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Biologists have long observed periodic-like oxygen consumption oscillations in yeast populations under certain conditions, and several unsatisfactory explanations for this phenomenon have been proposed. These ‘autonomous oscillations’ have often appeared with periods that are nearly integer divisors of the calculated doubling time of the culture. We hypothesize that these oscillations could be caused by a form of cell cycle synchronization that we call clustering. We develop some novel ordinary differential equation models of the cell cycle. For these models, and for random and stochastic perturbations, we give both rigorous proofs and simulations showing that both positive and negative growth rate feedback within the cell cycle are possible agents that can cause clustering of populations within the cell cycle. It occurs for a variety of models and for a broad selection of parameter values. These results suggest that the clustering phenomenon is robust and is likely to be observed in nature. Since there are necessarily an integer number of clusters, clustering would lead to periodic-like behaviour with periods that are nearly integer divisors of the period of the cell cycle. Related experiments have shown conclusively that cell cycle clustering occurs in some oscillating yeast cultures.

Keywords: autonomous oscillations; cell cycle; budding yeast

AMS Subject Classification: 37N25; 34C25; 34F05; 92D25

1. The yeast cell cycle and experimental observations

Klevecz [36], McKnight [46], and others have observed oscillations in the dissolved O2 levels of various strains of budding yeast under certain conditions. For instance, in Figure 1, we show data from such an experiment for the yeast strain Cen.PK 113 [38,42,43]. It is seen in this experiment that dissolved O2 levels oscillate between about 5% and 55%, with a period of a little less than 4 h. Since the doubling time (as calculated from the dilution rate) for this culture was 7.8 h, it is natural to suppose that the doubling time and O2 oscillations are causally related.

Autonomous oscillations have been the subject of speculation and analysis [7,11,20–22,37,38,43,46,50]. Cultures exhibiting O2 oscillations, although planktonic, are dense; the average distance between cells has been reported to be of the order of 1 cell diameter [36]. Observations such
Figure 1. (Available in colour online). Percent dissolved O\textsubscript{2} (green), bud index (blue) and cell density (red). The bud index (percentage of cells with buds) and cell density are both synchronized with the oscillation in the level of dissolved O\textsubscript{2}. Moreover, time periods of sharp increases in bud index percentage show that a cluster of cell (consisting of almost 50\% of the cells) is passing into S phase together. The doubling time of this culture is 7.8 h. Two oscillations per 7.8 h period also is consistent with two clusters of cells.

as these suggest the involvement of quorum sensing. Indeed, Klevecz and others have suggested that signalling is responsible for the O\textsubscript{2} oscillations based on the observation that acetyaldehyde and H\textsubscript{2}S pulses reset the phase of the oscillations [31,36]. Alternative explanations involving signalling have been proposed. For instance, Hjortso and Nielsen [22] have suggested an elegant mechanism in which the periodic consumption of metabolites can stabilize population synchrony.

Because the mitotic cell cycle of the budding yeast plays an important role as a coordinate in the ensuing analysis and theory, we briefly describe some of the relevant facts and nomenclature. Detailed reviews of the physiology and genetics of the budding yeast cell cycle abound [4,18,19,24,29], to list only a few. The mitotic cell cycle entails the process of DNA duplication, daughter cell growth and the subsequent cell division. The division processes in the budding yeast is asymmetric in the sense that it results in a distinguishable mother and daughter cell. The daughter cell at division is typically smaller in volume than the mother cell [48]. Only the mother cell carries an observable and permanent chitanous bud scar at the site of daughter cell growth. The number of bud scars determines the replicative age of a yeast cell which is to be distinguished from its chronological age. The bud scar equivalence classes are henceforth called generations. The emergence of a single bud, as opposed to multiple buds, per division cycle is tightly regulated.

