding of oxygen to the different sites of hemoglobin and the allosteric transitions of this protein between two different states. Rather than modeling oxygen binding, a protein with \( i \) phosphorylations is assumed to change between an active conformation \( A_i \) and an inactive conformation \( I_i \). Each of these forms can also be phosphorylated or dephosphorylated at the rates...
given in the diagram in Figure 2B. Although the model is interpreted in a different way from MWC, from a mathematical point of view it is simply assumed that the protein has a single site. Here \( n = 0, \ v = 0.1, P = 12 \text{ nM}, F = 1 \text{ nM}, L_1 = e^{-n/2},\ x = \beta = L_2 = 1 \) for the detailed model, and \( x = n/2, \beta = 1, \gamma = -n \) in \( x, \beta = L_1 \) for the MF approximation. D. Comparison of the error between the MF and the MWC model, for different values of \( n \) and \( \epsilon \) and the remaining parameters computed as above. E. Calculation of the Hill exponent of the MF model for different values of \( n, \epsilon \). In each case \( L_1 = e^{-n/2} \) to allow for half maximal activation with \( n/2 \) phosphorylations, and \( x = \beta = L_2 = 1 \). F. The default equations for the MF approximation. The active protein concentration \( P \) is a function of the fraction of active sites \( p \) and the total protein \( P \). The values of \( p \) are calculated via a simple chemical reaction, and a form for the function \( h(x) \) is suggested.

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At steady state, we assume that this exchange is balanced and calculate \( L_1L_2A_1 = L_2I_1 \). Then the fraction of active sites with \( i \) phosphorylations at steady state is

\[
h(i/n) = \frac{A_1}{L_1 + A_1} = \frac{L_1e^{-i}}{L_1e^{-i} + L_2} = \frac{e^{-i}}{L_1/L_2 + e^{-i}} = \frac{e^{-i/n\ln(\epsilon)}}{L_1/L_2 + e^{-i/n\ln(\epsilon)}}.
\]

In other words \( h(x) = e^{x^{\gamma}}/(\delta + e^{x^{\gamma}}) \), where \( \gamma = -n \ln \epsilon \) and \( \delta = L_1/L_2 \). See Text S1.3 for more details. In particular, the ultrasensitive behavior of the function generally increases with the number of sites.
At any given time, the active protein concentration of the full model is defined as $P_{MWC} = A_0 + \ldots + A_n$. In Figure 2C we compare the full 12-variable MWC model for $n=6$ with the corresponding MF approximation. For every value of the input kinase concentration, the resulting concentration of active protein $P$ is plotted at steady state. Notice the close similarity between the two graphs, which is even more surprising since MF is essentially a one-variable model.

For comparison, we also plot the behavior of an overly simplified but all too commonly used model, in which the substrate is assumed to have a single phosphorylation site instead of $n$ sites, and it is modeled according to the reaction

$$P_{\text{single site}} \xrightarrow{k_F} P_{\text{single site}}$$

using linear reaction rates. Notice that the behavior of this single-site model in Figure 2C is very different from that of the MWC model, and that any switch-like behavior in the response is lost. This can have important consequences regarding the existence of multiple steady states, hysteresis, oscillations etc in the context of larger systems, which will be illustrated below. It is easy to show that $P_{\text{single site}} \approx P_{\text{MF}}$, i.e. it corresponds to $P_{\text{MF}}$ when $h(x) = x$. It should be noted that if the single site system is modeled using Michaelis-Menten reactions rather than linear rates, it could have strongly ultrasensitive behavior in the saturation regime via zero-order ultrasmallity [33]; see the Discussion section for more details.

We carried out a calculation of the distance between $P_{\text{MF}}$ and $P_{\text{MWC}}$ for many different combinations of the number of sites $n$ and the allosteric parameter $e$. For every such set of parameters, the two graphs were plotted at steady state as a function of $E$, and the error $\max(E) | P_{\text{MWC}}(E) - P_{\text{MF}}(E) | / P$ was calculated in Figure 2D. Notice that the approximation is within 1% precision for arbitrary $n$ and $e>0.2$. On the other hand, in order to obtain high ultrasmallity it is required that $n$ be relatively large and/or $e$ be small (Figure 2E). See also Figure S2, where additional parameter variations are explored over four orders of magnitude using the same type of graphs, with similar results.

