A modular approach for modeling the cell cycle based on functional response curves

Jolan De Boeck^{1,2}, Jan Rombouts¹, Lendert Gelens^{1*},

1 Laboratory of Dynamics in Biological Systems, Department of Cellular and Molecular Medicine, University of Leuven, 3000 Leuven, Belgium

 ${\bf 2}$ Stem Cell Institute Leuven, Department of Development and Regeneration, University of Leuven, 3000 Leuven, Belgium

* lendert.gelens@kuleuven.be

Abstract

Modeling biochemical reactions by means of differential equations often results in systems with a large number of variables and parameters. As this might complicate the interpretation and generalization of the obtained results, it is often desirable to reduce the complexity of the model. One way to accomplish this is by replacing the detailed reaction mechanisms of certain modules in the model by a mathematical expression that qualitatively describes the dynamical behavior of these modules. Such an approach has been widely adopted for ultrasensitive responses, for which underlying reaction mechanisms are often replaced by a single Hill equation. Also time delays are usually accounted for by using an explicit delay in delay differential equations. In contrast, however, S-shaped response curves, which are often encountered in bistable systems, are not easily modeled in such an explicit way. Here, we extend the classical Hill function into a mathematical expression that can be used to describe both ultrasensitive and S-shaped responses. We show how three ubiquitous modules (ultrasensitive responses, S-shaped responses and time delays) can be combined in different configurations and explore the dynamics of these systems. As an example, we apply our strategy to set up a model of the cell cycle consisting of multiple bistable switches, which can account for events such as DNA damage and coupling to the circadian clock in a phenomenological way.

Author summary

Bistability plays an important role in many biochemical processes and typically emerges from complex interaction patterns such as positive and double negative feedback loops. Here, we propose to theoretically study the effect of bistability in a larger interaction network. We explicitly incorporate a functional expression describing an S-shaped input-output curve in the model equations, without the need for considering the underlying biochemical events. This expression can be converted into a functional module for an ultrasensitive response, and a time delay is easily included as well. Exploiting the fact that several of these modules can easily be combined in larger networks, we construct a cell cycle model consisting of multiple bistable switches and show how this approach can account for a number of known properties of the cell cycle.

Keywords

Bistability, Ultrasensitivity, Hill function, Delay differential equations, Ordinary differential equations, Modularity, Cell cycle oscillations, Cell cycle checkpoints, Circadian clock

Introduction

Cell division and the correct separation of genomic material among daughter cells is fundamental for 2 the proper development, growth and reproduction of a living organism. The molecular mechanisms that 3 underlie these processes are highly evolutionarily conserved. Incorrect cell division can have detrimental 4 effects, ranging from developmental defects to the transformation of healthy somatic cells into cancer cells. 5 Because of this, tight regulatory mechanisms are established early in embryonic development to ensure the 6 correct replication of DNA and cell division. These control mechanisms, or cellular checkpoints, ensure that the cell cycle only progresses to its next phase if appropriate intra- and extracellular conditions are 8 fulfilled. These conditions include the absence of DNA damage, proper alignment of the chromosomes in the metaphase plane, and abundance of nutrients and growth factors [1]. The sequential nature of cell 10 cycle progression where the start of one phase depends on the completion of a previous phase resulted in 11 the view of the cell cycle as a 'domino-like' process [2,3]. 12

Over the years, both theoretical and experimental studies have demonstrated how the mechanisms 13 that control the 'domino-like' nature of cell cycle progression are centered around bistable switches. Such 14 bistability is often encountered in systems that possess an S-shaped steady-state response curve and means 15 that the system may settle in two different stable states depending on the initial conditions (Fig. 1A). 16 These observable stable steady states correspond to the lower and top branch of the S-curve, while the 17 middle part of the S-curve is unstable and cannot be measured experimentally. The full S-shaped response 18 curve can therefore not be obtained experimentally, and bistability typically manifests itself through sharp 19 jumps and hysteresis in the measurements. Hysteresis appears when the threshold for switching from low 20 to high response levels is different from the threshold for switching from high to low response levels. In 21 the cell cycle, these all-or-none responses ensure robust transitions between different cell cycle phases, 22 while hysteresis prevents the cell cycle from returning to previous phases without having been through 23 the whole cell cycle. Control mechanisms at checkpoints can prevent such transitions, either by keeping 24 the input at sub-threshold levels or by shifting the right threshold to higher input levels [4–6]. 25

One of the first cell cycle transitions for which the role of an underlying bistable switch was established, 26 was mitotic entry in early embryonic cells of Xenopus frogs. This cell cycle transition is characterized 27 by the switch-like phosphorylation of numerous proteins, referred to as mitotic substrates. Throughout 28 interphase, cyclin B (CycB) molecules are gradually produced and bind to cyclin-dependent kinase 1 29 (Cdk1). At the onset of mitosis, the phosphorylation state of these CycB-Cdk1 complexes abruptly changes, 30 resulting in their switch-like activation and subsequent phosphorylation of the mitotic substrates [7]. The 31 sudden changes in phosphorylation state of CycB-Cdk1 are generated by positive and double negative 32 feedback loops with the phosphatase Cdc25 and kinase Wee1, respectively [8]. Theoretical work proposed 33 that bistability, resulting from these feedback loops, plays an important role in cell cycle progression [9,10]. 34 Afterwards, this was experimentally validated [11–13]. Not only CycB-Cdk1, but also counteracting 35 phosphatases such as PP2A help in regulating the phosphorylation state of mitotic substrates, and 36 thus entry into and exit from mitosis. Recent findings showed how PP2A too is regulated in a bistable 37 manner [14–16]. Furthermore, the phosphatases PP1 and Fcp1 have been implicated in regulating the exit 38 from mitosis, but how exactly these phosphatases interact with the different cell cycle regulators remains 39 a topic of active research [17, 18]. 40

The observation that, just like M-phase entry, other phase transitions of the cell cycle are irreversible 41 under normal physiological conditions, suggested that these too are built on bistable switches. For 42 example, the transition from G1 to S phase is governed by a complex interplay between extracellular 43 signals, CycD-Cdk4/6, the transcription factor E2F, retinoblastoma protein (Rb) and CycE-Cdk2. These 44 interactions lead to bistability in the activity of E2F [19]. Appropriate conditions, such as a sufficient 45 concentration of extracellular growth factors and nutrients, will push the cell across the threshold of the 46 switch to a 'high E2F' state. At this point the cell irreversibly commits to the cell cycle, i.e. it will finish 47 the started round of cell division even if nutrient levels drop [5, 20]. Another cell cycle transition for 48 which bistability has been proposed to fulfill an important role is the metaphase-to-anaphase transition 49 and the accompanying spindle assembly checkpoint (SAC). During this transition, microtubules in the 50 mitotic spindle need to correctly attach to the sister chromatids. Once these are correctly attached, the 51 cohesin rings that are keeping the sister chromatids together can be cleaved, upon which the chromatids 52



Fig 1. Modularity in the cell cycle (A) An S-shaped response is characterized by the coexistence of two stable and one unstable steady states for a certain range of input levels (blue shaded area). Cell cycle transitions can be described as sudden jumps from the lower to upper branch of the curve (or vice versa). Checkpoints act by fixing the input at sub-threshold levels (x) or shifting the threshold value to a different input level (blue curve). (B) Different phase transitions in the cell cycle have been shown to be centered around bistable switches for which S-shaped response curves have been experimentally measured. (C) Entry into mitosis (and other cell cycle transitions) can be isolated as a biological module from the bigger network of other cell signaling events in the cell (i). Whereas the dynamical features of these modules have been studied extensively by considering detailed reaction schemes (ii), we propose a phenomenological approach based on three easily combinable 'functional modules': time delay, ultrasensitive and S-shaped responses (iii).

are separated by the mitotic spindle. Although some experimental studies question the all-or-none nature of the SAC [21,22], indirect experimental and theoretical findings support the idea that this transition is also centered around a bistable switch [23–26]. Given the recurring occurrence of all-or-none transitions throughout the cell cycle, the latter has been envisioned as a chain of interlinked bistable switchs (Fig. 1B) [26,27].

Although a chain of bistable switches can account for control mechanisms at cell cycle transitions 58 and checkpoints, an additional mechanism is still needed to drive the cycle forward and reset it back to 59 its initial state at the end — thus putting the dominoes back up after toppling them. This is provided 60 by the periodic production and degradation of cyclins [28]. The CycB-Cdk1 complexes that accumulate 61 during interphase are activated at mitotic entry. In turn, they will activate the Anaphase-Promoting 62 Complex/Cyclosome (APC/C), a ubiquitin ligase. This protein complex then induces the degradation 63 of the cyclins [29]. APC/C regulation is believed to be a complex multi-step mechanism in which 64 time delays play an important role, hence introducing a delayed negative feedback loop in the reaction 65 network [30, 31]. The latter is generally known to allow for robust and sustained oscillatory behavior of 66 dynamical systems [32]. Even when the cellular checkpoints are absent, this negative feedback loop can 67 drive autonomous biochemical oscillations in a 'clock-like' manner, as for example seen in Xenopus and 68 sea urchin eggs which rapidly alternate between phases of DNA replication (S-phase) and segregation of the chromosomes (M-phase) [3,33].

The cell cycle is a complex process with many interacting parts. Dividing it into a sequence of discrete relation modules such as bistable switches might seem artificial. However, since its introduction in biology by Hartwell *et al.*, [34], such a modular approach describing biological processes has been justified by advances in synthetic biology, genomics, cell signalling and single-cell techniques [35–38]. Furthermore, studying the cell cycle based on discrete modules is warranted by the temporal segregation of the different cell recycle phases, and the presence of bistability itself [39,40].

Even though separating the full system into different modules greatly reduces the complexity, under-77 standing the dynamical behavior of those modules often requires mathematical models instead of mere 78 intuition [41]. Even for a single module, a biochemically detailed study results in a large number of 79 variables and parameters, many of which are difficult to determine experimentally. As this might hamper 80 interpretation of the results, it is often desirable to reduce the complexity of the mathematical model. 81 One way to accomplish this is by focusing on a core subnetwork and omitting all other reactions, thus 82 (hopefully) capturing the key qualitative behavior of the system [42]. Although this approach certainly 83 has its merits, some dynamical properties such as bistability may be lost when reducing the model too 84 much [43,44]. Of note, this does not mean that models of small reaction networks imply less interesting 85 dynamical features: for example, even a system containing a single molecule can behave in a bistable 86 manner [45]. 87

Another strategy to reduce the complexity of a model, while retaining much of the system's dynamical 88 behavior, is by making certain simplifying assumptions about the underlying reactions, rather than 89 omitting them. For example, using timescale separation methods, one can identify variables that evolve on 90 a much faster timescale than others. The equations for these variables can be replaced by expressions for 91 the steady-state response curves, which can be introduced into the equations for the slow variables. This 92 process is often responsible for the introduction of nonlinearities in a reaction system. Examples include 93 Michaelis-Menten enzyme kinetics, zero-order ultrasensitivity, multisite phosphorylations, cooperative 94 binding events and stoichiometric inhibition [46], whose net result is often summarized by their steady-95 state response curve. These steady state response curves can be described by reaction-specific formulas, 96 such as the Michaelis-Menten equation for enzyme-kinetics [47] or Goldbeter-Koshland function for 97 zero-order ultrasensitivity [48]. Alternatively, steep responses —of different molecular origins— can often 98 be adequately approximated by a Hill equation [49–51]. The Hill equation is therefore used to introduce 99 steep responses, without regarding the underlying reactions that generated them. Another example 100 of dynamical behavior that can be explicitly incorporated are time delays, which arise in biochemical 101 reactions due to the non-zero time to complete physical and/or biochemical processes, such as molecular 102 transport or intermediate reactions [52, 53]. Such delays can be accounted for via delay differential 103 equations. In addition to being simpler than mechanistic models, a phenomenological model based on 104 explicit mathematical expressions is often the sole option to describe experimentally observed responses 105 whenever the underlying molecular mechanisms are unknown. 106

S-shaped responses typically emerge from regulatory mechanisms like positive or double negative 107 feedback loops [54]. Many cell cycles models include these feedback loops to generate the S-shaped 108 response [10,55–57]. Steep responses and time delays have been explicitly included in cell cycle models 109 using a simple mathematical form. Given the recurring occurrence of bistable switches in the cell cycle, it 110 is remarkable that such a direct mathematical formulation of S-shaped response curves has never been, as 111 far as we know, explicitly incorporated into cell cycle models. Here we want to close this gap (Fig. 1C) by 112 providing an easily tunable, phenomenological expression for such an S-shaped response. This expression 113 can easily be converted into a functional module for ultrasensitivity by tuning a few parameters, and can 114 be combined with a time delay. As such, we provide a toolbox of three functional modules (ultrasensitive 115 response, S-shaped response, time delay) that can be combined in different configurations to model a 116 variety of biological processes. We will illustrate this approach with models that include different bistable 117 switches in the cell cycle. 118

Results

Ultrasensitive responses, S-shaped responses and delay as functional modules 120

Before considering actual models of the cell cycle, we start with a general overview of the functional ¹²¹ modules that will be used in the rest of the paper: ultrasensitivity, S-shaped response and delay. Nonlinear ¹²²

response curves are often described by a Hill equation of the form $\text{Output} = \frac{\text{Input}^n}{K^n + \text{Input}^n}$. For n > 1, ¹²³

this gives a sigmoidal response curve, with higher values of n corresponding to steeper responses (Fig. 2A). ¹²⁴ Such steep responses are often said to be 'ultrasensitive' [51]. Our ultrasensitive module thus consists of a ¹²⁵ Hill function. ¹²⁶



Fig 2. Ultrasensitive responses, S-shaped responses and delay as functional modules. (A) An ultrasensitive input-output response is characterized by the threshold value K and the Hill coefficient n. (B) An ultrasensitive function can be converted into a S-shaped response of different widths: for $\alpha = 0$, an ultrasensitive response is obtained, while increasing α increases the width of the S-shaped region. (C) A certain lag time τ between input and output is encountered in several biochemical reactions. (D) Several combinations of the three functional modules, always in the presence of negative feedback, to describe oscillations in the cell cycle.