As depicted in Figure 2, the mitotic cell cycle is typically partitioned into four phases, canonically enumerated in the sequence: G\textsubscript{1}, S, G\textsubscript{2} and M. The DNA is replicated during the S phase, and the temporal duration of the S phase is very regular [2,8,40,41]. The G\textsubscript{1} phase duration is observed to be the most variable. During the G\textsubscript{1} phase, environmental cues are integrated and synthetic precursors are obtained. The volumetric growth of the mother–daughter pair is observed to be monotone along the cell cycle in single cells [27,45,49], with growth and division jointly
Figure 2. Phases of the yeast cell cycle. DNA is replicated in the synthetic or S phase. DNA synthesis is preceded by the G1 phase, in which environmental cues are integrated and precursors are acquired. Our model explores the implications of the existence of a responsive R phase, situated within the G1 phase, during which cells can respond to signals by increasing or decreasing their growth rate. The M phase ends in cell division.

regulated by an as yet incompletely understood genetic network [30]. A complete causal description of the transition from G1 to S remains open [5,29]. This transition closely coincides with the phenomena of bud emergence.

In this article, we study the implications of the hypothesis that cells in the S phase can effect the growth rate of cells in a responsive phase (denoted as R) situated in the G1 phase. The growth rate modulation can be either positive or negative. We show rigorously and numerically in simple models that various forms of signalling among cells, with either positive or negative feedback, robustly lead the population density to become multimodal or clustered within the cell cycle.

We propose here that dissolved O2 oscillations in some strains could be due to a phenomenon that we will call clustering. In general, we will define clustering loosely as significant groups of cells going through cell cycle milestones at approximately the same time, i.e. there is a form of temporal synchronization. The clustering could cause fluctuations in the O2 dilution levels by clusters passing in and out of high oxygen metabolism phases of the cell cycle, i.e. G1 phase.

We will prove mathematically in idealized models that clustering occurs, and that the number of clusters formed depends on the widths |S| and |R| of the S phase and the hypothesized region R. Of key importance is the necessity that, if clustering occurs, then there are an integer number of clusters. This can provide the basis for a periodic-looking behaviour with a period, which is approximately an integer divisor of the cell-cycle length. Models for which proofs are constructed include noisy or randomly perturbed models. We use simulations to demonstrate that the same type of behaviour occurs in more detailed models. We note that an analogous form of clustering has been observed in all-to-all coupled networks of integrate-and-fire oscillators, and recently a rigorous proof of clustering and the stability of such solutions in these models were obtained [34].

In prior work [38,42,43], we have shown through an analysis of the bud index and cell density time-series (Figure 1) that clustering occurs, and have proposed how the periodic population phenomena could be exploited to enhance bioprocess.

In this paper, we focus on models that assume that the cell cycles of different generations are identical, or nearly so. Although this assumption is not justified in general yeast cultures, there is some evidence for it in the cultures under study. In conditions where the oscillations have been observed, the doubling time of the culture is extended significantly, by a factor of about four. It is known, and confirmed in [38,42,43], that when the cell cycle is extended, most of the extension occurs in the G1 phase. Age dependent differences in cell cycle duration have been observed and quantified [48], and are attributed to differences in G1 duration, although this will require further quantization at the single cell level [45,49]. Regardless, during autonomous oscillation, extending
the doubling time effectively elongates the cycles of all the generations to such an extent that the relative differences become insignificant. The age distribution provides further evidence that the assumption of equal cell cycle length for different generations is approximately true in the cited experiments.

In addition to models with the assumptions of identical or nearly identical cell cycles, we also briefly treat models with relative differences across generations and cases where even larger differences could still lead to clustering. In [42,43], we have explored a model with stratified generations under various assumptions and using biologically relevant values for all the parameters that can be determined. In all these models, clustering emerges as a common and robust phenomenon.