It is worth comparing this methodology with the approach known in the literature as rule-based modeling, where a series of chemical reactions is defined using a streamlined algorithm, and high-powered computing is used to handle the resulting large number of variables; see e.g. BioNetGen [34]. The advantage of this method is that a large number of reactions can be defined and handled this way, including complex parameter optimizations. One disadvantage is that the combinatorial explosion resulting from combining reactions can sometimes exceed the computational power. Another is that the large number of equations makes any mathematical analysis difficult, if at all possible.

It is interesting that the MWC model can actually be described in terms of rule-based modeling. In Figure S1A we describe a series of chemical reactions, over all possible phosphoform states, and we show in Text S1 that this system is in fact equivalent to the MWC model. Thus MF can also be seen as the 1-variable reduction of a system with $2^{n+1}$ variables and a much larger number of reactions.

Control theoretic analysis of the pathway reveals two robust switches

In this section, we will embed the MF system within increasingly complex systems of equations. We consistently use upper case for proteins and lower case for modified fractions of sites. However, we will first provide some technical experimental background regarding this specific pathway.

**Background: The cell size checkpoint pathway.** Any theory for cell size checkpoints should ideally account for several mechanistic features. First, cell size checkpoints should translate growth into a proportional checkpoint signal. Second, they should read the signal to detect when it reaches a threshold that indicates when sufficient growth has occurred. Third, when the threshold is reached the checkpoint should trigger a switch-like cell cycle transition. All of these mechanisms should be robust and adaptable to function in cells of diverse size and shape, and under conditions of fluctuating growth rates.

It was recently hypothesized that the timing of the polar to isotropic growth transition described in the introduction is controlled by a mitotic cell size checkpoint. Mitotic cell size checkpoints are controlled by Wee1 and Cdc25 [25]. Wee1 is a protein kinase that delays entry into mitosis by phosphorylating and inhibiting Cdk1, while Cdc25 is a phosphatase that promotes entry into mitosis by dephosphorylating Cdk1. The budding yeast homologs of Wee1 and Cdc25 are known as Swel and Mih1; however, for clarity we will use their more commonly known names. In this model, a checkpoint signal originates at the site of polar membrane growth in the bud, and downstream components read the strength of the signal and trigger activation of Cdk1 when it reaches a threshold, using the pathway described in Figure 1B. A switch-like increase in Cdk1 during entry into mitosis is postulated based on evidence of this qualitative behavior of Cdk1 in other systems. Also, blocking polar membrane growth causes a Wee1-dependent arrest before mitosis, which indicates that entry into mitosis is linked to membrane growth [25]. The timing of the switch from polar to isotropic growth determines the duration of polar growth, which influences cell size and shape. In this sense, it is a cell size checkpoint.

Signaling is thought to be initiated by delivery of the GTPase Rhod to the site of membrane growth. Rhod is delivered by vesicles that are sent to the bud and fuse with the membrane to increase its size [35]. Therefore the inflow of Rhod is proportional with the rate of polar membrane growth. Rhod on vesicles is inactive and it undergoes activation when vesicles fuse at the site of growth [35]. Active Rhod binds protein kinase C (Pkc1), a member of the protein kinase C family, and induces it to undergo autophosphorylation [36]. Importantly, phosphorylation of Pkc1 is dependent upon and appears to be proportional to membrane growth, which suggests that Rhod relays proportional checkpoint signals regarding the extent of membrane growth [25]. This scenario describes the upstream component of the pathway in Figure 1B.