In order to produce a mathematical expression for our second module, the S-shaped response, we would like to 'bend' the ultrasensitive curve just described such that it becomes S-shaped (Fig. 2B). Mathematically, this can be achieved by starting from a Hill function, and multiplying the threshold value K with a scaling function ξ (Out):

$$\operatorname{Out} = \frac{\operatorname{In}^{n}}{K^{n} + \operatorname{In}^{n}} \qquad \Rightarrow \qquad \operatorname{Out} = \frac{\operatorname{In}^{n}}{[\xi(\operatorname{Out}) \cdot K]^{n} + \operatorname{In}^{n}}.$$
 (1)

Note that the latter expression does not define a function — which is impossible, as per definition a 131 S-shaped response has multiple outputs for one input. Instead, it should be interpreted as the steady 132 state response of an ordinary differential equation (ODE): 133

$$\frac{d\operatorname{Out}}{dt} = \frac{\operatorname{In}^{n}}{[\xi(\operatorname{Out}) \cdot K]^{n} + \operatorname{In}^{n}} - \operatorname{Out}.$$
(2)

Although different options for ξ are possible, a straightforward choice in analogy with, for example, the Fitzhugh-Nagumo model would be a cubic function [58]: 134

$$\xi(\operatorname{Out}) = 1 + \alpha \cdot \operatorname{Out} \left(\operatorname{Out} - 1\right) \left(\operatorname{Out} - r\right).$$

Changing the parameter α allows for a smooth transition between an ultrasensitive and S-shaped response: for $\alpha = 0$ we retrieve the original ultrasensitive response, while increasing its value results in wider S-shaped regions (Fig. 2B). The parameter r can be used to make the response curve asymmetric. In the remainder of the text, we keep r = 0.5 (as in Fig. 2B), and we will not interpret the parameter rfurther biologically. The effect of the different parameters on the shape of the scaling function is discussed in more detail in S1 Text and S1 Fig.

For the last module, i.e. delay, a lag time τ between input and output of the system (Fig. 2C) can be explicitly modeled by setting $\operatorname{Out}(t) = \operatorname{In}(t - \tau)$. This turns the equation into a delay differential equation. In what follows, we will combine these three functional modules in several configurations, always in combination with a negative feedback (Fig. 2D), to model different aspects of the cell cycle. We will explain how changing the mathematical properties and function parameters influence the overall dynamical behavior of the model.

S-shaped, but not ultrasensitive, responses cause a two-dimensional system of the cell cycle to oscillate 149

The most straightforward strategy for combining the different functional modules into actual cell cycle 150 models, would be to start with a simple — but biologically relevant — reaction network and gradually add 151 additional modules to increase the scope of the model. The simplest cell cycle models arguably describe 152 the 'clock-like' cell cycle oscillations in *Xenopus laevis* eggs, compared with the more complex 'domino-like' 153 mechanism in somatic cells. Indeed, the early embryonic cell cycle in X. laevis simply cycles between S-154 and M-phase and lacks checkpoints and gap phases [33]. Furthermore, cell cycles 2 till 12 after fertilization 155 of the egg are characterized by an increased activity of the phosphatase Cdc25 relative to the kinase Weel, 156 resulting in quick activation of CycB-Cdk1 and subsequent APC/C activation [59]. Taken together, the 157 reaction network at the core of the early embryonic cell cycle can be simplified by a negative feedback 158 loop, where active CycB-Cdk1 phosphorylates and activates APC/C, which then causes degradation of 159 CycB molecules. A two-variable phenomenological model of this system is given by (Fig. 3A): 160

$$\begin{cases} \frac{d[\operatorname{Cdk1}]}{dt} = b_{\operatorname{syn}} - b_{\operatorname{deg}}[\operatorname{Cdk1}] \cdot [\operatorname{APC}]^* \\ \frac{d[\operatorname{APC}]^*}{dt} = \frac{1}{\epsilon_{\operatorname{apc}}} \left(\frac{[\operatorname{Cdk1}]^n}{K_{\operatorname{cdk,apc}}^n + [\operatorname{Cdk1}]^n} - [\operatorname{APC}]^* \right). \quad (\operatorname{system} (\operatorname{i-a})) \end{cases}$$

Here, the two variables [Cdk1] and [APC]^{*} represent the concentrations of activated CycB-Cdk1 ¹⁶¹ complexes and the ratio of activated APC/C to total APC/C, respectively. The first equation describes ¹⁶²

the rate of change of activated CycB-Cdk1 complexes and consists of two terms: the constant synthesis of cyclins and their APC/C-dependent degradation. In this model all synthesized cyclin immediately binds to Cdk1 and activates it. This is why cyclin synthesis is directly included in the equation for [Cdk1]. The second equation in system (i-a) states that the rate of change of [APC]* is proportional to the difference between the experimentally determined [APC]* ultrasensitive steady state response and its current level (similar as in Eq. 2) [30, 59], which it approaches at a time scale $1/\epsilon_{apc}$.



Fig 3. S-shaped, but not ultrasensitive, responses cause a two-dimensional system of the cell cycle to oscillate. (A) Block diagram of the ultrasensitive, negative feedback system (i-a). (B) Effect of relative synthesis c on the location of the [Cdk1] nullcline in the phase plane. (C) Time traces for two different initial conditions of system (i-a), denoted by either continuous or dotted line. Only one steady state exists. (D) Block diagram of the bistable, negative feedback system (i-b). (E) For wide bistable [APC]* nullclines, three intersections in the phase plane can exist, resulting in bistability of the overall ODE system. Closed black dots depict stable steady states, while open dots denote unstable steady states. (F) Time traces for system (i-b). Depending on the initial conditions, the system will settle in one of two stable steady states. (G) Number and position of the steady states for system (i-b). Osc = oscillations, Bi = bistable, Mono Low = one stable steady-state on lower branch of [APC]* nullcline, Mono Top = one stable steady-state on top branch of [APC]* nullcline intersects the [APC]* nullcline in between its folds, one unstable steady state exists, resulting in oscillations. The used parameter values can be found in the panels.

Because system (i-a) has two variables, its behavior can be conveniently studied in the phase plane. 169 To determine the steady state behavior of system (i-a), one needs to find the number and location of 170 intersections of the [Cdk1] and [APC]* nullclines. These are the curves in the [Cdk1]-[APC]* phase plane 171 for which $d[Cdk_1]/dt = 0$ and $d[APC]^*/dt = 0$ respectively. Here, only one intersection can exist, whose 172 location is determined by the relative position of both nullclines (Fig. 3B-C). Moreover, this steady state is 173 linearly stable (see S1 Text for details). While parameters n and $K_{cdk,apc}$ affect the ultrasensitive [APC]* 174 nullcline, the location of the [Cdk1] nullcline is determined by the ratio of $b_{\rm syn}$ and $b_{\rm deg}$. However, after 175 non-dimensionalizing the system to facilitate mathematical analysis (see Methods for details), this ratio 176 can be re-expressed as the product of $K_{cdk,apc}$ with a newly introduced dimensionless parameter c, termed 177 the relative synthesis rate: 178

$$c = \frac{b_{\rm syn}}{K_{\rm cdk, apc} \cdot b_{\rm deg}}.$$

Changing c thus changes the location of the [Cdk1] nullcline in the phase plane: whereas low values shift it to the left, high c values shift the [Cdk1] nullcline to the right (Fig. 3B). It is mainly this parameter that will be used in the remainder of the text when assessing the effect of the [Cdk1] nullcline on the dynamical behavior of the overall system.

As the steady state is stable, this system can not serve as a basic model to describe cell cycle oscillations. To make the system oscillate, either a time delay or a S-shaped response can be introduced. The effect of a time delay in combination with ultrasensitivity has previously been studied in detail [60] (see also S2 Fig). Here we will focus on the effect of converting the ultrasensitive module from system (i-a) into a S-shaped module, which is in line with recent experimental findings that measured hysteresis in APC/C response curves [15]. As discussed above, the threshold $K_{cdk,apc}$ can be multiplied with a cubic scaling function $\xi([APC]^*) = 1 + \alpha_{apc}[APC]^*([APC]^* - 1)([APC]^* - r)$ (Fig. 3D):

$$\begin{cases} \frac{d[\operatorname{Cdk1}]}{dt} = b_{\operatorname{syn}} - b_{\operatorname{deg}}[\operatorname{Cdk1}] \cdot [\operatorname{APC}]^* \\ \frac{d[\operatorname{APC}]^*}{dt} = \frac{1}{\epsilon_{\operatorname{apc}}} \left(\frac{[\operatorname{Cdk1}]^n}{\left(\xi([\operatorname{APC}]^*) \cdot K_{\operatorname{cdk,apc}}\right)^n + [\operatorname{Cdk1}]^n} - [\operatorname{APC}]^* \right). \quad (\text{system (i-b)}) \end{cases}$$

Note that by multiplying the threshold $K_{\text{cdk,apc}}$ with ξ in system (i-b), the [APC]* nullcline becomes 190 S-shaped when $\alpha_{\rm apc} > 0$ (as introduced in Eq. 2 and Fig. 2B). Systems exhibiting such a S-shaped response 191 are good candidates for showing bistability, in which case the system can evolve over time to two steady 192 states. Whether this behavior is indeed observed, depends on the number of intersections between the 193 [Cdk1] and [APC]^{*} nullclines in the two-dimensional phase plane. As the [APC]^{*} nullcline is now S-shaped 194 instead of sigmoidal, either one or three intersections can exist with the [Cdk1] nullcline (Fig. 3E,G,H). 195 The system is bistable when three intersections exist, with one steady state being unstable and the other 196 two stable. It is the initial condition of the system that determines in which of the two stable ones the 197 system will settle (Fig. 3F). Whenever the system has only one steady state (i.e. one intersection of the 198 nullclines), the latter can either be stable (on the upper or lower branch of the S-shaped nullcline) or 199 unstable. Only in the latter case, i.e. if both nullclines intersect in between the fold points of the [APC]^{*} 200 nullcline (Fig. 3H), sustained oscillations around this steady state appear. These oscillations can be 201 thought of as 'orbiting around the nullclines', i.e. they show increasing [Cdk1] levels on the lower branch 202 of the response curve until the [APC]^{*} activation threshold is reached. At this point, [APC]^{*} levels jump 203 to a higher value, which triggers the decrease of [Cdk1]. The system then proceeds along the upper branch 204 until the inactivation threshold is reached and the cycle is complete. These oscillations, characterized by 205 slow progress along the branches of the nullclines and quick jumps between them, are called relaxation 206 oscillations. They are similar to oscillations observed in, for example, the FitzHugh-Nagumo equations [58]. 207

From the observation that the system oscillates if the nullclines have a single intersection in the middle, we can derive some qualitative conditions for oscillations to exist: the $[APC]^*$ nullcline should not be too wide and the [Cdk1] nullcline should not be located too far to the left nor to the right (relative to the other nullcline). Indeed, from Fig 3G, we see that oscillations are favored for *c* values around 0.5 and narrow S-shaped regions. As the S-shaped region becomes wider, the period of oscillations and [Cdk1]

amplitude increases (S3 Fig), but the range of relative synthesis c for which oscillations can be sustained decreases. For [APC]^{*}, the effect of the width of the S-shaped region on the amplitude is more moderate, and the amplitude is near-maximal for most oscillations. From the time profiles, we further see that the overall shape of the oscillations has a typical sawtooth-like waveform (Fig. 3I). It is interesting to note that, in this system, the effect of the S-shaped regions is mixed: on the one hand, it is required for the system to oscillate, but on the other hand a wider S-shaped region makes oscillations less likely, as in that case the system settles in a steady state on the upper or lower branch. 213

Asymmetric S-shaped APC/C nullclines slow down APC/C (in)activation

The use of a cubic function ξ with r = 0.5 results in what we would call 'symmetric' S-shaped responses, 221 meaning that the vertical distance between the left folding point and lower branch equals the vertical 222 distance between the right folding point and the upper branch of the response curve (S4 Fig). One might 223 then wonder what would happen if the assumption of such symmetry is relaxed. After all, the lower and 224 top branch of experimentally observed S-shaped response curves (recall that the middle part cannot be 225 measured experimentally) will typically not be symmetric. Examining some of the experimentally measured 226 responses in [15], such as the one for APC/C phosphorylation levels, indeed shows that asymmetry can 227 occur. 228

To explore the effect of asymmetry, one needs to shift the folding points of the S-shaped response 229 up or down. Using the cubic ξ function, this can be accomplished by altering the value of parameter r. 230 However, this approach would shift the response curve horizontally too, making it impossible to isolate 231 the effect of vertical asymmetry from other effects. Similar difficulties exist when ξ is defined by other 232 smooth functions, such as a quadratic function (S4 Fig). As a solution to this problem, we approximated 233 the cubic scaling function by a piecewise linear function in combination with large values of n (n = 300). 234 The piecewise approach enables us to exactly position the extrema of $\xi([APC]^*)$ at the desired location, 235 while the high values of n ensure that these extrema correspond to the folds of the S-shaped nullcline (see 236 S1 Text for further details), thus giving precise control over their location. 237

The approximation of the cubic scaling function $\xi([APC]^*)$ (with arbitrarily chosen values for r and 238 $\alpha_{\rm apc}$) by a piecewise fit of linear functions through its extrema (inset Fig. 4A), leads to the response 239 curve in Fig. 4A. When comparing the dynamics of the system with the piecewise approximation to the 240 dynamics with a smooth cubic function, similar oscillatory regions in parameter space can be observed 241 (Fig. 4B). We then exploited the fact that we can now independently change the [APC]^{*} coordinate of 242 the local extrema of $\xi([APC]^*)$ to explore the effect of an asymmetric response curve: by moving the 243 extrema of $\xi([APC]^*)$ to the right or left, the corresponding folds of the S-shaped response shifted up 244 or down in the phase plane respectively, while their [Cdk1] coordinate was kept constant at the levels 245 from Fig 4A. Again, the system can be either bistable or monostable, or have an unstable steady state 246 leading to oscillations. The period of these oscillations decreases as the vertical distance between the folds 247 increases, i.e. towards the lower right corner of Fig 4C. 248

The position of the [Cdk1] nullcline (set by the parameter c) determines the positions of the fold points for which the system can oscillate. For smaller c values, oscillations can occur for low left folds, while these need to be higher as the [Cdk1] nullcline shifts to the right (higher c). The further apart the folds, the more likely the [Cdk1] nullcline intersects the [APC]* nullcline in between the folds. This can be seen from the overlap of the oscillatory region for all c values in the lower right corner of the parameter space (Fig. 4D).

The vertical position of the folds also determines the shape of the time traces of [APC]^{*} activity. For 255 symmetric nullclines, here meaning $[APC]^*_{RF} = 1 - [APC]^*_{LF}$, with vertically well-separated folds, the 256 time trace resembles a square wave (Fig. 4E, F i). This leads to rapid and abrupt interconversion between 257 active and inactive [APC]^{*}, leading to well-defined transitions between interphase and M-phase. Shifting 258 the right fold upwards generates an asymmetric nullcline and introduces a period of slow exponential-like 259 build-up in [APC]^{*} activity before it switches to its maximal activity (Fig. 4E, F ii). Hence, the activation 260 of [APC]^{*} is slowed down as the time needed to convert completely inactive [APC]^{*} into its fully activated 261 form is lengthened. Likewise, shifting the left fold downwards results in a slow exponential-like decay in 262 [APC]^{*} activity prior to complete inactivation and thus in a longer time required to completely inactivate 263

bioRxiv preprint doi: https://doi.org/10.1101/2021.04.28.441747; this version posted April 28, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.