2. Models of cell cycle dynamics and clustering

2.1. A general model for the cell cycle of an individual cell

We will begin by defining a normalized logarithmic scale to represent the cell cycle. Customarily, the cell cycle is delineated by volume milestones. For a given cell, indexed by $i$, let $v_i(t)$ denote its volume. It is usually assumed that the volume growth of a cell in a culture is proportional to its volume, i.e.

$$\frac{dv_i}{dt} = c_i(t)v_i,$$

where the relative growth rate $c_i(t)$ may depend on many factors, such as available resources, chemical composition of the culture substrate, etc [10,26,28,47]. We will allow that the growth rate may also be influenced by the state of the cell itself, thus the cell may react to environmental factors in differing ways at different stages of its cycle. The environmental factors in turn may be influenced by the cells in the culture and their history. Finally, the individual cells may have individual differences. Without any loss of generality, we may include all of these factors in the rate $c_i(t)$.

Let $V_{b,i}$ denote the volume of the cell at the beginning of its cell cycle and $V_{d,i}$ its volume at division. In order to have $x_i \in [0, 1)$, we make the change of variables:

$$x_i = \ln\left(\frac{v_i}{V_{b,i}}\right)\frac{1}{\ln(\frac{V_{d,i}}{V_{b,i}})}.$$

With this change, birth has the coordinate $x_i = 0$ and division occurs at $x_i = 1$. We have then that $x_i$ satisfies

$$\frac{dx_i}{dr} = d_i(t) \equiv \frac{c_i(t)}{\ln(\frac{V_{d,i}}{V_{b,i}})}.$$

Next, we may rescale time so that the average time span of a cell cycle in the culture is normalized to 1, i.e. scale by a factor $d(t)$ which is the average (over all cells) growth rate in a culture. In making this change of time, we obtain

$$\frac{dx_i}{dr} = \frac{d_i(t)}{d(t)}.$$

If the variation between individual cells is not too great, then the right-hand side of this equation is approximately 1. Next, we distinguish between those influences on the growth rate that are common to all the cells in the culture and those due to individual differences. We allow the common features to depend on the state, $x_i$ of the cell itself, as well as the conglomerate history
of the whole culture, which we denote by \( \bar{x} \). Denote the common part of the growth rate as \( a \) and the individual part as \( g_i \). We can thus write the equation as follows:

\[
\frac{dx_i}{dt} = a(t, x_i, \bar{x}) + g_i(t). \tag{1}
\]

The part of the growth rate due solely to individual differences in cells is contained in the term \( g_i \). In many applications, this term will be relatively small and ‘random’ in the sense that differences are due to details of the cell process that are far too minute and complex to model. It contains both variations in the growth rate and differences in \( V_{b,i} \) and \( V_{d,i} \). Thus, we may view Equation (3) as a random differential equation (RDE) [3]. A reasonable approximation of this equation in some circumstances is to replace \( g_i \) by a stochastic term:

\[
\frac{dx_i}{dt} = a(t, x_i, \bar{x}) \, dt + \sqrt{\sigma} \, dW_i, \tag{2}
\]

where \( dW_i \) represents an independent noise term. This stochastic differential equation (SDE) must be interpreted in the usual way as an Ito integral equation. For other possible stochastic approximations see [1]. It is also reasonable under certain circumstances to consider Equations (1) and (2) as perturbations of an ordinary differential equation (ODE)

\[
\frac{dx_i}{dt} = a(t, x_i, \bar{x}). \tag{3}
\]

This model allows for variation of the growth on state within the cell cycle and the overall state of the system, but ignores differences between individual cells. We will consider versions of this model in the rest of this section.

### 2.2. Model of the culture

First of all, we note that in a culture, cells are constantly dividing, dying and perhaps being harvested. Thus, the equations above must be applied to a changing set of cells, indexed by a changing finite set of positive integers \( S(t) \subset \mathbb{N} \). If the \( i \)th cell dies or is harvested, then \( i \) is dropped from \( S \). When a cell divides, one of the new cells could continue being denoted by \( i \), with \( x_i \) resetting to 0, and the other (daughter cell) would be identified by a new index. This would be appropriate in budding yeast where the mother and daughter cells are distinguishable. The conglomerate history \( \bar{x} \) would contain each \( x_i \) over its respective lifespan.