Regarding the downstream component in Figure 1B, it is thought that hyperphosphorylation of Cdc25 by a poorly understood kinase inhibits its activity [37,38]. Thus, a key event necessary for initiation of early mitotic events is dephosphorylation of Cdc25. This dephosphorylation is carried out by the phosphatase PP2A (more precisely, the heterotrimeric complex PP2A Cbeta5, which includes the Cdc25 regulatory subunit [25]). PP2A is in turn activated by the upstream Pkc1 protein. Activation of PP2A is also dependent upon an as yet identified protein named Zds1 [25,38] (Zds1 is a redundant paralog named Zds2, which will be identified with Zds1 in the model). Zds1 forms a tight stoichiometric complex with PP2A [25]. It also binds to Pkc1, although that bond is not included explicitly in the model. PP2A is also believed to dephosphorylate Pkc1. Thus, inactivation of PP2A causes hyperphosphorylation of Pkc1 [25]. Although the functions of Pkc1 phosphorylation are not yet fully known, this observation suggests that PP2A restrains activation of Pkc1 by checkpoint
signals. Similarly, PP2A also dephosphorylates associated Zds1, which suggests that activation of PP2A by Pkc1 leads to dephosphorylation of Zds1 and activation of targeted dephosphorylation of Cdc25 [39]. In this way PP2A is at the heart of checkpoint signaling: it opposes phosphorylation of both Zds1 and Pkc1, and it is responsible for the critical step of dephosphorylating Cdc25. There is also good evidence that PP2A regulates Wee1, although it is less clear whether this function of PP2A is controlled by signals from Pkc1 [4,39].

A bistable switch between Zds1 and PP2A. We can now analyze a model involving two different multisite proteins, Zds1 and PP2A, each of them described according to the MF framework. This forms the downstream component of a larger model for the signal transduction of Rho1, and it is described inside the lower dashed rectangle in Figure 1B.

The activity of each of the two proteins can be modified through phosphorylation and dephosphorylation, and both proteins bind together to form a dimer. Only the dimer configuration of these molecules is active as an enzyme, and only to the extent that the Zds1 and PP2A components have been modified appropriately through phosphorylation or dephosphorylation. Moreover, as shown in the figure and justified by experimental data, the active dimer is itself involved in the modification of the Zds1 protein.

Denote by \( Z, P \) the total monomer concentration of Zds1 and PP2A respectively, regardless of their phosphorylation state, \( Z, P \) their active concentration, \( z, p \) the modified fraction of Zds1 and PP2A sites, and \( D \) the total Zds1/PP2A dimer concentration. Call \( C \) the active Rho1/Pkc1 dimer concentration, which can be treated as a constant input to the system for now. The dimerization reaction can be written as

\[
Z + P \xrightarrow{q_3} D.
\]

Now, following the argument in the previous section and given that both Zds1 and PP2A have been found to have 5 to 10 phosphorylation sites each, we keep track of \( z, p \) using the equations

\[
\begin{align*}
q_1 D & \xrightarrow{q_2} Z, \\
q_3 & \xrightarrow{q_4} P.
\end{align*}
\]

Since Zds1 is activated through dephosphorylation, here \( z \) represents the fraction of dephosphorylated Zds1 sites, while \( p \) represents the fraction of phosphorylated PP2A sites. Using the MF equation one can then estimate

\[
Z = Zh_1(z), \quad P = Ph_2(p).
\]

See Figure 3A for a graph of these two functions. The level of ultrasensitivity of each curve is dependent on the estimated number of sites in each protein (see the Methods section), but notice that the graphs are roughly consistent with the measurements done for example in the Ste5 protein. A simple mathematical analysis of this 3-dimensional model in Text S1.1 shows that the solutions of the system converge towards the steady states, and that the steady state equations can actually be reduced to a single equation for \( z \),

\[
\Gamma_1 z = h_2 \left( \frac{C}{C + q_3/q_1} \right) h_1(z)(1 - z),
\]

where \( \Gamma_1 \) can be calculated from the system parameters. For a given value of \( C \) this equation can have one or possibly three solutions, depending on the parameter values. Plotting the left and right hand sides separately can be helpful (Figure 3B). As \( C \) increases, the function \( h_1(z)(1 - z) \) is rescaled vertically, which can control the number of solutions and create a hysteretic response on the variable \( z \).

Given a solution for \( z \) in this equation one can solve for the concentration of \( Z, P \) and every other protein. In particular one can calculate the concentration of the active form of the Zds1/PP2A dimer which we call \( D \), and which constitutes the natural output of this system. For specific values of the parameters in the model, a bifurcation graph of the output \( D \) as a function of the input \( C \) can be found in Figure 3C (solid line).

One can interpret this graph in the following way. There is a positive feedback loop in the system consisting of Zds1 promoting its own activation via dimerization with PP2A. This positive feedback allows the possibility of two stable steady states in the system. When \( C = 0 \), there can be binding between Zds1 and PP2A but since PP2A is inactive, most of the PP2A/Zds1 dimer is inactive. For high values of \( C \), PP2A is forced to become active, however the PP2A/Zds1 dimer may still have two steady states or only one, depending on the parameters of the system.