Fig 4. Asymmetric bistable APC/C nullclines slow down APC/C (in)activation. (A) Piecewise approximation of cubic scaling function $\xi([APC]^*)$ (inset) and derived bistable response curve. (B) Comparison of oscillatory region for large *n* between a cubic $\xi([APC]^*)$ (grey contourline) and its piecewise approximation (colormap). The green contour line indicates the oscillatory region for a cubic scaling function with n = 15. (C) [Cdk1] period for different [APC]^{*} coordinates of the folds ([Cdk1] coordinates were kept constant at the levels from panel A). Inset: number and location of steady-states. (D) Oscillatory regions for different values of c. (E,F) Time traces and phase plane for parameter values indicated in panel C. The used parameter values can be found in the panels.

all $[APC]^*$ (Fig. 4E,F iii).

264

265

The S-shaped module reproduces the behavior of mass-action models

In the previous sections, we showed how the S-shaped module can be used to conveniently model oscillatory 266 systems and showed that the obtained oscillations resemble those of the early embryonic cell cycle in a 267 qualitative way. The way we introduced the S-shaped response, through the modified Hill function, is 268 artificial, and not based on biochemical interactions. To show that our approach can represent the behavior 269 of an actual biochemical system, we compare our module-based system to an existing cell cycle model based 270 on mass-action kinetics. For this purpose, we first extended a previously described mass-action model of 271 the PP2A-ENSA-GWL network [61], by incorporating synthesis and APC/C-mediated degradation of 272 CycB (Fig. 5A). This system is known to generate S-shaped APC/C response curves via a double negative 273 feedback loop. More specifically, GWL (which is phosphorylated by CycB-Cdk1) indirectly inhibits PP2A 274 by phosphorylating ENSA, which is both substrate and inhibitor of PP2A [62]. PP2A itself, when active, 275 dephosphorylates and inactivates GWL, thus closing the double negative feedback loop. This leads to 276 an S-shaped response of PP2A activity as function of CycB-Cdk1 activity. As APC/C is a substrate of 277 PP2A, the S-shaped steady-state response of PP2A is translated into a similar response for APC/C as 278 function of CycB-Cdk1. 279

Once the S-shaped steady-state response curve of the mass-action model (for a fixed parameter set) 280 was calculated, we approximated this response using a piecewise linear scaling function $\xi([APC]^*)$ (Fig. 281 5D, see Methods for details). Next, we screened different values for the synthesis and degradation rate of 282 CycB for both modeling frameworks (note how these parameters do not affect the shape of the response 283 curve itself, but only alter the way the time series orbit around it). Even though the steady-state response 284 curves do not exactly match, the region in parameter space for which oscillations can be observed do 285 correspond to a good extent. Furthermore, the periods of the oscillations for both models are very similar, 286 as can also be seen from the time traces in Fig. 5C. This behavior is not unique to the chosen parameter 287 set, as also other response curves and their corresponding oscillations can be approximated well (Fig. 288 5G-I). Hence, the S-shaped module can faithfully represent the dynamical behavior of more detailed 289 mass-action models. 290

Delay increases the period, amplitude and robustness of oscillations

So far, we have described the regulation of APC/C by CycB-Cdk1 as a module in which changes in 292 CycB-Cdk1 immediately affect APC/C activity. In reality, however, this regulation is believed to be 293 a complex multi-step mechanism in which time delays play an important role [30, 60]. Two types of 294 delays can be distinguished. First, a delay τ_1 between the activation of CycB-Cdk1 and the subsequent 295 activation of APC/C can exist, i.e. a lag time between [Cdk1] levels reaching the right fold of the [APC]* 296 nullcline and $[APC]^*$ levels actually jumping to the upper branch (Fig. 6C-F). Secondly, a delay τ_2 might 297 occur between the inactivation of CycB-Cdk1 and APC/C, respectively. The delays τ_1 and τ_2 do not 298 necessarily need to be the same. For example, if the activation of APC/C were to take much longer than 299 its inactivation, delay τ_1 would have a larger value than τ_2 [60]. This time delay can be included directly 300 into the model equations (Fig. 6A): 301

$$\begin{cases} \frac{d[\mathrm{Cdk1}]}{dt} = b_{\mathrm{syn}} - b_{\mathrm{deg}}[\mathrm{Cdk1}] \cdot [\mathrm{APC}]^{*} \\ \frac{d[\mathrm{APC}]^{*}}{dt} = \frac{1}{\epsilon_{\mathrm{apc}}} \left(\frac{[\mathrm{Cdk1}]^{n}(t-\tau)}{(\xi([\mathrm{APC}]^{*}) \cdot K_{\mathrm{cdk,apc}})^{n} + [\mathrm{Cdk1}]^{n}(t-\tau)} - [\mathrm{APC}]^{*} \right) \\ \text{with } \tau = \tau_{1} = \tau_{2} \quad \text{or} \quad \tau = \tau([\mathrm{APC}]^{*}) = \tau_{1} + (\tau_{2} - \tau_{1}) \frac{[\mathrm{APC}]^{*p}}{0.5^{p} + [\mathrm{APC}]^{*p}}. \end{cases}$$

The expression for $\tau([APC]^*)$ can be used in case $\tau_1 \neq \tau_2$, as then the value of the delay is not a constant but depends on the level of $[APC]^*$ in the system. Here, we use a Hill equation in $\tau([APC]^*)$ to ensure that the time delay is a smooth function of $[APC]^*$. However, for high values of the Hill coefficient p, the Hill function approaches a step function with $\tau \approx \tau_1$ if $[APC]^*$ is smaller than 0.5 and $\tau \approx \tau_2$ when $[APC]^*$ activity is larger than 0.5, the situation we want to capture with the model.

For $\xi([APC]^*) = 1$ (i.e. $\alpha_{apc} = 0$) we retrieve the combination of a delay and ultrasensitive module 307 (S2 Fig). As before, increasing $\alpha_{\rm apc}$ converts the ultrasensitive response into a S-shaped one. Without a 308 time delay, i.e. $\tau = 0$, system (ii) is identical to system (i-b): oscillations can be observed for intermediate 309 values of the relative synthesis c (c = 0.5) whenever the S-shaped region is not too wide (as otherwise the 310 overall system becomes bistable). Low or high c values result in one stable steady state and no oscillations 311 (compare Fig. 6B and Fig. 3G). However, adding a sufficiently large time delay τ can cause the system 312 to start oscillating, even for widths of the S-shaped region and values of c where the system is mono-313 or bistable if $\tau = 0$. The wider the S-shaped region becomes, the larger the delay required to induce 314 oscillations. A maximal width exists beyond which no oscillations can be sustained, independent of the 315 time delay, corresponding to the right vertical boundary of the oscillatory region in Fig 6B (especially 316 clear for c = 0.2). 317

Both the period and [Cdk1] amplitude grow as the time delay increases (Fig. 6C-E, S5 Fig). Activation of [APC]*, on the other hand, is an all-or-none process for the majority of parameter values, with longer delays prolonging the time during which the [APC]* activity is at its maximal/minimal level, resulting in plateau phases in the time profiles (Fig. 6C-E, S5 Fig).



Fig 5. The S-shaped module reproduces the behavior of mass-action models (A) Mass-action model of the PP2A-ENSA-GWL network in which double negative feedback can lead to S-shaped response of APC/C. (B) The S-shaped module replacing the detailed reaction network in panel (A). (C) Time traces showing the resemblance between the modular approach and mass-action model. Parameters as in panel (D), with synthesis and degradation as indicated in (E,F). (D) Steady state response curve for mass-action model and fitted S-shaped module. Parameter values as given in Table 1, association constant of ENSAp with PP2A $k_{ass} = 5.62 \text{ nM}^{-1}\text{min}^{-1}$ and APC/C dephosphorylation constant $k_{da} = 3.16 \text{ nM}^{-1}\text{min}^{-1}$. (E,F) Oscillatory region as a function of CycB synthesis and degradation rates for the piecewise fit and mass-action model. Black dot denotes rate constants used in panel (C). (G,H,I) Similar as panels (D,E,F) with larger association constant ($k_{ass} = 7.94 \text{ nM}^{-1}\text{min}^{-1}$), making the S-shaped region wider, and larger APC/C dephosphorylation constant ($k_{da} = 10 \text{ nM}^{-1}\text{min}^{-1}$), shifting the response curve downwards.

Large amplitude oscillations are facilitated by two bistable switches

In the previous section we considered CycB-Cdk1 activity to be equivalent to CycB levels. This is justified for cycles 2-12 of the embryonic cell cycle of X. *laevis*, where all CycB quickly binds to Cdk1 and the complexes are rapidly activated by the phosphatase Cdc25 whose activity is dominant over the inhibitory kinase Wee1 [59]. However, in the first embryonic cycle and extracts of Xenopus eggs, a S-shaped response of CycB-Cdk1 activity as function of total CycB concentrations is typically observed [11, 12, 59]. Hence, a



Fig 6. Delay increases the period, amplitude and robustness of oscillations. (A) Block diagram of the bistable, delayed negative feedback network. (B) Oscillatory regions, located above and to the left of the plotted lines, as a function of the time delay and bistable width for different values of the relative synthesis rate c. See figure for parameter values. (C) Phase plane representation of the oscillations for different values of τ . Note how one can only conclude that $\tau_1 = \tau_2$ from time traces, not directly from the phase plane. (D,E) Time traces for parameter values indicated in panel B. τ_1 denotes the delay between activation of [Cdk1] and activation of [APC]^{*}, while τ_2 represents the delay between their inactivations. Dashed lines depict the [Cdk1] levels at the folds of the bistable [APC]^{*} nullcline. (F) State dependent time delays can account for differences in the activation and inactivation delays (p = 5).

logical next step in adjusting our cell cycle model is to add a separate S-shaped module to account for this regulation (Fig. 7A): 329

$$\begin{cases} \frac{d[\operatorname{CycB}]}{dt} = b_{\operatorname{syn}} - b_{\operatorname{deg}}[\operatorname{CycB}] \cdot [\operatorname{APC}]^{*} \\ \frac{d[\operatorname{Cdk1}]}{dt} = \frac{1}{\epsilon_{\operatorname{cdk}}} \left(\frac{[\operatorname{CycB}]^{n}}{[\xi([\operatorname{Cdk1}]/[\operatorname{CycB}]) \cdot K_{\operatorname{cyc,cdk}}]^{n} + [\operatorname{CycB}]^{n}} [\operatorname{CycB}] - [\operatorname{Cdk1}] \right) \\ \frac{d[\operatorname{APC}]^{*}}{dt} = \frac{1}{\epsilon_{\operatorname{apc}}} \left(\frac{[\operatorname{Cdk1}]^{n}}{(\xi([\operatorname{APC}]^{*}) \cdot K_{\operatorname{cdk,apc}})^{n} + [\operatorname{Cdk1}]^{n}} - [\operatorname{APC}]^{*} \right) \end{cases}$$
 (system (iii))

Here, the variable [CycB] represents the total concentration of CycB-Cdk1 complexes, [Cdk1] represents activated CycB-Cdk1 complexes and [APC]* the ratio of active APC/C molecules to the total amount of APC/C. We can still consider the total amount of CycB-Cdk1 (but not its activity) to be equivalent to the total amount of CycB, as Cdk1 is present in excess and it is assumed that free CycB molecules immediately bind free Cdk1 molecules [63]. Note how for the rate of change of [Cdk1], the Hill-like expression for the S-shaped response is multiplied by [CycB], the reason for which is explained in detail in the Methods.



Fig 7. Large amplitude oscillations are facilitated by two bistable switches. See figure for parameter values. (A) [Cdk1] amplitude for system (iii-a), with relative synthesis c/d. The green contour represents the oscillatory region for system (i), with relative synthesis c. (B) [Cdk1] amplitude for system (iii-b) and (iii-c). (C-F) Time trajectories (black lines) orbiting around the nullclines (grey lines) showing the distinct features of low and high amplitude oscillations for parameters as shown in panel B. (G-I) [Cdk1] amplitudes for the different system configurations as a function of the threshold values K. Parameters as in panel B or as indicated, with c = 0.5. (J-L) Phase plane representation of the oscillatory regions in G-I, as indicated by the dotted lines.

Adding this second module for CycB-Cdk1 regulation greatly affects the position and shape of the 336 nullclines, which now become surfaces in the three dimensional space (although they can be projected in a 2D plane, Fig. 7C-F). As a result, the parameter space sustaining oscillations changes compared 338

with system (i-b), even if the added response of [Cdk1] as a function of [CycB] is only ultrasensitive, as 339 in system (iii-a) which can be derived from system (iii) by setting the prefactor α_{cdk} in $\xi([Cdk1]/[CycB])$ 340 equal to zero (Fig. 7A). In particular, oscillations can be observed for wider [APC]* nullclines. Note that 341 whereas the relative synthesis (i.e. $b_{syn}/K_{cdk,apc}b_{deg}$) was given by the previously introduced parameter c342 for system (i-b), the non-dimensionalization of system (iii) (see Methods for details) causes the relative 343 synthesis to become the ratio of c and a newly introduced parameter $d = \frac{K_{\text{cyc,cdk}}}{K_{\text{cdk,apc}}}$. The [Cdk1] 344 amplitude is directly proportional to the width of the [APC]^{*} nullcline (Fig. 7A and S6 Fig), and the 345 amplitude is minimally affected when moving the nullclines in the phase plane by altering their threshold 346 values K (Fig. 7G,J and S6 Fig). 347

Whenever the [Cdk1] response as a function of [CycB] becomes bistable by making α_{cdk} in system (iii) 348 greater than zero, the behavior of the system drastically changes, with two distinct types of oscillations 349 emerging corresponding to either low or high [Cdk1] amplitudes (Fig. 7B, system (iii-b) for $W_{cdk,apc} = 0$ 350 and (iii-c) for $W_{cdk,apc} > 0$). Low amplitude oscillations arise from trajectories orbiting around the 351 [APC]* nullcline, but moving along the linear part of the [Cdk1] nullcline (Fig. 7C-D). Consequently, 352 their amplitude is determined by the width of the [APC]^{*} nullcline, similar as when the [Cdk1] nullcline 353 was ultrasensitive. High amplitude oscillations, on the other hand, do orbit around the [Cdk1] nullcline 354 (Fig. 7E-F), the linear part of which then becomes instructive for the amplitude of the oscillations. 355 Therefore, the amplitude will increase if the [Cdk1] nullcline is shifted to higher threshold levels $K_{\rm cvc.cdk}$ 356 (Fig. 7H-I and S6 Fig). 357

The high amplitude oscillations generated by two bistable switches induce a sufficiently large increase 358 in CycB-Cdk1 activity followed by sufficiently large inactivation required for correct entry into and exit 359 from mitosis. Furthermore, the two bistable switches make such oscillations more robust, meaning that 360 oscillations can be observed for a larger set of parameter values. From a biological perspective, this means 361 that cell cycle oscillations can be ensured even if physiological conditions within the cell (e.g. enzyme 362 activity) would fluctuate, a situation which might otherwise disrupt correct cell cycle progression. This 363 behavior can be appreciated from the larger yellow region in Fig. 7I,L compared with Fig. 7H,K. Note that 364 the nullclines need to intersect in between the folds of the S-shaped response in system (iii-b) (Fig. 7K) 365 and the previously considered two-dimensional system (i) to get oscillations, whereas this is not the case 366 for system (iii-c) with two bistable switches (Fig. 7L). Here, oscillations can be observed even if the second 367 nullcline intersects the [Cdk1] nullcline at its linear part, as long as this second nullcline reaches [Cdk1] 368 values lower than the left fold of the [Cdk1] nullcline. 369

The cell cycle can be represented as a chain of interlinked bistable switches

The results on the G2-M transition coupled to a (delayed) negative feedback in the previous sections are 371 mainly applicable to the early embryonic cell cycles that rapidly cycle between S- and M-phase without 372 intermittent gap phases. Many insects, amphibians and fish that lay their eggs externally carry out 373 multiple rounds of such rapid cell cycles following fertilization. However, all of them then pass through 374 the so-called midblastula transition (MBT) after approximately ten cell cycles [64–67]. This transition is 375 characterized by the establishment of gap phases and slowing down of the cell cycle, resulting in a higher 376 resemblance to the cell cycle as typically studied in yeast and mammalian somatic cells. Remarkably, 377 many of the transitions between these additional cell cycle phases are — just like the G2-M transition – 378 governed by bistable switches [26]. 379

During the G1-S transition in cultured mammalian cells for example, it is the activity of the E2F transcription factor that is regulated in a bistable manner. The external presence of growth factors can induce the expression of CycD and the activation of CycD-Cdk4/6 complexes, followed by the activation of E2F. E2F then induces the expression of several target genes, one of which is CycE. As CycE can further enhance the activation of E2F, a positive feedback loop is established, ultimately leading to the bistable response of E2F [19]. Consequently, the G1-S transition can be depicted as a S-shaped module, similar as was done for the G2-M transition.