Now suppose that we are considering the unperturbed equation (3). Since in this model there are no terms that differentiate the progression of different cells, when a cell divides its two descendants will both start at \( x_i = 0 \) and remain completely synchronized for the rest of their lifespans. Thus, tracking the evolution of both cells is redundant in this model and we are inevitably led to consider a fixed set of cells. If the culture is in a steady or periodic state, then there is also a probabilistic interpretation of this simplification; given a living cell at time \( t \), the expected number of cells descended from that cell at any later time \( t + n \) is exactly 1. Thus, the original model with a changing index set \( S \) can be replaced by a fixed system, each variable \( x_i(t) \) representing the state of its expected descendant at time \( t \). A PDE justification of considering a fixed index set appears in the appendix.

With a fixed index set, this model is easily amenable to numerical investigation with currently available computer speeds. For instance, one can easily investigate the behaviour of a conglomeration of 10,000 or more cells over several cell cycles on a desktop workstation. Further, under some additional assumptions on the dependence of \( a \) and \( g_i \), we can investigate properties of the solutions of Equations (1) and (3) rigorously as we show below.
2.3. Modelling of the growth term $a$

The standard biological assumption on $a$ is that it does not depend on $x_i$, i.e. it is independent of the cell’s current state within the cycle. Rather it is assumed to depend mostly on the nutrients available and other environmental factors. With these assumptions $a = a(t, \bar{x})$.

As already described earlier, we hypothesize a more sophisticated form for $a$, in which the location of cells in a cell cycle may influence the growth rate of themselves and other cells, i.e.

$$a = a(t, x_i, \bar{x}).$$

For example, in budding yeast, it has been hypothesized that cells in the S phase may influence the growth of cells in the pre-budded G1 phase through some (unspecified) signalling mechanism [6,16,23,35].

In our models, the population of cells in the S phase will be assumed to effect the growth of cells in a preceding portion of the cell cycle, which we denote by $R = [r, s]$, i.e. a portion of the G1 phase. As explained in the previous section and in the appendix, we will consider a fixed finite population of $N$ cells, each of them progressing via the following form of Equation (3)

$$\frac{dx_i}{dt} = \begin{cases} 1 & \text{if } x_i \not\in R \\ 1 + F(\#\{\text{cells in } S\}) & \text{if } x_i \in R. \end{cases}$$

(4)

When a cell in the model reaches division at $x = 1$, it returns to the beginning of the cycle, $x = 0$. The phase space for each individual cell is thus the unit circle and the phase space of the entire culture is the $N$ torus. In this section, we will consider two idealized (and discontinuous) forms of positive and negative $F$ that we call the advancing and blocking. Our motivation for considering such models is that they are analytically tractable. Also, biologically it is known that the introduction of certain compounds has been shown to arrest or rapidly accelerate cell processes in budding yeast and other micro-organisms [7,15,25,36]. Note that periodic blocking at division was considered by [39].

By the advancing model, we will mean that if the fraction of cells in $S$ exceeds some threshold $\tau$, then all cells in $R$ are instantaneously advanced to the beginning of $S$, from which they resume the normal growth. This corresponds to a limit $F \rightarrow \infty$ in Equation (4) when $\#\{\text{cells in } S\} \geq \tau$ and $F = 0$ when $\#\{\text{cells in } S\} < \tau$.

By the blocking model, we mean that if the fraction of cells in $S$ equals or exceeds some threshold $\tau$, then cells at $s$ are blocked from proceeding into the S phase, until the fraction of cells in $S$ drops below the threshold. This corresponds to choice $F = -1$ when $x_i = s$ and $\#\{\text{cells in } S\} \geq \tau$ and $F = 0$ when $\#\{\text{cells in } S\} < \tau$. If more than $\tau$ cells have accumulated at $s$, then all of those cells will enter $S$ together when the fraction of cells drops below $\tau$.