A bistable switch between Rho1 and Pkc1. In a similar way as it was carried out for the interaction between Zds1 and PP2A, one can study the activation of Rho1 and its binding with Pkc1. This upstream system includes all the variables and interactions that are not in the Zds1/PP2A module. Once again following the biological evidence and the diagram in Figure 1B, we define the biochemical reactions

\[
Rin \xrightarrow{q_6} R, \quad \phi \xrightarrow{q_9} R_{in}, \quad R \rightarrow \phi,
\]

where \( R_{in}, R \) denote inactive and active Rho1 in monomer form respectively and \( \phi \) the rate of vesicle fusion to the membrane. That is, the protein Rho1 is activated at a linear rate, the vesicle flow increases Rho1 concentration in inactive form, and both active and inactive Rho1 are degraded. The protein Pkc1 is modeled using a MF mechanism and denoted by the variables \( k, K \).

\[
k \xrightarrow{q_5} k, \quad K = Kh_3(k).
\]

For the study of this subsystem, \( D \) will be considered a fixed input parameter; it constitutes the feedback from the downstream Zds1/PP2A system. Calling \( C \) the total Pkc1/PP2A dimer regardless of activity,

\[
R + K \xrightarrow{q_9} C.
\]

Notice that Rho1 needs to be active to bind with Pkc1. However active Rho1 binds with Pkc1 regardless of the activity or inactivity of the latter, Pkc1 does need to be active in order for the dimer to become active as a kinase, so that \( C/K = Kh_3(k) \), or
\[ C = Ch_3(k). \]

Once again, one can carry out a complete mathematical analysis of this multi-dimensional system at steady state and essentially reduce its solutions to one equation for \( C \). The resulting equation is

\[ \Gamma_2 C = \nu \left( S_K h_3 \left( \frac{C}{C + Q_5 D} \right) - C \right), \]

where \( Q_5 = q_{-5}/q_5 \), \( \Gamma_2 \) can be calculated from the parameters and constants of the system (other than \( \nu \), \( D \)), and \( S_K \) is constant. See Text S1.2 for a full analysis.

Figure 3D illustrates the solutions of the above equation in the general case, plotting right and left hand sides separately. For fixed \( \nu \) and \( D \) values, \( C=0 \) is always a solution, however there may also be solutions \( C<0 \). For increasing values of \( D \), the values decrease on the right hand side of the equation, and the steady state \( C<0 \) eventually disappears. Figure 3E illustrates a typical bifurcation graph of the steady states of \( C \) on the input parameter \( D \) and fixed \( \nu \). In the case \( D=0 \) there is actually only one solution for \( C \), since equation (3) becomes linear.

The bifurcation graph can be interpreted as follows. For all but small values of the PP2A/Zds1 dimer \( D \), there is the steady state with \( C=k=R=0 \), in which the pathway is inactive since the lack of active Rho1 inhibits the activation of Pkc1. For middle concentrations of \( D \), the system may be bistable, again due to the positive feedback from Rho1/Pkc1 to the activation of Pkc1. For high concentrations of \( D \), Pkc1 is almost fully inactivated by \( D \) and the pathway is shut down.

The Rho1 pathway can implement a cell size checkpoint

In order to find the steady states of both subsystems together, recall that each one can be reduced to a single equation, so that the steady states correspond to the joint solution of the two equations. For fixed \( \nu \), the solutions of the full model form the intersection of the graphs for the equations (2), (3). This is illustrated in Figure 3F, where the graphs in Figure 3C and Figure 3E are superimposed on the same plane. From a control perspective, the upstream and downstream systems have each an input and an output, and they feed back into each other (see the two dotted boxes in Figure 1B). The active Zds1/PP2A dimer \( D \) also acts as the overall output of the system, since it triggers the downstream response to cell cycle regulatory proteins. Although it is natural that an increase in the Rho1 flow can eventually trigger the activation of the pathway, the main focus here is not in the flow but in the overall Rho1 concentration at the bud tip. Given that Rho1 has a rate of growth proportional to \( \nu \) and a linear rate of degradation, at steady state one can show that \( \nu \) and \( R = R_{in} \) are proportional, \( R = R_{in} = \nu P/R_{-9} \). This follows from adding the ODE rate equations at steady state, \( 0 = R' + R_{in} + C' = q_{5} \nu = \nu P / R_{-9} \). In this way one can use the total Rho1 concentration \( R = R_{in} + R_{out} \) as a bifurcation parameter at steady state even though it is simultaneously a
variable in the system. Alternatively, since enzymatic reactions and dimer formation are fast processes compared with Rho1 flux and Rho1 degradation, one can let $R$ be the slow variable in the system and carry out a timescale decomposition analysis using $R$ as a constant [39].