It is possible to combine the different transitions — each represented by a S-shaped module — into a phenomenological model of the overall cell cycle. For this, we still need to link the 'input' and 'output' of each module in a suitable way. We link E2F activity (the output of the G1-S switch) to CycB (the input 389

of the G2-M switch) by recognizing that CycA, which is a target of E2F, drives the activation of the transcription factor FoxM1, which subsequently induces the expression of CycB [68] (Fig 8A, see Methods for associated equations). Similar as before, cyclins then need to be degraded to reset the system to the lower branch of the S-shaped response curves, after which a new round of cell division can start. Here, it is assumed that the degradation of all cyclins is induced in M-phase by activated APC/C.



Fig 8. The cell cycle can be represented as a chain of interlinked bistable switches. Parameter values from Table 2 or as indicated in the figure. (A) Block scheme of interlinked S-shaped modules representing the overall cell cycle. (B) Oscillating time traces of the cell cycle model. (C,D) Time traces from B orbiting around the bistable response curves. As in B, concentrations are normalized for their maximal value. (E,F,G) Overall period of the oscillations as a function of CycD (E) or CycB (F) synthesis and degradation rates and the width of the S-shaped responses shown in C and D (G). No oscillations exist in the white regions. Grey regions represent oscillations that are not orbiting around all three bistable switches. White dots denote parameter values used in B-D.

As seen from Fig 8B, this modular representation of the cell cycle can produce oscillations in the concentration levels of the biochemical components. Synthesis of [CycD] (here used to directly model the effect of external growth factors), results in a linear increase of its concentration levels. Once [CycD]levels reach the $[E2F]^*$ activation threshold, $[E2F]^*$ activity suddenly rises (Fig 8B,C). Activated $[E2F]^*$ then (indirectly) induces the expression of [CycB], followed by the activation of [Cdk1] when [CycB] levels cross the [Cdk1] activation threshold (Fig 8B,D). As in the previous model of the G2-M transition, active CycB-Cdk1 induces APC/C activation and subsequent degradation of the cyclins (Fig. 8C,D, S1 Video).

The oscillatory behavior of this system depends on how the time traces orbit around the steady-state 402 response curves, and hence are determined by the synthesis and degradation rates of the cyclins, together 403 with the shape of the response curves themselves. For certain combinations of parameters, no oscillations 404 can be observed (white regions in Fig 8E-G). For other combinations oscillations do exist, but these do 405 not circle around all three response curves (grey regions in Fig 8E-G; the system might for example move 406 along the top branch of the [E2F]^{*} response curve, while orbiting around the [Cdk1] nullcline), resulting 407 in irregular oscillatory patterns and hence irregular cell cycle progression (S7 Fig). The duration of the 408 different phases depends on the values of production and degradation rates. A smaller synthesis rate $d_{\rm syn}$ 409

of [CycD] and larger degradation rate d_{deg} , entails a longer time needed for [CycD] levels to reach the threshold value for [E2F]^{*} activation. This leads to oscillations with elongated G1 phases and thus longer overall periods (Fig 8E and S8 Fig). A decrease in the synthesis rate b_{syn} or degradation rate b_{deg} of [CycB] brings about oscillations with increased periods (Fig 8F). Whereas the effect of the synthesis rate can mainly be attributed to an elongation of the S-G2 phase, a diminished degradation rate causes an extension of the M phase (S8 Fig).

Control mechanisms affecting the bistable switches shape the dynamics of the cell cycle 416

Unlike the early embryonic cell cycle, the somatic cell cycle is no 'clock-like' oscillator with a fixed period, but instead a 'domino-like' oscillator in which tight control mechanisms or 'checkpoints' have been established that safeguard correct DNA replication and cell division. Hence, the cell cycle representation from the previous section — in which no such checkpoints were considered — needs to be refined. Interestingly, the bistable nature of the cell cycle transitions itself can provide the means for this regulation by shifting the (in)activation thresholds (i.e. folding points) of the S-shaped response curves to lower or higher levels [13, 69, 70].

As a first example of a checkpoint, let us consider the so-called restriction point (RP). The traditional 425 view of this checkpoint states that cells in G1 phase are predetermined to stay in this phase and not 426 enter S phase. Only if sufficiently high concentrations of growth factors are present in the extracellular 427 environment, leading to sufficiently high synthesis rates of CycD, the cell will progress into S-phase and 428 irreversibly commit to complete the started round of cell division [71]. Interlinked bistable switches can 429 account for such behavior as — once activated — $[E2F]^*$ activity can remain high even if [CycD] synthesis 430 is strongly reduced (grey area in Fig 9A). This high [E2F]* activity then further induces [CycB] expression 431 to reach the [Cdk1] activation threshold followed by activation of [APC]^{*}. 432

A second example of cellular control is the way cells preserve genomic integrity by arresting the cell 433 cycle whenever deleterious DNA lesions are encountered. During G1 phase, DNA damage can prevent 434 further cell cycle progression by inducing the degradation of CycD [72]. This behavior can be understood 435 in our model by looking at the bistable switch governing the G1-S transition. The increased degradation 436 of [CycD] (grey area in Fig 9B) keeps its levels below the [E2F]* activation threshold, thus blocking S 437 phase entry, until DNA repair re-establishes normal degradation rates and [CycD] levels can rise beyond 438 the threshold. The extent to which interphase is lengthened depends on when in G1 the damage was 439 imposed, the duration of the damage and the time required to reach the [E2F]* activation threshold after 440 the damage was repaired (Fig 9B.C). 441

DNA repair mechanisms during G2 phase have been associated with the activation of the kinase 442 Wee1, which plays an important role in inactivating CycB-Cdk1 complexes and thus preventing entry 443 into M-phase (Fig 9D) [73]. Exploiting the fact that increased Weel activity shifts the Cdk1 activation 444 threshold to higher CycB levels [13, 59], the effect of DNA damage can straightforwardly be implemented 445 by tuning the α parameter of the S-shaped module. As long as DNA damage is present, the [Cdk1] 446 activation threshold is shifted to the right (grey area in Fig 9E) and [CycB] levels keep rising until a 447 steady state is reached. When extensive damage is encountered, a permanent raise of the threshold might 448 result in permanent cell cycle arrest. However, if DNA damage is repaired, Wee1 gets inhibited again, the 449 activation threshold shifts back to lower [CycB] levels and [Cdk1] is activated, resulting in subsequent 450 [APC]^{*} activation. From the time traces (Fig 9E), it can be observed that cells enter mitosis with higher 451 [CycB] levels (and thus also higher [Cdk1] levels) when recovering from DNA damage in comparison 452 with unperturbed cells. Consequently, a longer time period is required to reach the [APC]^{*} inactivation 453 threshold and M-phase is prolonged. In contrast with the DNA checkpoint in G1 phase, modification of 454 the bistable response curve in G2 can keep the total duration of interphase unaffected after DNA damage. 455 If the damage is repaired before [CycB] levels reach the original [Cdk1] activation threshold, shifting it to 456 higher levels has no effect on interphase duration (Fig 9F). 457

These examples show that a modular description can be used to implement biological events such as DNA damage in a phenomenological way: by adjusting the overall effect on the shape of the S-shaped response, without needing to know the exact molecular mechanisms. We provide another example of this 460



Fig 9. Control mechanisms affecting the bistable switches shape the dynamics of the cell cycle. Parameter values as in Table 2 or as indicated. (A) Time traces comparing unperturbed cell cycle progression (bottom) with progression after reduction of the [CycD] synthesis rate. In line with the classical interpretation of the RP, a started round of cell division can still be completed (as seen from the activation of [APC]*) even if [CycD] synthesis is reduced (grey area). (B) Time traces comparing unperturbed cell cycle progression (bottom) with progression after DNA damage in G1 phase. Dotted line represents the [E2F]* activation threshold and grey area marks region of increased [CycD] degradation. (C) Elongation of interphase after DNA damage in G1. (D) The kinase Wee1 is controlled by DNA damage and circadian rhythms and affects the Cdk1 activation threshold. (E) Time traces comparing unperturbed cell cycle progression (bottom) with progression after DNA damage in G2 phase. Dotted line represents the [Cdk1] activation threshold. Similar as in A and B, concentrations are normalized for their maximal value in the unperturbed case. (F) Elongation of interphase after DNA damage in G2. (G) Arnold tongues showing p:q phase locking between [Cdk1] oscillations and the circadian clock. The coupling strength A_{cdk} is a measure for the extent to which the [Cdk1] activation threshold is shifted to higher [CycB] levels. (H) Time traces of phase locking for a constant natural frequency of the cell cycle (i.e. $2\pi/24$ h⁻¹) but different frequencies of the circadian clock ((i) $\omega_{\text{circadian}} \approx \frac{2\pi}{31} \text{ h}^{-1}$, (ii) $\omega_{\text{circadian}} \approx \frac{2\pi}{39} \text{ h}^{-1}$, (iii) $\omega_{\text{circadian}} \approx 2\pi/12 \text{ h}^{-1}$). (I) Time traces of phase locking for constant circadian frequency (i.e. $2\pi/24 \text{ h}^{-1}$) but different natural frequencies of the cell cycle ((i) $\omega_{cdk} \approx 2\pi/24 \text{ h}^{-1}$, (ii) $\omega_{cdk} \approx 2\pi/32 \text{ h}^{-1}$, (iii) $\omega_{cdk} \approx 2\pi/19 \text{ h}^{-1}$). A coupling strength of $A_{cdk} = 8$ was used in H and I.

approach by linking the cell cycle to the 24h circadian clock. Expression of the Wee1 kinase is not only 461 affected by DNA damage but also tightly regulated by the Clock-Bmall transcription factor complex, a 462 master regulator of the circadian clock. This couples the cell cycle to the circadian clock (Fig 9D) [74]. 463 We explored the dynamics of such unidirectional coupling by introducing the circadian regulation of 464 Wee1 into the model. More specifically, the [Cdk1] activation threshold — which is regulated by Wee1 465 was periodically shifted between its basal [CycB] level and a predefined maximal level by periodically 466 altering parameter α_{cdk} (see Methods). When two oscillators are coupled like this, one can expect to 467 observe so-called p:q phase locking, meaning that p cycles of the cell cycle are completed for q cycles of 468 the circadian clock. Whether such behavior indeed occurs depends on two factors [75]: (1) the relative 469 frequencies of the circadian clock and the unforced cell cycle (i.e. the frequency in the absence of any 470 circadian control, also called the natural frequency), and (2) on the strength of the coupling (here $A_{\rm cdk}$, 471 i.e. the extent to which the [Cdk1] activation threshold is shifted to higher [CycB] levels over one period). 472 When plotting regions of locking as function of the frequencies and coupling strength, one often finds 473 so-called Arnold tongues (see Methods). 474

Typically, Arnold tongues span wedge-shaped regions in parameter space (at least below a certain 475 critical coupling strength): for low coupling strengths, p:q phase locking would only occur when the natural 476 frequency of the cell cycle is close to the frequency of the circadian clock, whereas a larger mismatch 477 between the two frequencies may still result in synchronization if the coupling strength increases. In 478 our case, we do find these regions, but the Arnold tongues solely widen to the left hand side (Fig. 9G). 479 This indicates that the circadian clock only seems capable of lengthening the cell cycle and not shorten 480 it. Indeed, examining the time traces in Fig. 9H, a 24h natural period of the cell cycle is elongated till 481 31h (i), 38h (ii) and 47h (iii) by a circadian rhythm of 31h, 19h and 12h respectively (see S9 Fig for 482 additional time traces). Similarly, in Fig. 9I, a constant circadian rhythm of 24h elongates cell cycles 483 with a natural period of 24h (i), 32h (ii) and 19h (iii) to forced periods of 48h, 48h and 24h, respectively. 484 Considering that the circadian clock in our model only shifts the [Cdk1] activation threshold to higher 485 [CycB] levels, relative to those of the unforced cell cycle, explains why the forced cell cycle period cannot 486 become shorter than the unforced one. Of note, this observation is in agreement with the findings in [76], 487 where unidirectional regulation of Wee1 by the circadian clock was analyzed using a mechanistic model. 488 There too, Arnold tongues were found to only widen to the left, at least when a basal Wee1 synthesis rate 489 was included. 490

Discussion

In an attempt to unravel the complexity of living systems, it is convenient to envision them as a hierarchical 492 structure of interlinked biological 'modules'. Under the premise that the functionalities of these individual 493 modules do not change when combined with each other, knowing their behavior allows to predict the 494 behavior of the overall system [35]. Two proteins (or protein domains), for example, can be combined to 495 generate recombinant proteins such as fused fluorescent reporters. Additionally, transcriptional promoters 496 can be combined in synthetic networks that possess desired dynamical features, such as the oscillatory 497 behavior of the 'representation', which consists of three negative feedback modules [77]. Although many 498 examples of modularity exist, it has been recognized for a long time [78] that complex living systems can 499 not always be understood as the sum of their constituent parts. Indeed, mechanisms have been described 500 that can hamper the validity of a modular approach, such as off-target effects of transcription factors or 501 the competition for cellular resources. At the same time however, methods to overcome these problems in 502 engineered biological circuits have been proposed [35]. 503

To gain insight into the dynamics of individual or interlinked modules that are difficult to understand in an intuitive way, mathematical models can be very useful. As such, they also provide helpful tools to guide the design of synthetic biological systems [79]. One way to set up a mathematical model is to start from known biochemical interactions and use the law of mass action to derive kinetic rate equations for the concentrations of the biochemical components. However, such mechanistic models often contain a large numbers of variables and parameters, many of which can be difficult to measure experimentally. This can impede generalization of the obtained results, which might strongly depend on the particular set

of chosen parameter values. Therefore, techniques for reducing the complexity of mathematical models while preserving their most fundamental characteristics have been a topic of great interest for several decades, ranging from the well-established quasi-steady-state assumption to more advanced mathematical algorithms [80–83].