Note that the advancing and blocking models are defined in terms of their flow directly, thus the solution flow is unique. Since the phase space is compact, the flow is global.

2.4. Clustering

Under either the advancing or blocking models, it is clear that some cells may become synchronized. Consider the advancing model; whenever the fraction of cells in $S$ exceeds the threshold $\tau$, then all cells in $R$ are instantaneously synchronized at $s$. There being no mechanism in the model to subsequently differentiate them, they will remain synchronized from then on. Similarly, for the blocking model, during a period when the threshold is exceeded, all cells arriving at $s$ will be thereafter synchronized. We will call a group of synchronized cells a cluster. If the number of cells in a cluster is large enough to exceed the threshold of the model, we call it a critical cluster.
Given an initial (discrete) distribution \( \tilde{x}(0) \) or \( w(s) = \sum_{i=1}^{N} \delta_{q_i}(s) \) the evolution of such distribution under either model will be called a trajectory starting at \( w(s) \) and denoted by \( \phi(t, w(s)) \). It is clear that different trajectories can merge.

A trajectory is called an equilibrium if it is stationary in the coordinate frame moving with the speed 1. In other words, \( \phi(t, w(s)) = w(x - t) \mod 1 \) for all \( t \geq 0 \). A periodic orbit is a trajectory that is periodic in the moving frame. In such a case, there exists \( T \) with \( \phi(T + t, w(x)) = \phi(t, w((x))) \) for all \( t \in [0, T] \); if \( T \) is the smallest number with this property, it is called a period.

### 2.5. Advancing model

Denote by \( [x] \) the largest integer \( \leq x \) and by \( \lfloor x \rfloor \) the smallest integer \( \geq x \).

**Theorem 2.1** Consider the advancing model.

1. If the initial \( w(x) \) exceeds the threshold on all intervals of length \( |S| \), except possibly inside the interval \( R \), then the trajectory converges to a periodic orbit.
2. Any initial \( w(x) \) that is below threshold on all intervals of length \( |S| \), is an equilibrium point.
3. Every other initial condition converges to an equilibrium \( e \) with a finite number of critical clusters, separated by voids of length at least \( |R| + |S| \).

**Proof** Let

\[
q(t, x) = \int_{[x, x+|S|] \mod 1} \phi(t, w(z))dz. \tag{5}
\]

The function \( q(t, x) \) is a form of local density of the discrete distribution of cells. Observe that if \( q(0, x) < \tau \) for all \( x \), then this is a fixed point of the dynamics. This corresponds to the case (2) above.

The case (1) corresponds to the case when \( q(0, x) \geq \tau \) for all \( x \), except for \( x \in R \). At \( t = 0 \), all cells in \( R \) are advanced to the point \( s \). Since the \( q(0, x) \geq \tau \) for all \( x \) the cells arriving at \( r \), the start of the interval \( R \), are immediately transferred to the point \( s \). This continues indefinitely. This trajectory is a periodic orbit in the moving frame.

Now we do the analysis of the case (3). Let \( q(i, x), i = 0, 1, \ldots \) be the mass function after \( i \) passes through the cell cycle. We decompose the domain \( I = [0, 1] \) into intervals \( I_j^0 \cup J_j^1 \cup \cdots \cup J_j^{n-1} \cup I_j^n \) where \( q(i, x) \geq \tau \) for all \( x \in I_j^i \) and \( q(i, x) < \tau \) for all \( x \in J_j^i \). Set \( I_j^i := [d_{j, i}, c_{j, i}] \). We compare the intervals \( I_j^i \) from iteration to iteration. There are several possibilities that can happen to \( I_j^i \) after a passage through a cell cycle:

1. The interval \( I_j^i \) will shorten by \( |R| \) and \( I_j^{i+1} = [d_{j, i} + |R|, c_{j, i}] \).
2. The intervals \( I_j^i, \ldots, I_{j-k} \) may merge, if all intervals in between fall within distance \( |R| \), i.e. \( c_{j, i} - c_{j-k, i} < |R| \).
3. If an interval \( I_{j-1} \) follows the interval \( I_j^i \) by a distance closer, then \( |R| + |S| \); i.e. \( d_{j, i} - c_{j-1, i} \leq |R| + |S| \), then the interval \( I_{j-1} \), or a portion of that interval, will be promoted across \( R \). The resulting distance between the two new intervals will be less then \( |S| \). After each subsequent pass, the distance between these two intervals will shorten by \( |R| \) and in finite number of steps they will be within \( |R| \) of each other and they will merge. The number of steps this takes is uniformly bounded above by \( l := \frac{|S|}{|R|} + 1 \). This process may result in a split of an interval \( I_j^i \) into two intervals with the total mass conserved in the transaction. Note that even though temporarily the number of intervals \( I_j^{i+1} \) may increase over the number \( I_j^i \), after \( k \) subsequent passes through \( S \) the number of intervals \( I_j^{i+k} \) is less than or equal to the number of intervals \( I_j^i \).
(4) The interval $I_j^i$ may shorten, if the preceding interval $I_{j+1}^i$ has critical mass in $S$, and therefore promotes part of the mass ahead of $c_j^i$ across $R$. Since a mass ahead of $c_j^i$ is lost, $c_j^{i+1} < c_j^i$. The difference between this case and case (3) is that here $|R| + |S| \leq d_j^i - c_j^{i-1} \leq |R| + |S| + |S|$. Observe that the transferred cells will not form a new interval $I_{j+1}^i$, since they do not have a sufficient mass.

In summary, the number of intervals $I_j^i$ is greater than or equal to the number of intervals $I_j^{i+k}$ for every $i$, and the length of each intervals is a non-increasing function of $i$. Finally, it is easy to see that the only interval $I_j^i$ that will not change under the advance operator is a singleton $I_j^i = [c_j^i, c_j^i]$. Since the population is finite, after a finite number of passes through $S$, each $I_j$ is a singleton. If two such singletons follow each other closer, then $|R| + |S|$, by the step 3 above, and they will merge in finite number of steps. Eventually, all remaining singletons $I_j$ are followed by an empty void of length $|R| + |S|$.

**Corollary 2.2**  
In the advancing model no more than $\lceil (|R| + |S|)^{-1} \rceil$ critical clusters can persist.

**Proof**  
By the previous theorem, each critical cluster has to be followed by a gap of length at least $|R| + |S|$.

### 2.6. Blocking model

**Theorem 2.3**  
In the blocking model no more than $\lceil |S|^{-1} \rceil$ critical clusters can persist.

**Proof**  
When two consecutive critical clusters pass through $S$, the second cluster must wait at $s$ until the first cluster has passed $t$. Thus, the distance between them must then be at least $|S|$ and will remain at least $|S|$ until the first cluster hits $r$. Then, the first cluster may have to wait to enter $S$, possibly decreasing the distance between the two clusters in question.

Assume that the number $n$ of persistent clusters is constant along some trajectory. Then either all these clusters are separated by a distance at least $|S|$, or there are $n-1$ clusters separated by a distance $|S|$, and there is an additional cluster inside $S$ whose distance to a waiting cluster at $s$ is smaller than $|S|$.

This minimal spacing implies that

$$(n-1)|S| \leq 1.$$ 

The result then follows.

We have shown that a cluster cannot persistently follow a critical cluster closer than $|S|$ (except while the critical cluster is being blocked at $s$). Also, it is clear that as many as $\lceil |S|^{-1} \rceil$ critical clusters may persist if the overall number $N$ of cell satisfies:

$$N > \tau \lceil |S|^{-1} \rceil.$$ 

Recall the definition of $q(t, x)$ in Equation (5).