Let's look at how the system has a hysteretic response for increasing values of the flow signal $v$ and the corresponding total Rho1 concentration at steady state. An increase in these values has the effect of raising the graph associated with the upstream system, as shown in Figure 4A. For smaller values of $R$, the intersection of both graphs includes three positive steady states (notice the two graphs don't quite intersect at the origin). But when $R$ increases over a certain threshold, the intersection of the two graphs contains a single positive steady state, with a large value of $D$. This can cause an abrupt change in the qualitative behavior of the system, triggering a sudden increase in the Zds1/PP2A output. Once this change has taken place, the concentration of the output stays high even if the input decreases.

Figure 4B shows a sample timecourse of the system for a time-variable vesicle flow (dotted line). The total Rho1 concentration increases over time with the inflow of vesicles. At a certain timepoint the active Rho1 concentration abruptly increases, due to the switch at the Pkc1/Rho1 upstream level. An increase in Rho1/Pkc1 concentration some time before this can be seen in Figure 4C. At a later time the switch between Zds1 and PP2A is also triggered, leading to a sudden increase in PP2A/Zds1 concentration. Even under variable flow, the total Rho1 concentration roughly corresponds to the membrane accumulated at the bud, except for a certain amount of variability due to Rho1 degradation. Lowering the Rho1 degradation rate can decrease this difference. Notice that the vesicle flow oscillations do not correspond to cell division, but to oscillations in the rate of growth, for instance due to varying food availability.

In Figure 4D we plot the output signal PP2A/Zds1 as a function of total Rho1 at steady state and overlay the solution of the timecourse simulation (red stars). This graph also illustrates the hysteretic behavior of the system, in that once a critical threshold of Rho1 concentration is reached, the output signal is dramatically increased. This change would constitute a clear signal that the bud has reached a large enough size for crossing the polar/isotropic growth checkpoint.

Since both the downstream (PP2A, Zds1) switch and the upstream (Rho1,Pkc1) switch are driven by positive feedback loops, it is valid to ask which of the two loops is more relevant for the overall system behavior. We argue that it is the downstream loop that is more essential, using the bifurcation analysis in Figure 3.

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**Figure 4. Checkpoint analysis.** A: Qualitative analysis of the steady states of the system for different values of the total Rho1 concentration $R$, as a result of changing the value of the fixed flux rate $v$. When $R$ reaches a sufficiently large value there is a single positive steady state and it has a large $D$ concentration. B,C: Response of the checkpoint pathway to a variable vesicle flow input (dashed line). For simplicity the membrane is modeled using the equation $m'(t) = v(t)$. D: The steady state values of the output $D$ as a function of $R$ (solid line), and plot of the timecourse in 4C over time (stars). doi:10.1371/journal.pcbi.1003443.g004
and Figure 4A. If the upstream system is not bistable but has a single steady state for every input \( D \), then the graph in Figure 3E is replaced by a single-valued decreasing function. Nevertheless this (green) line can still have one or three intersections with the (blue) downstream dose response in Figure 4A, indicating hysteresis for the overall system. On the other hand if the downstream system is not bistable, then the blue curve in Figure 4A is replaced by a single-valued, increasing function, which would be unlikely to have three intersection points with the (green) upstream dose response. Thus the downstream switch is essential, while the upstream switch is not.

Notice that this system contains the standard elements of a signal transduction pathway, including an initiating signal (vesicles), a sensor (Rho1), a series of transducers (Pkc1, PP2A, etc), and an effector (active Zds1/PP2A). Total Rho1 is a proxy for the membrane concentration, even if bud growth slows for a period of time, and the cascade of reactions allows the signal to be transduced from the membrane to Zds1/PP2A and ultimately the cyclin dependent kinase. To ensure the high fidelity of the signal transmission [40], the downstream signal is sent abruptly after total Rho1 concentration reaches a particular size. Notice that longer periods of inactivity can potentially reduce the Rho1 concentration significantly – one possible prediction is that after such a period the bud grows longer than expected.