A conceptually easy approach to reduce the complexity of biochemical equations is to replace the 515 detailed reaction mechanisms of certain biological modules by a mathematical function that explicitly 516 describes their resulting dynamical behavior. Such a functional or phenomenological approach not only 517 reduces the complexity of mathematical models, but it can also be a valuable alternative to model 518 biochemical networks for which the underlying interaction patterns have not been identified completely. 519 Furthermore, parameters of functional modules correspond directly to observable measurements that are 520 often easier to determine experimentally than the values of biochemical rate constants. In our example of a 521 bistable switch, the parameter α correlates with the width of the bistable region. By choosing the function 522 ξ appropriately, the activation and inactivation threshold can thus be set directly in accordance with a 523 response curve obtained in the lab. Moreover, the fact that these features are directly tunable allows 524 us to answer questions such as 'how does the activation threshold influence the period of oscillations?'. 525 In a mechanistic model, answering such questions would depend on knowledge about what biochemical 526 parameters determine the activation threshold, which might be less straightforward. The same argument 527 holds for other properties, such as steepness of a response or a time delay. Thus, a direct intuitive link 528 between the model and experimentally determined response curves exists when using functional modules. 529 This is an important advantage of this strategy over other frameworks that are suitable for modeling 530 biochemical reactions without considering mechanistic details, such as Boolean networks [84], especially 531 for biologists without mathematical background. 532

Functional modules based on the Hill equation have been extensively used to describe steep sigmoidal 533 response curves, which can originate from various biochemical mechanisms (e.g. multisite phosphorylations, 534 cooperative binding events and stoichiometric inhibition). All of the molecular details that may generate 535 such a response are then hidden in a few parameters such as the exponent n and a threshold K. In 536 contrast, S-shaped response curves are typically not included explicitly in mathematical models. An 537 important reason for this might be that an S-shaped response is not a function, because there are multiple 538 output values for a single input. One contribution of this paper is to show that, in fact, a Hill function 539 can be smoothly changed into an S-shaped response curve by making the threshold dependent on the 540 output variable. In this way, the equation 541

$$\frac{dY}{dt} = \frac{X(t-\tau)^n}{(K\xi(Y))^n + X(t-\tau)^n} - Y$$
(3)

provides an easy way to model a system with input X and output Y that includes the three modules of ultrasensitive, S-shaped and time delayed responses. If the function ξ is equal to 1, we recover an ultrasensitive response for the steady state of Y as function of X. By choosing and tuning ξ , many different shapes of S-shaped response curves can be obtained. Whenever an experimentally determined steady state curve is available, the function ξ could also be fitted to such data (similar as what we did when fitting the S-shaped module to the mass-action model). 542

In the first part of the paper, we used this extended Hill function in combination with a negative 548 feedback loop to model the early embryonic cell cycle oscillator and showed how the shape of the steady 549 state response affects the oscillations. For ultrasensitive responses, no oscillations can be observed unless 550 a sufficiently large time delay is present (S2 Fig, [60]). On the other hand, a system containing a S-shaped 551 module readily sustains oscillations. Whether oscillations occur depends on the width of the S-region, 552 and oscillations are more likely if there are time delays in the system. A second bistable switch can also 553 facilitate oscillations, as we showed in a model that incorporates the two mitotic switches in the cell 554 cycle. Furthermore, we demonstrated that the S-shaped module can faithfully reproduce the oscillatory 555 dynamics of more detailed mass-action models. 556

In the second part, we extended the phenomenological model of the early embryonic cell cycle with an additional bistable switch for the G1-S transition. Furthermore, these bistable switches were exploited to model several cell cycle checkpoints, thus making the model representative for the somatic cell cycle. Even if the molecular underpinnings of these checkpoints are not incorporated into the model, the functional 550 approach can give some basic understanding of events such as the restriction point (RP), DNA damage checkpoints and even coupling of the cell cycle with the circadian clock. 562

The restriction point as depicted here corresponds to the classical interpretation where newborn 563 cells have to cross a threshold in order to activate E2F and commit to S-phase. It should be noted 564 that such representation is a simplification, as it is known that additional bistable switches control 565 entry into S-phase [85–87]. Furthermore, some cells retain hyperphosphorylated Rb proteins, which are 566 incapable of inhibiting E2F, after division. These cells can directly start S-phase even in suboptimal 567 conditions, challenging the classical view of the RP [71]. Interestingly, irregular cell cycle progression in 568 which E2F activity remained high could also be observed for several combinations of parameters in our 569 phenomenological model (S7 Fig). It would be interesting to see whether the biochemical mechanisms 570 enabling cells to bypass the RP correspond with changes in functional responses identified here. 571

By implementing DNA damage in G2 via manipulation of the Cdk1 activation threshold, we found 572 that cells enter M-phase with higher CycB-Cdk1 levels after DNA damage in G2 phase. This is in line 573 with recently published experimental results [88]. Whether the subsequent M-phase takes longer than 574 in unperturbed cells, as is predicted by the model, was not assessed in the cited study. If this would 575 not be the case, it could indicate that either CycB degradation is accelerated after DNA damage or 576 that the inactivation threshold too is shifted to higher CycB levels. Another interesting study which 577 looked at the effects of DNA repair mechanisms was published by Chao et al., [89]. There, the authors 578 found that elongation of interphase after DNA damage in G2 depends on the severity of the damage, but 579 is independent of when the cells were damaged within G2. This finding is in contrast with our model 580 predictions that DNA damage in G2 can actually be repaired without interphase elongation. It should 581 be noted that here we supposed that DNA damage can be detected and repaired at any time during 582 G2, not only at a checkpoint at the end of G2 phase. Furthermore, our model does not account for 583 regulatory processes such as decreased CycB synthesis [90] or increased CycB degradation [91], which might 584 explain the discrepancies. Even if we cannot account for molecular details and nuances, the possibility to 585 incorporate DNA damage in a phenomenological manner can be helpful to understand these events in a 586 dynamical way. 587

Although in this work we combined the S-shaped/ultrasensitive/delay modules to generate a phe-588 nomenological model of the overall cell cycle, these modules are also applicable for modelling other 589 processes based on their experimentally determined response curves. Bistable switches for example, are 590 known to play a role during epithelial-to-mesenchymal transition [92] and differentiation in cell types 591 ranging from embryonic stem cells [93] till osteogenic precursors [94] and Drosophila eves [95]. Furthermore, 592 the convenient way of combining several functional modules can be exploited to reconstitute interaction 593 patterns with complex dynamical behavior, such as a combination of the 'representation' with bistable 594 switches that has been observed during neural tube development [96]. 595

The modeling strategy based on functional modules put forth in this work fits in the mindset of reducing the complexity of mathematical models and has the advantage of preserving an intuitive link with the experimentally measured response curves. Furthermore, the modular approach provides the flexibility for combining the modules in a multitude of different ways. As such, it can provide a toolbox for other researchers who want to generate phenomenological modular models, not only of the cell cycle but also of many other biological processes.

Materials and methods

Converting an ultrasensitive function into a S-shaped one

Starting from an (increasing) ultrasensitive response curve, one can obtain a S-shaped response by shifting the lower and upper part of the curve to the right and left respectively. For APC/C, the ultrasensitive response curve can be expressed as a Hill equation representing the fraction of activated APC/C to total APC/C molecules as a function of [Cdk1] (Fig. 10A):

$$[APC]^* = \frac{[APC/C]}{[APC/C]_{tot}} = \frac{[Cdk1]^n}{K_{cdk,apc}^n + [Cdk1]^n}$$
(4)

602



Fig 10. An ultrasensitive response curve (A) can be converted into a S-shaped one (D) by inverting the product (E) of a scaling function (C) with the inverted ultrasensitive response (B).

The S-shaped $[APC]^*$ response cannot be expressed as a function of [Cdk1] (since it would have multiple $[APC]^*$ values for a certain range of [Cdk1]). However, even if the function is S-shaped, there would be only one [Cdk1] value for each $[APC]^*$ value. This motivates us to invert the function (Fig. 10B): 610

$$[\mathrm{Cdk1}] = K_{\mathrm{cdk,apc}} \left(\frac{[\mathrm{APC}]^*}{1 - [\mathrm{APC}]^*}\right)^{1/\tau}$$

Subsequently, this inverted response can be multiplied by the scaling function $\xi([APC]^*)$ (Fig. 10C), ⁶¹¹ such that [Cdk1] values increase for low [APC]^{*} values and decrease for high [APC]^{*} (Fig. 10E): ⁶¹²

$$[Cdk1] = \xi([APC]^*) \cdot K_{cdk,apc} \left(\frac{[APC]^*}{1 - [APC]^*}\right)^{1/n}$$
(5)

with $\xi([APC]^*) = 1 + \alpha_{apc} \cdot [APC]^* ([APC]^* - 1) ([APC]^* - r)$. Rearranging Eq. 5 leads to an expression for a S-shaped response (Fig. 10D), analogous with Eq. 1 in the main text. This is not a function, but the expression can be put in a differential equation such that the steady state of this equation follows the S-shaped response:

$$[APC]^* = \frac{[Cdk1]^n}{(\xi([APC]^*) \cdot K_{cdk,apc})^n + [Cdk1]^n}$$

$$\Rightarrow \frac{d[APC]^*}{dt} = \frac{1}{\epsilon_{apc}} \left(\frac{[Cdk1]^n}{(\xi([APC]^*) \cdot K_{cdk,apc})^n + [Cdk1]^n} - [APC]^* \right)$$
(6)

A similar approach can be followed for the bistable switch from total CycB-Cdk1 (i.e. [CycB]) to active CycB-Cdk1 (i.e. [Cdk1]). The ultrasensitive response is given by

$$\frac{[\text{Cdk1}]}{[\text{CycB}]} = \frac{[\text{CycB}]^n}{K_{\text{cyc,cdk}}^n + [\text{CycB}]^n}$$
(7)

Rearranging and multiplying by $\xi([Cdk1]/[CycB]) = 1 + \alpha_{cdk} \cdot [Cdk1]/[CycB] ([Cdk1]/[CycB] - 1) ([Cdk1]/[CycB] - r)$ 619 gives:

$$[CycB] = \xi ([Cdk1]/[CycB]) \cdot K_{cyc,cdk} \left(\frac{[Cdk1]/[CycB]}{1 - [Cdk1]/[CycB]}\right)^{1/n}$$

$$\Rightarrow \frac{[Cdk1]}{[CycB]} = \frac{[CycB]^n}{(\xi ([Cdk1]/[CycB]) \cdot K_{cyc,cdk})^n + [CycB]^n}$$
(8)

This means that

$$[Cdk1] = \frac{[CycB]^n}{(\xi([Cdk1]/[CycB]) \cdot K_{cyc,cdk})^n + [CycB]^n} [CycB]$$
(9)

Again, this is not a function, but can be made the steady state of a differential equation (system (iii) in the main text):

$$\frac{d[\mathrm{Cdk1}]}{dt} = \frac{1}{\epsilon_{\mathrm{cdk}}} \left(\frac{[\mathrm{CycB}]^n}{(\xi([\mathrm{Cdk1}]/[\mathrm{CycB}]) \cdot K_{\mathrm{cyc,cdk}})^n + [\mathrm{CycB}]^n} [\mathrm{CycB}] - [\mathrm{Cdk1}] \right)$$

It needs to be emphasized that both Eq. 4 and 7 describe an ultrasensitive response of which the 622 output is given by the concentration level of a compound Y, normalized for the total amount $Y_{\rm tot}$. 623 Accordingly, both expressions are limited to the interval $Y/Y_{\text{tot}} \in [0, 1]$. Similarly, Eq. 6 and 8 describe 624 S-shaped responses for $Y/Y_{\text{tot}} \in [0, 1]$. When incorporating these steady state expressions into the 625 differential equations however, a distinction should be made. As $[APC/C]_{tot}$ is a constant, the ratio 626 $[APC]^* = [APC/C]/[APC/C]_{tot}$ can directly be incorporated into the ODE for the rate of change of $[APC]^*$. In 627 contrast, [CycB] is not a constant and therefore, the rate of change of ^[Cdk1]/_[CycB] would need to be given 628 by the quotient rule for differentiation, as both numerator and denominator are variables. Alternatively, 629 [CycB] can first be moved to the right hand side of Eq. 8, resulting in Eq. 9. Subsequently, this expression 630 for the steady state of [Cdk1] can be incorporated into an ODE describing the rate of change of [Cdk1]. 631 Of note, Eq. 9 does no longer saturate at [Cdk1] = 1 (as was the case for Eq. 8 at [Cdk1]/[CycB] = 1) but 632 keeps rising indefinitely (see for example Fig. 7C,E). 633

Non-dimensionalization of system equations

To facilitate mathematical analysis, the system equations for the two-variable models described in the main text were non-dimensionalized by introducing the new variables:

$$[\mathrm{Cdk1}]^* = \frac{[\mathrm{Cdk1}]}{K_{\mathrm{cdk,apc}}} \qquad [\mathrm{APC}]^* = \frac{[\mathrm{APC/C}]}{[\mathrm{APC/C}]_{\mathrm{tot}}} \qquad t^* = b_{\mathrm{deg}} \cdot t \qquad \tau^* = b_{\mathrm{deg}} \cdot \tau$$

where $b_{\text{deg}} \text{[min}^{-1]}$ is the reaction rate of the apparent first order degradation of CycB-Cdk1 complexes (i.e. variable [Cdk1]) by [APC]*. As an example, system (ii) then becomes:

$$\begin{cases} \frac{d[\text{Cdk1}]^{*}}{dt^{*}} = \frac{b_{\text{syn}}}{b_{\text{deg}} \cdot K_{\text{cdk,apc}}} - [\text{Cdk1}]^{*} \cdot [\text{APC}]^{*} \\ \frac{d[\text{APC}]^{*}}{dt^{*}} = \frac{1}{\epsilon_{\text{apc}} \cdot b_{\text{deg}}} \left(\frac{[\text{Cdk1}]^{*n}(t^{*} - \tau^{*})}{\xi([\text{APC}]^{*})^{n} + [\text{Cdk1}]^{*n}(t^{*} - \tau^{*})} - [\text{APC}]^{*} \right) \end{cases}$$
(10)

after which we can define $c = \frac{b_{\text{syn}}}{b_{\text{deg}} \cdot K_{\text{cdk,apc}}}, \epsilon_{\text{apc}}^* = \epsilon_{\text{apc}} \cdot b_{\text{deg}}$ and $\xi([\text{APC}]^*)$ remains unaffected:

$$\xi([\mathrm{APC}]^*) = 1 + \alpha_{\mathrm{apc}} \cdot [\mathrm{APC}]^* ([\mathrm{APC}]^* - 1) ([\mathrm{APC}]^* - r)$$

Standard parameter values used are: $\epsilon_{apc}^* = 0.01$, n = 15, $b_{deg} = 0.1 \text{ min}^{-1}$, $K_{cdk,apc} = 20 \text{ nM}$, of which the latter three are based on experimental observations in *Xenopus laevis* eggs [12, 15, 30]. Obtained simulation results were scaled back to dimensional values where possible. For APC/C however, no experimental estimates for the total amount [APC/C]_{tot} were found and results are presented as [APC]^{*}.