Discussion

In this paper we have introduced a simple and compact framework to describe the dynamics of allosteric multisite phosphorylation systems, and we have applied this tool to a new molecular model of a size checkpoint in budding yeast. Multisite phosphorylation modeling can be problematic because ignoring the multiple sites can have significant effects in the dynamics, while introducing many auxiliary phosphoform variables can be cumbersome in more realistic models. The modified fraction approach is intuitive and flexible (model the sites instead of the protein), and it only introduces one additional variable per protein.

The components of the MF model, namely the function \( h(x) \) and the rates of phosphorylation and dephosphorylation of individual sites, can potentially be subject to direct experimental measurement, unlike the use of more abstract Hill function terms. This can allow to carry out ‘raw-data modeling’ e.g. to use an experimentally measured activity function \( h(x) \) directly in the model rather than using it to derive parameters. This methodology is also useful out of equilibrium when the timescale of phosphorylation is sufficiently fast compared with other timescales in the system.

There are several reasons why the MF method might be particularly suitable for modeling many multisite phosphorylation systems. Nonsequential phosphorylation is likely more common in nature than the more often modeled sequential systems, since enforcing sequential phosphorylations would require an additional mechanistic effort. Bioinformatic data suggests that most phosphorylation sites in multisite proteins are located in unstructured and unconserved protein regions [6], suggesting that often it is the collective effect that matters rather than the individual sites. There is also experimental evidence in yeast signal transduction that certain proteins, such as Ste5, are activated in a concerted and redundant manner, although this type of information is still unknown for most proteins. Notice that the approximation formula would still hold if the protein activation is not concerted or redundant. In that case the formula will just approximate a dose response that may not be ultrasensitive.

One of the best known mechanisms for ultrasensitive dose responses is zero-order ultrasensitivity, as suggested by Goldbeter and Koshland [33,41]. Its main assumption is that substrate concentration needs to be in the saturation regime i.e. large compared to the \( K_m \) value of the enzymes. The MF method does not pose any constraint on \( K_m \), in fact the linear regime we used can be found when substrate concentrations are small compared to \( K_m \) values. Moreover, MF also applies when the enzymatic reactions involve complex formation, by writing a Michaelis-Menten equation for \( dp/dt \). Therefore zero-order ultrasensitivity can be used in synergy with MF in the saturation regime, and MF can be used regardless of \( K_m \) value. A zero-order dose response could likely replicate the behavior of the MWC model as shown in Figure 2C, however it could not be considered a short hand notation for MWC since the two mechanisms are fundamentally different.

In the case of the checkpoint pathway, the active proteins Pkc1 and PP2A have been found to have an approximate \( K_m \) of 0.5 \( \mu M \) [42] and 1.2 \( \mu M \) [43], respectively, for specific targets. The overall concentrations of their substrates in the cell are much lower - however these proteins tend to localize at the bud, so that the resulting local concentrations are unknown and it is unclear whether a zero-order approach would apply. A recent paper by Martins and Swain [44] points out that zero-order ultrasensitivity often results from low enzyme to substrate ratios, and localized proteins that act as enzymes and substrates for each other would likely not satisfy such ratios. That paper proposes instead a mechanism involving an allosteric model analogous to MWC, using enzyme sequestration to obtain ultrasensitivity. The paper by Kapuy et al [32] also proposes a mechanism for bistability through ultrasensitive effects, and this mechanism is applied to a detailed model of the budding yeast G1-S transition in Barik et al [45]. Other mechanisms for ultrasensitivity involve competition among substrates for the same enzyme [46] and protein localization [12], among others [28,47].

More generally, in cell regulatory networks there is a need to implement nontrivial dynamics such as bistable switches and hysteresis, which requires some form of nonlinear response in addition to the right feedback interconnections. It has been observed that several regulatory proteins have multiple phosphorylation sites, and there are many open questions regarding their intended function. Together with the onerous nature of modeling several multisite proteins using sequential networks and multiple variables each, it can be seen why a one-variable reduction such as MF can allow for much-needed simplicity.

The actual mechanisms regulating the interactions between cell size and cell division remain largely unanswered in many cases. This has left few alternative options apart from somewhat heuristic approaches in otherwise very detailed models, see e.g. [48]. The present model is an attempt, based on recent experiments, to construct a detailed mechanistic model in the context of the polar to isotropic bud transition in yeast. Notice that if the proteins PP2A, Pkc1, Zds1 had only one site each, then \( h(x) = x \) according to the argument in the first Results section, and the downstream and upstream models could never be bistable (see equations (2) and (3)). The multiple sites are providing the underlying nonlinearity so that the models can have interesting dynamical behaviors. This is consistent with the work by Yang et al [49], which reached the same conclusion through randomized parameter searches in multisite cell cycle models. Also, the equations (2), (3), which represent the steady states of the downstream and upstream systems, have the same qualitative behavior for a wide range of parameters. In this sense one can say that the switch-like nature of the checkpoint is robust to many parameter changes, provided that a few key qualities are satisfied. The bistability in each subsystem is due in part to positive feedback loops in each.
subsystem, one between Pkc1 and the Rho1/Pkc1 dimer, and another between Zds1 and the PP2A/Zds1 dimer. Notice that while Rho1/Pkc1 activates PP2A, the downstream PP2A/Zds1 inactivates Pkc1, forming a negative feedback loop. This feedback could serve to reduce the activity of the pathway before a sufficient Rho1 signal has accumulated. Notice that the switch-like activation of Cdk1 is a complex process that may well be regulated by other mechanisms in conjunction with the switch discussed, and that this overall regulation also depends on the organism studied.

The MF framework eliminates several parameters such as the number of phosphorylation sites $n$ (as long as it is sufficiently large), the transition rates $L_i$ and the cooperativity coefficient $c$. The remaining parameters, such as the shape of the activity function $h(x)$, can potentially be measured in the lab using site-directed mutagenesis and activity assays. However this is still a formidable task and one that is yet to be done for most proteins involved in cell cycle regulation.

Since it is assumed in the derivation of the formula (1) that the sites are roughly independent from each other, one might think that the MF framework doesn’t work for allosteric or cooperative systems. However the detailed model in Figure 2 is allosteric, and yet the model closely describes its dynamics. In simulations we find that the accuracy of the representation is increased when $n$ is large (e.g. $n > 10$) and/or the cooperativity is weak. The use of a MWC-type model for multisite phosphorylation has been pointed out in the past, see for instance [5] and the more recent [44].

Questions for future work include: if phosphorylation and dephosphorylation of the multisite protein $P$ is not faster than other processes in the system, can one still approximate $P(t)$ away from equilibrium? This might be possible by defining a simple differential equation for $P$ instead of the algebraic equation (1). Also, the linear dynamics used to calculate the fraction $p$ can be replaced by more complex models such as a Michaelis-Menten reaction, which may be explored in detail, including the interaction with zero-order mechanisms. This might raise the question of how the corresponding model reduction might be, possibly involving multivalued functions $h(x)$.

**Methods**

For convenience we include in one location all chemical reactions of the model, mass conservation laws, the definition of auxiliary variables following the multisite modeling formalism, and a self contained set of differential equations after eliminating additional variables. Recall that for multisite proteins $x$ represents the fraction of active sites, $X$ the active monomer concentration, $X$ the total concentration including active and inactive forms, and $S_X$ the total amount of $X$ in the system including dimer and monomer forms. Also recall that Rho1 and Pkc1 are denoted by $R$, $K$, $Zds1$ and $PP2A$ by $Z$, $P$, and the Rho1/Pkc1 and Zds1/PP2A dimers by $C$, $D$, respectively.