The system containing two S-shaped response (Eq. system (iii)) was non-dimensionalized in a similar way:

$$[CycB]^* = \frac{[CycB]}{K_{cyc,cdk}} \qquad [Cdk1]^* = \frac{[Cdk1]}{K_{cdk,apc}} \qquad [APC]^* = \frac{[APC/C]}{[APC/C]_{tot}} \qquad t^* = b_{deg} \cdot t$$

$$\frac{d[\operatorname{CycB}]^*}{dt^*} = \frac{b_{\operatorname{syn}}}{b_{\operatorname{deg}} \cdot K_{\operatorname{cyc,cdk}}} - [\operatorname{CycB}]^* \cdot [\operatorname{APC}]^*$$

$$\begin{cases}
\frac{d[\operatorname{Cdk1}]^{*}}{dt^{*}} = \frac{1}{\epsilon_{\operatorname{cdk}} \cdot b_{\operatorname{deg}}} \left(\frac{K_{\operatorname{cyc,cdk}}}{K_{\operatorname{cdk,apc}}} \cdot \frac{[\operatorname{CycB}]^{*n+1}}{\xi \left([\operatorname{Cdk1}]^{*}/d \cdot [\operatorname{CycB}]^{*} \right)^{n} + [\operatorname{CycB}]^{*n}} - [\operatorname{Cdk1}]^{*} \right) \\
\frac{d[\operatorname{APC}]^{*}}{dt^{*}} = \frac{1}{\epsilon_{\operatorname{apc}} \cdot b_{\operatorname{deg}}} \left(\frac{[\operatorname{Cdk1}]^{*n}}{\xi ([\operatorname{APC}]^{*})^{n} + [\operatorname{Cdk1}]^{*n}} - [\operatorname{APC}]^{*} \right)
\end{cases} \tag{11}$$

If $\frac{K_{\text{cyc,cdk}}}{K_{\text{cdk,apc}}} = d$, we get $\frac{b_{\text{syn}}}{b_{\text{deg}} \cdot K_{\text{cyc,cdk}}} = \frac{c}{d}$. For the scaling functions we have $\xi([\text{APC}]^*)$ as before and

$$\xi([\mathrm{Cdk1}]^*/d\cdot[\mathrm{CycB}]^*) = 1 + \alpha_{\mathrm{cdk}} \cdot \frac{[\mathrm{Cdk1}]^*}{d\cdot[\mathrm{CycB}]^*} \left(\frac{[\mathrm{Cdk1}]^*}{d\cdot[\mathrm{CycB}]^*} - 1\right) \left(\frac{[\mathrm{Cdk1}]^*}{d\cdot[\mathrm{CycB}]^*} - r\right)$$

The same standard parameter values as the two-dimensional system were used, with the additional parameters being: $b_{\text{deg}}\epsilon_{\text{cdk}} = b_{\text{deg}}\epsilon_{\text{apc}} = 0.01$ and $K_{\text{cyc,cdk}} = 40$ nM [11].

Conversion of parameter α to the width of the S-shaped region

644

641

All screens for which the width of the bistable region was altered, were performed by screening different values of α and linking this value to the width of the S-shaped region. The expression for the non-dimensionalized S-shaped response curve, i.e.

$$y = \frac{x^n}{\xi(y)^n + x^n} \qquad \text{with} \begin{cases} x = [\text{Cdk1}]^*, y = [\text{APC}]^* \\ x = [\text{CycB}]^*, y = [\text{Cdk1}]^* / d \cdot [\text{CycB}]^* \end{cases}$$

can be inverted, resulting in:

$$x = \xi(y) \left(\frac{y}{1-y}\right)^{1/n} \qquad \text{with} \begin{cases} x = [\operatorname{Cdk1}]^*, y = [\operatorname{APC}]^* \\ x = [\operatorname{CycB}]^*, y = [\operatorname{Cdk1}]^*/d \cdot [\operatorname{CycB}]^* \end{cases}$$

The width of the S-shaped region is given by the difference of x-values at the extrema of this inverted S-shaped response, which can be calculated as the roots of the derivative. For the cubic scaling function $\xi(y) = 1 + \alpha \left[y^3 - (1+r)y^2 + r \cdot y\right]$, we have:

$$\begin{aligned} \frac{dx}{dy} &= \frac{d\xi}{dy} \cdot \left(\frac{y}{1-y}\right)^{1/n} + \xi(y) \frac{d}{dy} \left[\left(\frac{y}{1-y}\right)^{1/n} \right] \\ &= \alpha \left[3y^2 - 2(1+r)y + r \right] \left(\frac{y}{1-y}\right)^{1/n} \\ &+ \left[1 + \alpha y(y-1)(y-r) \right] \frac{1}{n(1-y)^2} \left(\frac{y}{1-y}\right)^{\frac{1-n}{n}} \end{aligned}$$

The roots of this function on the interval $y \in [0, 1]$ were calculated numerically and used to determine the values of x at the extrema of the inverted S-shaped response. Either two extrema were found for which x > 0 (this was ensured by the choice of parameter α , see S1 Text), in which case the width of the S-shaped response curve was determined by their difference, or no extrema were found, in which case the width equaled zero. 649 640 649 649 649 649 649 649 649 649

Mass-action model for the PP2A-ENSA-GWL network

The model of the PP2A-ENSA-GWL network was largely based on the network described in [61], where ⁶⁵¹ the double negative feedback between GWL and PP2A can give rise to bistability. GWL indirectly ⁶⁵²

inhibits PP2A by phosphorylating ENSA, which is both a substrate and inhibitor of PP2A and binds it in a complex C [62]. Here, the model was converted into an oscillator by incorporating synthesis and degradation of [Cdk1]:

$$\begin{cases} \frac{d[\operatorname{Cdk1}]}{dt} = b_{\operatorname{syn}} - b_{\operatorname{deg}}[\operatorname{Cdk1}][\operatorname{APC}]^{*} \\ \frac{d[\operatorname{GWLp}]}{dt} = k_{\operatorname{pg}}[\operatorname{GWL}][\operatorname{Cdk1}] - k_{\operatorname{dg}}[\operatorname{GWLp}][\operatorname{PP2A}] \\ \frac{d[\operatorname{C}]}{dt} = k_{\operatorname{ass}}[\operatorname{ENSAp}][\operatorname{PP2A}] - (k_{\operatorname{dis}} + k_{\operatorname{cat}})[\operatorname{C}] \\ \frac{d[\operatorname{ENSAp}]}{dt} = k_{\operatorname{dis}}[\operatorname{C}] + k_{\operatorname{pe}}[\operatorname{ENSA}][\operatorname{GWLp}] - k_{\operatorname{ass}}[\operatorname{ENSAp}][\operatorname{PP2A}] \\ \frac{d[\operatorname{APC}]^{*}}{dt} = k_{\operatorname{pa}}(1 - [\operatorname{APC}]^{*})[\operatorname{Cdk1}] - k_{\operatorname{da}}[\operatorname{APC}]^{*}[\operatorname{PP2A}] \end{cases}$$
(12)

with conservation of mass giving:

$$\begin{split} [GWL] &= [GWL_{tot}] - [GWLp] \\ [PP2A] &= [PP2A_{tot}] - [C] \\ [ENSA] &= [ENSA_{tot}] - [ENSAp] - [C] \end{split}$$

Parameter values were manually screened to obtain a steady-state response curve centered around [Cdk1] ≈ 20 nM (to be in line with $K_{\rm cdk,apc} = 20$ nM used elsewhere in this paper) and obtain oscillations with biologically relevant periods (Table 1). Steady-state response curves were determined via a custom Python script performing numerical continuation [97].

			_
$k_{ m pg}$	0.03	1/(nM min)	Phosphorylation constant of GWL
$k_{ m dg}$	3.16	1/(nM min)	Dephosphorylation constant of GWL
$k_{ m pe}$	3.16	1/(nM min)	Phosphorylation constant of ENSA
$k_{ m pa}$	0.63	1/(nM min)	Phosphorylation constant of APC/C
$k_{ m dis}$	1	$1/\min$	Dissociation constant of ENSA-PP2A complex C
k_{cat}	3.98	$1/\min$	Catalytic constant of ENSA-PP2A complex C
$\mathrm{GWL}_{\mathrm{tot}}$	20	nM	Total GWL concentration
$\mathrm{ENSA}_{\mathrm{tot}}$	40	nM	Total ENSA concentration
$PP2A_{tot}$	20	nM	Total PP2A concentration

 Table 1. Default parameter values for the mass-action model of the PP2A-ENSA-GWL network.

 Parameter
 Value
 Units
 Description

Fitting the S-shaped module to the mass-action model

Once the steady-state response of the mass-action model was determined, we manually fitted a piecewise linear scaling function $\xi([APC]^*)$ for use in system (i-b) (after non-dimensionalization).

$$\xi([APC]^*) = \begin{cases} \frac{\xi_{RF} - 1}{[APC]^*_{RF}} [APC]^* + 1 & \text{if } [APC]^* \leq [APC]^*_{RF} \\ \frac{\xi_{LF} - \xi_{RF}}{[APC]^*_{LF} - [APC]^*_{RF}} ([APC]^* - [APC]^*_{RF}) + \xi_{RF} & \text{if } [APC]^*_{RF} < [APC]^* \leq [APC]^*_{LF} \\ \frac{1 - \xi_{LF}}{1 - [APC]^*_{LF}} ([APC]^* - [APC]^*_{LF}) + \xi_{LF} & \text{if } [APC]^*_{LF} < [APC]^* \end{cases}$$

Here the subscripts RF and LF indicate right and left fold respectively (similar as in Fig. 4). For Fig. 5D, ⁶⁶¹ the best fit was obtained for $[APC]^*_{LF} = 0.65$, $[APC]^*_{RF} = 0.39$, $\xi_{LF} = 0.76$, and $\xi_{RF} = 1.25$. In Fig. ⁶⁶² 5G, the chosen values were $[APC]^*_{LF} = 0.39$, $[APC]^*_{RF} = 0.16$, $\xi_{LF} = 0.73$, and $\xi_{RF} = 1.62$. In each ⁶⁶³

case, the location of the upper branch was manually shifted by multiplying the Hill-like term in system (i-b) (with n = 5) by a correction factor equaling 1 for Fig. 5D and 0.75 for Fig. 5G. For the time traces, $\epsilon_{apc}^* = 0.1$ gave the best correspondence with the mass-action model.

System equations for interlinked switches

The equations for the interlinked switches were derived following the same principles as described before, resulting in:

$$\begin{cases}
\frac{d[\operatorname{CycD}]}{dt} = d_{\operatorname{syn}} - d_{\operatorname{deg}}[\operatorname{CycD}]([\operatorname{APC}]^* + \delta_d) \\
\frac{d[\operatorname{E2F}]^*}{dt} = \frac{1}{\epsilon_{\operatorname{e2f}}} \left(\frac{[\operatorname{CycD}]^n}{(\xi([\operatorname{E2F}]^*) \cdot K_{\operatorname{cyc,e2f}})^n + [\operatorname{CycD}]^n} - [\operatorname{E2F}]^* \right) \\
\frac{d[\operatorname{CycB}]}{dt} = b_{\operatorname{syn}} \cdot [\operatorname{E2F}]^* - b_{\operatorname{deg}}[\operatorname{CycB}]([\operatorname{APC}]^* + \delta_b) \\
\frac{d[\operatorname{Cdk1}]}{dt} = \frac{1}{\epsilon_{\operatorname{cdk}}} \left(\frac{[\operatorname{CycB}]^n}{(\xi([\operatorname{Cdk1}]/[\operatorname{CycB}]) \cdot K_{\operatorname{cyc,cdk}})^n + [\operatorname{CycB}]^n} [\operatorname{CycB}] - [\operatorname{Cdk1}] \right) \\
\frac{d[\operatorname{APC}]^*}{dt} = \frac{1}{\epsilon_{\operatorname{apc}}} \left(\frac{[\operatorname{Cdk1}]^n}{(\xi([\operatorname{APC}]^*) \cdot K_{\operatorname{cdk,apc}})^n + [\operatorname{Cdk1}]^n} - [\operatorname{APC}]^* \right)
\end{cases}$$
(13)

with

$$\begin{split} \xi([\text{APC}]^*) &= 1 + \alpha_{\text{apc}}[\text{APC}]^* ([\text{APC}]^* - 1)([\text{APC}]^* - r) \\ \xi([\text{Cdk1}]/[\text{CycB}]) &= 1 + \alpha_{\text{cdk}} \cdot \frac{[\text{Cdk1}]}{[\text{CycB}]} \left(\frac{[\text{Cdk1}]}{[\text{CycB}]} - 1\right) \left(\frac{[\text{Cdk1}]}{[\text{CycB}]} - r\right) \\ \xi([\text{E2F}]^*) &= 1 + \alpha_{\text{e2f}}[\text{E2F}]^* ([\text{E2F}]^* - 1)([\text{E2F}]^* - r) \end{split}$$

As before, $[APC]^* = [APC/C]/[APC/C]_{tot}$ and similarly $[E2F]^* = [E2F]/[E2F]_{tot}$. For all other variables and parameters, the original dimensions were retained. Default parameter values were manually screened so that oscillations with biologically relevant periods were obtained (Table 2). The threshold value $K_{cyc,e2f}$ at which [CycD] activates $[E2F]^*$ was assumed to be 3 times higher than the threshold $K_{cyc,cdk}$ at which [CycB] activates [Cdk1], i.e. 120 nM, based on simulations from [56] where CycD levels are about three times higher than CycB levels and on simulations from [19] where CycD levels rise up to ~ 100 nM.

To distinguish the different cell cycle phases in this model, cells were considered to be in M-phase wherever $[APC]^* > 0.95$, in G1 wherever $[E2F]^* < 0.95$ and $[APC]^* < 0.95$, and in S/G2 wherever $[E2F]^* > 0.95$ and $[APC]^* < 0.95$.