**Model reactions**

\[
\begin{align*}
&z_m \xrightarrow{q_1 D} z + P \xrightarrow{q_2} D \xrightarrow{q_3 C} p
\end{align*}
\]

**Mass conservation laws**

\[
\begin{align*}
z_m + z & = 1 \\
p_m + p & = 1 \\
&k_m + k = 1 \\
&Z + D = S_Z \\
P + D = S_P \\
&K + C = S_K
\end{align*}
\]

**MF framework equations**

\[
\begin{align*}
Z &= Zh_1(z) = (S_Z - D)h_1(z) \\
P &= Ph_2(p) = (S_p - D)h_2(p) \\
K &= Kh_3(k) = (S_k - C)h_3(k) \\
C &= Ch_1(k) \\
D &= Dh_1(z)h_2(p)
\end{align*}
\]

**Differential equations**

\[
\begin{align*}
\frac{dz}{dt} &= q_1 Dh_1(z)h_2(p)(1 - z) - q_{-1} z \\
\frac{dD}{dt} &= q_2(S_Z - D)(S_P - D) - q_{-2} D \\
\frac{dp}{dt} &= q_3 Ch_3(k)(1 - p) - q_{-3} p \\
\frac{dR}{dt} &= -q_4 R(S_K - C) + q_{-4} C + q_6 R_m - q_{-6} R - q_{-9} R \\
\frac{dC}{dt} &= q_4 R(S_K - C) - q_{-4} C \\
\frac{dk}{dt} &= q_5 Ch_3(k)(1 - k) - q_{-5} D h_1(z)h_2(p)k \\
\frac{dR_m}{dt} &= -q_6 R_m + q_{-6} R + q_{9} v - q_{-9} R_m
\end{align*}
\]

**Parameter values**

Since most quantitative information about the pathway is unknown, we make educated estimates on the order of magnitude of the parameters. Since parameters are clustered in equations (2)
Table 1. Parameter values in the cell size checkpoint model.

| Parameter | Value |
|-----------|-------|
| \( S_Z \) | 0.1 \( \mu M \) |
| \( S_P \) | 0.1 \( \mu M \) |
| \( S_K \) | 0.01 \( \mu M \) |
| \( \xi_{\text{high}} \) | \( 10^{-4} \text{nM s}^{-1} \) |
| \( \xi_{\text{low}} \) | \( 8 \times 10^{-6} \text{nM s}^{-1} \) |
| \( q_{-1} \) | 1 \( \text{s}^{-1} \) |
| \( q_{+1} \) | 10 \( \mu M^{-1} \text{s}^{-1} \) |
| \( \delta_1 \) | 100 |
| \( \delta_2 \) | 10 |
| \( \delta_3 \) | 1000 |

The model parameters used in the simulation of the checkpoint pathway in Figures 3, 4 are detailed in this table, including total protein concentrations, binding and unbinding rates, linear reaction rates, rates of vesicle flow and the parameters for the activation functions \( h(x) \). We use the notation \( Q_2 = q_{-1}/q_{+1}. \) See the Methods section for an estimation of their values.

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Supporting Information

Figure S1 A: A rule-based approach for modeling multisite phosphorylation, which results in a system equivalent to MWC. Here \( I\in\{0,1\}^\beta \) represents any phosphoform state, and \( J \) is the result of adding one site phosphorylation to \( I \). B: Timecourse simulation for the MWC model in which the kinase concentration \( E \) is increased linearly over time, and the full MWC model is compared with the MF approximation and a model using a single site. Here \( n = 12, \ e = 0.3, P = 12 \text{nM}, F = 1 \text{nM}, L_1 = e^{-n/2}, \ a = \beta = L_2 = 1 \) for the detailed model, and \( n = -n \ln e, \ a = \beta = \gamma = \delta = L_1 = 1 \) for the MF approximation. (EPS)

Figure S2 For the MWC model in Figure 2B, a bootstrap parameter analysis of the corresponding MF system. Baseline parameter values are \( a = \beta = L_2 = 1, L_1 = e^{-n/2} \). A: The parameters \( a, \beta, L_1 \) are varied over four orders of magnitude each, and the percentage error with the MF model is calculated. B: A similar analysis of the Hill exponent of the MF model. (EPS)

Text S1 The supplementary material to this paper has three sections. In Section S1.1 and S1.2 we carry out a mathematical analysis of the downstream and upstream subsystems of the checkpoint model, respectively. Each analysis results in the reduction of the system at steady state to a single equation. In Section S1.3 we derive the MWC model using a system of equations involving all \( 2^\beta \) protein phosphoforms and a rule-based system of reactions. (PDF)

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Author Contributions

Conceived and designed the experiments: GE DRK. Analyzed the data: GE DRK. Wrote the paper: GE DRK. Implemented Matlab simulations: GE AV. Carried out mathematical proofs: GE AV. Wrote the supplementary text: GE.
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