To model the effect of the restriction point, reduced levels of external growth factors were incorporated 679 into the model by reducing the synthesis rate $d_{\rm syn}$ of [CycD] by a factor 10. This number was chosen 680 based on experimental measurements that demonstrated how growth factor stimulation increases CycD 681 levels between 4 and 20 fold [98,99]. DNA damage during G1 phase was accounted for by increasing 682 the basal degradation rate of [CycD] (i.e. δ_d) by a factor 3 and damage in G2 phase was modeled by 683 increasing the width of the bistable [Cdk1] response (i.e. $\alpha_{cdk} = 30$) to shift the right fold of the curve to 684 higher [CycB] levels. The value of $\alpha_{cdk} = 30$ was chosen so that the steady state of [CycB] in the standard 685 model is below the [Cdk1] activation threshold. 686

Simulations and analysis

All simulations were performed in Python 3.7. Ordinary differential equations were solved using the Python Scipy package solve_ivp (method = Radau). For equations including a delay, the JITCDDE package was used [100]. The amplitudes and periods were numerically determined from the extrema in the time series via a custom Python script. Small amplitude oscillations and damped oscillations were omitted from the analysis.

687

Parameter	Value	\mathbf{Units}	Description			
r	0.5		As defined in ξ			
$\alpha_{\rm e2f}$	5		As defined in ξ			
$\alpha_{\rm cdk}$	5		As defined in ξ			
$\alpha_{\rm apc}$	5		As defined in ξ			
n^{-}	15		Hill coefficient			
ϵ_{e2f}	0.01		Time constant E2F reaction			
$\epsilon_{ m cdk}$	0.01		Time constant Cdk1 reaction			
$\epsilon_{ m apc}$	0.01		Time constant APC/C reaction			
δ_d	0.05		Basal CycD degradation			
δ_b	0.05		Basal CycB degradation			
$K_{\rm cyc,e2f}$	120	nM	Threshold of E2F activation by CycD			
$K_{\rm cyc,cdk}$	40	nM	Threshold of Cdk1 activation by CycB			
$K_{\rm cdk,apc}$	20	nM	Threshold of APC/C activation by Cdk1			
$d_{\rm syn}$	0.15	nM/min	Synthesis rate of CycD			
$d_{ m deg}$	0.009	$1/\min$	Apparent first order degradation of CycD			
$b_{\rm syn}$	0.03	nM/min	Synthesis rate of CycB			
$b_{ m deg}$	0.003	$1/\min$	Apparent first order degradation of CycB			

	Demonster	X 7-1	T T ! +	D!-				
Table 2.	Default parameter	values for	modeling	the cell cyc	cle as i	nterlinked	bistable	switches.

Arnold tongues and phase locking

Phase locking occurs when the ratio of periods (or frequencies) from two oscillators equals a rational number p:q, meaning that p cycles of one oscillator are completed while the second oscillator completes q cycles. To check for phase locking between the cell cycle and circadian clock, the system of interlinked switches (Eq. 13) was reanalyzed (again with parameters from Table 2), but this time the prefactor α_{cdk} in $\xi([Cdk1]/[CycB])$ was periodically shifted between its basal level and a predefined maximal level (i.e. $\alpha_{cdk} + 2A_{cdk}$) at a forcing frequency $\omega_{circadian}$ ranging from 1/3 to 3 times the natural frequency of the [Cdk1] oscillations:

$$\xi([Cdk1]/[CycB]) = 1 + \alpha_{cdk}^* \frac{[Cdk1]}{[CycB]} \left(\frac{[Cdk1]}{[CycB]} - 1\right) \left(\frac{[Cdk1]}{[CycB]} - r\right)$$

with $\alpha_{cdk}^* = [\alpha_{cdk} + A_{cdk} + A_{cdk} \sin(\omega_{circadian}t)]$. Custom Python code was used to find a repeating pattern in the forced [Cdk1] oscillations, based on the difference between the simulated time series and a time shifted version of itself (similar to calculating the autocorrelation). The ratio of the periods from this repeating pattern and the circadian clock was compared with p:q ratios for p and q in [1,2,3,4,5] to decide whether phase locking occurred.

Supporting information

S1 Text. Supplementary information. This file contains additional mathematical analysis of the models and the supplemental figures listed below. 701

S1 Fig Effect of parameters on the scaling function and system response. Here we show in more detail how the different parameters affect the shape of the scaling function ξ and the overall dynamics of the system. 702

S2 Fig Oscillations for a time delayed, ultrasensitive cell cycle model. Ultrasensitivity by 705 itself cannot induce oscillatory behavior of a two-dimensional system, i.e. either bistability needs to be 706 incorporated or a sufficiently large time delay. In the main text, we focused on the effect of bistability, 707 whereas here we summarize how ultrasensitivity in combination with a time delay can sustain oscillations. 708

693

S3 Fig Oscillation period and amplitude for the bistable module. The period and amplitudes $_{709}$ of [Cdk1] and [APC]* oscillations for system (i-b) as a function of the relative synthesis *c* and the width $_{710}$ of the bistable response curve. Related to Fig 3 in the main text. $_{711}$

S4 Fig Alternative definitions of the scaling function ξ . Definitions for the scaling function ξ other than a cubic function can be used. The concept of 'symmetric' and 'asymmetric' bistable nullclines as used in the main text is visualised based on some alternative ξ functions.

S5 Fig Oscillation period and amplitude for the delayed bistable module. The period and 715 amplitudes of [Cdk1] and [APC]* oscillations for system (ii) as a function of the width of the bistable 716 response curve and the time delay. Related to Fig 6 in the main text. 717

S6 Fig The width of the bistable modules in the three dimensional model affects the amplitude of the oscillations. In Fig 7 in the main text we showed how the [Cdk1] amplitudes change values K of the bistable switches. Here, we compare these results for different vidths of the bistable response curves and show how the amplitude is proportional to the bistable width. The second seco

S7 Fig Irregular cell cycle oscillations in a chain of bistable switches. In Fig 8 in the main text we indicated grey regions in parameter space for which irregular oscillations were observed. Here, we show time traces of [E2F]* for such irregular patterns.

S8 Fig Effect of synthesis and degradation rates of CycD and CycB on the duration of different cell cycle phases. In Fig 8 in the main text we showed the effect of changing synthesis and degradation rates on the overall length of the cell cycle. Here, we separate the effects on the different cell cycle phases.

S9 Fig Phase locking between the circadian clock and the cell cycle. Time traces showing 729 the absence or presence of p:q phase locking between the cell cycle and the circadian clock for several 730 parameter combinations. Related to the Arnold tongues shown in Fig 9. 731

S1 Video The cell cycle can be represented as a chain of interlinked bistable switches. ⁷³² Video showing how the cell cycle progresses through the different bistable switches as shown in Fig 8. ⁷³³

Acknowledgments

We are grateful to Prof. Catherine Verfaillie and the members of the Gelens lab for useful comments on the manuscript. 735

Funding

This work was supported by the Research Foundation Flanders (FWO, www.fwo.be) with individual r38 support to J.D.B. (1189120N) and J.R. (11D0920N) and project support to L.G. (Grant GOA5317N) and r39 the KU Leuven Research Fund (No. C14/18/084) to L.G. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. 741

Competing interests

The authors have declared that no competing interests exist.

734

737

742

References

- Novák B, Sible JC, Tyson JJ. Checkpoints in the Cell Cycle. In: Encycl. Life Sci. Chichester, UK: John Wiley & Sons, Ltd; 2003.Available from: http://doi.wiley.com/10.1038/npg.els. 0001355.
- Hartwell LH, Weinert TA. Checkpoints: Controls that ensure the order of cell cycle events. Science (80-). 1989;246(4930):629–634. doi:10.1126/science.2683079.
- 3. Murray AW, Kirschner MW. Dominoes and clocks: The union of two views of the cell cycle. Science (80-). 1989;246(4930):614-621. doi:10.1126/science.2683077.
- 4. Verdugo A, Vinod PK, Tyson JJ, Novák B. Molecular mechanisms creating bistable switches at cell cycle transitions. Open Biol. 2013;3(3):120179. doi:10.1098/rsob.120179.
- 5. Stallaert W, Kedziora KM, Chao HX, Purvis JE. Bistable switches as integrators and actuators during cell cycle progression. FEBS Lett. 2019;593(20):2805-2816. doi:10.1002/1873-3468.13628.
- Tyson JJ, Csikasz-Nagy A, Novák B. The dynamics of cell cycle regulation. BioEssays. 2002;24(12):1095–1109. doi:10.1002/bies.10191.
- Solomon MJ, Glotzer M, Lee TH, Philippe M, Kirschner MW. Cyclin activation of p34cdc2. Cell. 1990;63(5). doi:10.1016/0092-8674(90)90504-8.
- 8. Murray AW. Turning on mitosis. Curr Biol. 1993;3(5). doi:10.1016/0960-9822(93)90182-N.
- Thron CD. A model for a bistable biochemical trigger of mitosis. Biophys Chem. 1996;57(2-3):239–251. doi:10.1016/0301-4622(95)00075-5.
- 10. Novák B, Tyson JJ. Numerical analysis of a comprehensive model of M-phase control in Xenopus oocyte extracts and intact embryos. J Cell Sci. 1993;106(4).
- 11. Pomerening JR, Sontag ED, Ferrell Jr JE. Building a cell cycle oscillator: Hysteresis and bistability in the activation of Cdc2. Nat Cell Biol. 2003;5(4):346–351. doi:10.1038/ncb954.
- Sha W, Moore J, Chen K, Lassaletta AD, Yi CS, Tyson JJ, et al. Hysteresis drives cell-cycle transitions in Xenopus laevis egg extracts. Proc Natl Acad Sci U S A. 2003;100(3):975–980. doi:10.1073/pnas.0235349100.
- Pomerening JR, Sun YK, Ferrell Jr JE. Systems-level dissection of the cell-cycle oscillator: Bypassing positive feedback produces damped oscillations. Cell. 2005;122(4):565–578. doi:10.1016/j.cell.2005.06.016.
- 14. Rata S, Suarez Peredo Rodriguez MF, Joseph S, Peter N, Echegaray Iturra F, Yang F, et al. Two Interlinked Bistable Switches Govern Mitotic Control in Mammalian Cells. Curr Biol. 2018;28(23):3824–3832.e6. doi:10.1016/j.cub.2018.09.059.
- 15. Kamenz J, Gelens L, Ferrell Jr JE. Bistable, Biphasic Regulation of PP2A-B55 Accounts for the Dynamics of Mitotic Substrate Phosphorylation. Curr Biol. 2020;doi:10.1016/j.cub.2020.11.058.
- 16. Mochida S, Rata S, Hino H, Nagai T, Novák B. Two Bistable Switches Govern M Phase Entry. Curr Biol. 2016;26(24):3361–3367. doi:10.1016/j.cub.2016.10.022.
- 17. Heim A, Konietzny A, Mayer TU. Protein phosphatase 1 is essential for Greatwall inactivation at mitotic exit. EMBO Rep. 2015;16(11):1501–1510. doi:10.15252/embr.201540876.
- Holder J, Poser E, Barr FA. Getting out of mitosis: spatial and temporal control of mitotic exit and cytokinesis by PP1 and PP2A. FEBS Lett. 2019;593(20):2908–2924. doi:10.1002/1873-3468.13595.

- 19. Yao G, Lee TJ, Mori S, Nevins JR, You L. A bistable Rb-E2F switch underlies the restriction point. Nat Cell Biol. 2008;10(4):476–482. doi:10.1038/ncb1711.
- 20. Blagosklonny MV, Pardee AB. The restriction point of the cell cycle. Cell Cycle. 2002;1(2):102–109. doi:10.4161/cc.1.2.108.
- 21. Collin P, Nashchekina O, Walker R, Pines J. The spindle assembly checkpoint works like a rheostat rather than a toggle switch. Nat Cell Biol. 2013;15(11):1378–1385. doi:10.1038/ncb2855.
- Dick AE, Gerlich DW. Kinetic framework of spindle assembly checkpoint signalling. Nat Cell Biol. 2013;15(11):1370–1377. doi:10.1038/ncb2842.
- Holt LJ, Krutchinsky AN, Morgan DO. Positive feedback sharpens the anaphase switch. Nature. 2008;454(7202):353–357. doi:10.1038/nature07050.
- 24. Kuhn J, Dumont S. Mammalian kinetochores count attached microtubules in a sensitive and switch-like manner. J Cell Biol. 2019;218(11):3583–3596. doi:10.1083/JCB.201902105.
- Rattani A, Vinod PK, Godwin J, Tachibana-Konwalski K, Wolna M, Malumbres M, et al. Dependency of the spindle assembly checkpoint on Cdk1 renders the anaphase transition irreversible. Curr Biol. 2014;24(6):630–637. doi:10.1016/j.cub.2014.01.033.
- Novák B, Tyson JJ. Mechanisms of signalling-memory governing progression through the eukaryotic cell cycle. Curr Opin Cell Biol. 2021;69:7–16. doi:10.1016/j.ceb.2020.12.003.
- Hutter LH, Rata S, Hochegger H, Novák B. Interlinked bistable mechanisms generate robust mitotic transitions. Cell Cycle. 2017;16(20):1885–1892. doi:10.1080/15384101.2017.1371885.
- Murray AW, Solomon MJ, Kirschner MW. The role of cyclin synthesis and degradation in the control of maturation promoting factor activity. Nature. 1989;339(6222):280–286. doi:10.1038/339280a0.
- King RW, Deshaies RJ, Peters JM, Kirschner MW. How proteolysis drives the cell cycle. Science (80-). 1996;274(5293):1652–1659. doi:10.1126/science.274.5293.1652.
- 30. Yang Q, Ferrell Jr JE. The Cdk1-APC/C cell cycle oscillator circuit functions as a time-delayed, ultrasensitive switch. Nat Cell Biol. 2013;15(5):519-525. doi:10.1038/ncb2737.
- 31. Alfieri C, Zhang S, Barford D. Visualizing the complex functions and mechanisms of the anaphase promoting complex/cyclosome (APC/C). Open Biol. 2017;7(11). doi:10.1098/rsob.170204.
- 32. Thomas RRR, D'Ari R. Biological feedback. Taylor & Francis Inc; 1990.
- Padgett J, Santos SDM. From clocks to dominoes: lessons on cell cycle remodelling from embryonic stem cells. FEBS Lett. 2020;594(13):2031–2045. doi:10.1002/1873-3468.13862.
- Hartwell LH, Hopfield JJ, Leibler S, Murray AW. From molecular to modular cell biology. Nature. 1999;402(6761 SUPPL. 1):C47–C52. doi:10.1038/35011540.
- Grunberg TW, Del Vecchio D. Modular Analysis and Design of Biological Circuits. Curr Opin Biotechnol. 2020;63:41–47. doi:10.1016/j.copbio.2019.11.015.
- Atay O, Skotheim JM. Modularity and predictability in cell signaling and decision making. Mol Biol Cell. 2014;25(22):3445–3450. doi:10.1091/mbc.E14-02-0718.
- Gyorgy A, Del Vecchio D. Modular Composition of Gene Transcription Networks. PLoS Comput Biol. 2014;10(3):1003486. doi:10.1371/journal.pcbi.1003486.
- Garcia S, Trinh CT. Modular design: Implementing proven engineering principles in biotechnology. Biotechnol Adv. 2019;37(7):107403. doi:10.1016/j.biotechadv.2019.06.002.

- 39. Atay O, Doncic A, Skotheim JM. Switch-like Transitions Insulate Network Motifs to Modularize Biological Networks. Cell Syst. 2016;3(2):121–132. doi:10.1016/j.cels.2016.06.010.
- Araujo AR, Gelens L, Sheriff RSM, Santos SDM. Positive Feedback Keeps Duration of Mitosis Temporally Insulated from Upstream Cell-Cycle Events. Mol Cell. 2016;64(2):362–375. doi:10.1016/j.molcel.2016.09.018.
- 41. Dalchau N, Szép G, Hernansaiz-Ballesteros R, Barnes CP, Cardelli L, Phillips A, et al. Computing with biological switches and clocks. Nat Comput. 2018;17(4):761–779. doi:10.1007/s11047-018-9686-x.
- 42. Aguda BD, Goryachev AB. From Pathways Databases to Network Models of Switching Behavior. PLoS Comput Biol. 2007;3(9):e152. doi:10.1371/journal.pcbi.0030152.
- 43. Gérard C, Goldbeter A. A skeleton model for the network of cyclin-dependent kinases driving the mammalian cell cycle. Interface Focus. 2011;1(1):24–35. doi:10.1098/rsfs.2010.0008.
- 44. Gérard C, Gonze D, Goldbeter A. Effect of positive feedback loops on the robustness of oscillations in the network of cyclin-dependent kinases driving the mammalian cell cycle. FEBS J. 2012;279(18):3411–3431. doi:10.1111/j.1742-4658.2012.08585.x.
- Hernansaiz-Ballesteros RD, Cardelli L, Csikász-Nagy A. Single molecules can operate as primitive biological sensors, switches and oscillators. BMC Syst Biol. 2018;12(1):70. doi:10.1186/s12918-018-0596-4.
- Novák B, Tyson JJ. Design principles of biochemical oscillators. Nat Rev Mol Cell Biol. 2008;9(12):981–991. doi:10.1038/nrm2530.
- 47. Johnson KA, Goody RS. The original Michaelis constant: Translation of the 1913 Michaelis-Menten Paper. Biochemistry. 2011;50(39):8264–8269. doi:10.1021/bi201284u.
- Goldbeter A, Koshland DE. An amplified sensitivity arising from covalent modification in biological systems. Proc Natl Acad Sci U S A. 1981;78(11 II):6840–6844. doi:10.1073/pnas.78.11.6840.
- Hill, V A. The possible effects of the aggregation of the molecules of haemoglobin on its dissociation curves. J Physiol. 1910;40:4–7.
- Gonze D, Abou-Jaoudé W. The Goodwin Model: Behind the Hill Function. PLoS One. 2013;8(8):69573. doi:10.1371/journal.pone.0069573.
- 51. Ferrell Jr JE, Ha SH. Ultrasensitivity part I: Michaelian responses and zero-order ultrasensitivity. Trends Biochem Sci. 2014;39(10):496–503. doi:10.1016/j.tibs.2014.08.003.
- Korsbo N, Jönsson H. It's about time: Analysing simplifying assumptions for modelling multi-step pathways in systems biology. PLoS Comput Biol. 2020;16(6):e1007982. doi:10.1371/journal.pcbi.1007982.
- Hinch R, Schnell S. Mechanism equivalence in enzyme-substrate reactions: Distributed differential delay in enzyme kinetics. J Math Chem. 2004;35(3):253–264. doi:10.1023/B:JOMC.0000033258.42803.60.
- Ferrell Jr JE, Ha SH. Ultrasensitivity part III: Cascades, bistable switches, and oscillators. Trends Biochem Sci. 2014;39(12):612–618. doi:10.1016/j.tibs.2014.10.002.
- Csikász-Nagy A, Battogtokh D, Chen KC, Novák B, Tyson JJ. Analysis of a generic model of eukaryotic cell-cycle regulation. Biophys J. 2006;90(12):4361–4379. doi:10.1529/biophysj.106.081240.
- Gérard C, Goldbeter A. Temporal self-organization of the cyclin/Cdk network driving the mammalian cell cycle. Proc Natl Acad Sci U S A. 2009;106(51):21643–21648. doi:10.1073/pnas.0903827106.

- 57. Terhune SS, Jung Y, Cataldo KM, Dash RK. Network mechanisms and dysfunction within an integrated computational model of progression through mitosis in the human cell cycle. PLoS Comput Biol. 2020;16(4):e1007733. doi:10.1371/journal.pcbi.1007733.
- 58. FitzHugh R. Impulses and Physiological States in Theoretical Models of Nerve Membrane. Biophys J. 1961;1(6):445–466. doi:10.1016/S0006-3495(61)86902-6.
- 59. Tsai TYC, Theriot JA, Ferrell Jr JE. Changes in Oscillatory Dynamics in the Cell Cycle of Early Xenopus laevis Embryos. PLoS Biol. 2014;12(2):e1001788. doi:10.1371/journal.pbio.1001788.
- 60. Rombouts J, Vandervelde A, Gelens L. Delay models for the early embryonic cell cycle oscillator. PLoS One. 2018;13(3):e0194769. doi:10.1371/journal.pone.0194769.
- Vinod PK, Novak B. Model scenarios for switch-like mitotic transitions. FEBS Lett. 2015;589(6):667–671. doi:10.1016/j.febslet.2015.02.007.
- 62. Williams BC, Filter JJ, Blake-Hodek KA, Wadzinski BE, Fuda NJ, Shalloway D, et al. Greatwall-phosphorylated Endosulfine is both an inhibitor and a substrate of PP2A-B55 heterotrimers. Elife. 2014;2014(3). doi:10.7554/eLife.01695.
- Arooz T, Yam CH, Siu WY, Lau A, Li KKW, Poon RYC. On the concentrations of cyclins and cyclin-dependent kinases in extracts of cultured human cells. Biochemistry. 2000;39(31):9494–9501. doi:10.1021/bi0009643.
- 64. Tadros W, Lipshitz HD. The maternal-to-zygotic transition: A play in two acts. Development. 2009;136(18):3033-3042. doi:10.1242/dev.033183.
- Farrell JA, O'Farrell PH. From egg to gastrula: How the cell cycle is remodeled during the drosophila mid-blastula transition. Annu Rev Genet. 2014;48:269–294. doi:10.1146/annurev-genet-111212-133531.
- Olivier N, Luengo-Oroz MA, Duloquin L, Faure E, Savy T, Veilleux I, et al. Cell lineage reconstruction of early zebrafish embryos using label-free nonlinear microscopy. Science (80-). 2010;329(5994):967–971. doi:10.1126/science.1189428.
- Anderson GA, Gelens L, Baker JC, Ferrell JE. Desynchronizing Embryonic Cell Division Waves Reveals the Robustness of Xenopus laevis Development. Cell Rep. 2017;21(1):37–46. doi:10.1016/j.celrep.2017.09.017.
- Lim S, Kaldis P. Cdks, cyclins and CKIs: Roles beyond cell cycle regulation. Dev. 2013;140(15):3079– 3093. doi:10.1242/dev.091744.
- 69. Rombouts J, Gelens L. Dynamic bistable switches enhance robustness and accuracy of cell cycle transitions. PLOS Comput Biol. 2021;17(1):e1008231. doi:10.1371/journal.pcbi.1008231.
- Kwon JS, Everetts NJ, Wang X, Wang W, Della Croce K, Xing J, et al. Controlling Depth of Cellular Quiescence by an Rb-E2F Network Switch. Cell Rep. 2017;20(13):3223–3235. doi:10.1016/j.celrep.2017.09.007.
- Moser J, Miller I, Carter D, Spencer SL. Control of the restriction point by rb and p21. Proc Natl Acad Sci U S A. 2018;115(35):E8219–E8227. doi:10.1073/pnas.1722446115.
- Jirawatnotai S, Hu Y, Livingston DM, Sicinski P. Proteomic identification of a direct role for cyclin D1 in DNA damage repair. Cancer Res. 2012;72(17):4289–4293. doi:10.1158/0008-5472.CAN-11-3549.
- Kousholt A, Menzel T, Sørensen C. Pathways for Genome Integrity in G2 Phase of the Cell Cycle. Biomolecules. 2012;2(4):579–607. doi:10.3390/biom2040579.

- Matsuo T, Yamaguchi S, Mitsui S, Emi A, Shimoda F, Okamura H. Control mechanism of the circadian clock for timing of cell division in vivo. Science (80-). 2003;302(5643):255–259. doi:10.1126/science.1086271.
- Heltberg ML, Jensen MH. Locked body clocks. Nat Phys. 2019;15(10):989–990. doi:10.1038/s41567-019-0617-2.
- 76. Gérard C, Goldbeter A. Entrainment of the Mammalian Cell Cycle by the Circadian Clock: Modeling Two Coupled Cellular Rhythms. PLoS Comput Biol. 2012;8(5):e1002516. doi:10.1371/journal.pcbi.1002516.
- 77. Elowitz MB, Leibier S. A synthetic oscillatory network of transcriptional regulators. Nature. 2000;403(6767):335–338. doi:10.1038/35002125.
- Novikoff AB. The concept of integrative levels and biology. Science (80-). 1945;101(2618):209–215. doi:10.1126/science.101.2618.209.
- Crook N, Alper HS. Model-based design of synthetic, biological systems. Chem Eng Sci. 2013;103:2– 11. doi:10.1016/j.ces.2012.12.022.
- Eshtewy NA, Scholz L. Model reduction for kinetic models of biological systems. Symmetry (Basel). 2020;12(5):863. doi:10.3390/SYM12050863.
- Gorban AN. Model reduction in chemical dynamics: slow invariant manifolds, singular perturbations, thermodynamic estimates, and analysis of reaction graph. Curr Opin Chem Eng. 2018;21:48–59. doi:10.1016/j.coche.2018.02.009.
- 82. Rao S, van der Schaft A, van Eunen K, Bakker BM, Jayawardhana B. A model reduction method for biochemical reaction networks. BMC Syst Biol. 2014;8(1):52. doi:10.1186/1752-0509-8-52.
- Pandey A, Murray R. Model Reduction Tools For Phenomenological Modeling of Input-Controlled Biological Circuits. bioRxiv. 2020; p. 2020.02.15.950840. doi:10.1101/2020.02.15.950840.
- Albert R, Robeva R. Signaling Networks: Asynchronous Boolean Models. In: Algebr. Discret. Math. Methods Mod. Biol. Elsevier; 2015. p. 65–91.
- Cappell S, Chung M, Jaimovich A, Spencer S, Meyer T. Irreversible APCCdh1 Inactivation Underlies the Point of No Return for Cell-Cycle Entry. Cell. 2016;166(1):167–180. doi:10.1016/j.cell.2016.05.077.
- Rubin SM, Sage J, Skotheim JM. Integrating Old and New Paradigms of G1/S Control. Mol Cell. 2020;80(2):183–192. doi:10.1016/j.molcel.2020.08.020.
- 87. Barr AR, Heldt FS, Zhang T, Bakal C, Novák B. A Dynamical Framework for the All-or-None G1/S Transition. Cell Syst. 2016;2(1):27–37. doi:10.1016/j.cels.2016.01.001.
- Lafranchi L, Müllers E, Rutishauser D, Lindqvist A. FRET-Based Sorting of Live Cells Reveals Shifted Balance between PLK1 and CDK1 Activities During Checkpoint Recovery. Cells. 2020;9(9):2126. doi:10.3390/cells9092126.
- Chao HX, Poovey CE, Privette AA, Grant GD, Chao HY, Cook JG, et al. Orchestration of DNA Damage Checkpoint Dynamics across the Human Cell Cycle. Cell Syst. 2017;5(5):445–459.e5. doi:10.1016/j.cels.2017.09.015.
- Taylor WR, DePrimo SE, Agarwal A, Agarwal ML, Schönthal AH, Katula KS, et al. Mechanisms of G2 arrest in response to overexpression of p53. Mol Biol Cell. 1999;10(11):3607–3622. doi:10.1091/mbc.10.11.3607.

- 91. Gillis LD, Leidal AM, Hill R, Lee PWK. p21Cip1/WAF1 mediates cyclin B1 degradation in response to DNA damage. Cell Cycle. 2009;8(2):253–256. doi:10.4161/cc.8.2.7550.
- 92. Tian XJ, Zhang H, Xing J. Coupled reversible and irreversible bistable switches underlying TGFβ-induced epithelial to mesenchymal transition. Biophys J. 2013;105(4):1079–1089. doi:10.1016/j.bpj.2013.07.011.
- 93. Gunne-Braden A, Sullivan A, Gharibi B, Sheriff RSM, Maity A, Wang YF, et al. GATA3 Mediates a Fast, Irreversible Commitment to BMP4-Driven Differentiation in Human Embryonic Stem Cells. Cell Stem Cell. 2020;26(5):693–706.e9. doi:10.1016/j.stem.2020.03.005.
- 94. Wang L, Walker BL, Iannaccone S, Bhatt D, Kennedy PJ, Tse WT. Bistable switches control memory and plasticity in cellular differentiation. Proc Natl Acad Sci U S A. 2009;106(16):6638–6643. doi:10.1073/pnas.0806137106.
- 95. Graham TGW, Tabei SMA, Dinner AR, Rebay I. Modeling bistable cell-fate choices in the Drosophila eye: Qualitative and quantitative perspectives. Development. 2010;137(14):2265–2278. doi:10.1242/dev.044826.
- 96. Balaskas N, Ribeiro A, Panovska J, Dessaud E, Sasai N, Page KM, et al. Gene regulatory logic for reading the sonic hedgehog signaling gradient in the vertebrate neural tube. Cell. 2012;148(1-2):273–284. doi:10.1016/j.cell.2011.10.047.
- Kuznetsov YA. Numerical Analysis of Bifurcations. In: Elem. Appl. Bifurc. Theory. 3rd ed.; 2004. p. 505–585.
- 98. Winston JT, Pledger WJ. Growth factor regulation of cyclin D1 mRNA expression through protein synthesis-dependent and -independent mechanisms. Mol Biol Cell. 1993;4(11):1133–1144. doi:10.1091/mbc.4.11.1133.
- 99. Perry JE, Grossmann ME, Tindall DJ. Epidermal growth factor induces cyclin D1 in a human prostate cancer cell line. Prostate. 1998;35(2):117–124. doi:10.1002/(SICI)1097-0045(19980501)35:2;117::AID-PROS5;3.0.CO;2-G.
- 100. Ansmann G. Efficiently and easily integrating differential equations with JiTCODE, JiTCDDE, and JiTCSDE. Chaos. 2018;28(4):043116. doi:10.1063/1.5019320.





