**Theorem 2.4**  
In the blocking model, if the cells are initially distributed so that the density $q(x, 0)$ everywhere exceeds twice the threshold, then exactly $\lceil |S|^{-1} \rceil$ clusters will develop and persist.
Proof  Cells at $s$ are initially stopped from entering $S$ until the fraction of cells in $S$ drops below the threshold. Since $d(|S| - t) = \tau$, this will occur at
\[ t = |S| - \frac{\tau}{d}. \]
At this time, the cells that have clustered at $s$ number $d|S|$ or $d(|S| - \tau)$. Since $d|S|$ is assumed to be at least twice $\tau$, the threshold is again exceeded as the cluster enters $S$. The threshold will continue to be exceeded until this cluster leaves $S$. At this time, a new cluster will have formed at $s$ with volume $d|S|$ which is by assumption above $\tau$. The second cluster is spaced exactly at a distance $|S|$ behind the first cluster. At third cluster, etc. will form in the same way until the first cluster returns to $s$. At that time, there is either a cluster in the interior of $S$ or at $t$. In the former case, the first cluster will wait less than time $|S|$ to enter $S$, ahead of when the second cluster arrives. In the latter case it enters $S$ immediately. This scenario will repeat itself indefinitely and the number of critical clusters thus produced is an easy calculation.

3. Small perturbations of the advancing and blocking models

3.1. Small perturbations

Next, we consider small perturbations of the advancing and blocking models by which we mean that the progression of each individual cell is independently perturbed by a small term which is independent of the other cells.

\[
\frac{dx_i}{dt} = \begin{cases} 
1 + \epsilon \xi_i(x_i, t) & \text{if } x_i \notin R \\
1 + F(\#\text{cells in } S) + \epsilon \xi_i(x_i, t) & \text{if } x_i \in R,
\end{cases}
\]

where $|\xi_i(x_i, t)| \leq 1$ and $\epsilon$ is small. We will take as our definition of small that $\epsilon$ is smaller than $\min(|S|, |R|)/6$. The effect of the perturbation is to cause initially synchronized cells to drift from each other, but by no more than $2\epsilon$ or $\min(|S|, |R|)/3$ within a single cell cycle. Note that under these assumptions, Equation (6) can be considered as a RDE with bounded noise. For general results about RDE with bounded noise see [24].

Note that this model can be viewed as a relaxation of the assumption that cells are indistinguishable. It also can be considered as incorporating random variations and noise.

With these perturbations, clusters will not remain precisely synchronized as in the unperturbed models, but rather will spread out as the cells proceed through the cell cycle. We now expand our definition of cluster to include any group of at least $\tau$ cells that are within $|S|/3$ of each other in the cell cycle.

Below, we will assume that the small perturbations act like noise in the ways we define below. We say that perturbations satisfy the \textit{maximum principle} if
\[
\max_x q(t_1, x) > \max_x q(t_2, x)
\]
and
\[
\min_x q(t_1, x) < \min_x q(t_2, x)
\]
for any $t_1 < t_2$ that are not separated by an activation or deactivation of blocking or advancing. We will say that the perturbations are \textit{diffusive} if:
- the perturbations satisfy the maximum principle.
- synchronized cells will immediately de-synchronize.
3.2. Perturbed blocking model

Theorem 3.1  In the blocking model with small diffusive perturbations such that $\epsilon < |S|/6$, and an initial distribution of cells that satisfies $q(0, x) > 2\tau + 2\epsilon$ for all $x$, the system will form either $|S|^{-1}$, $\lceil |S|^{-1} \rceil$ or $\lceil |S|^{-1} \rceil + 1$ critical clusters within one cell-cycle period and these clusters will persist indefinitely.

Proof  Denote $d = \min q(0, x)$. Cells at $s$ are initially blocked from entering $S$ until the fraction of cells in $S$ drops below the threshold. This will occur at time $t_0$ no smaller than

$$t_0 = \frac{|S| - \tau}{d} \cdot \frac{1}{1 + \epsilon}.$$  

At this time, the cells that have clustered at $s$ number at least $t_0d$ or $(d|S| - \tau)/(1 + \epsilon)$. This group of cells we will call cluster 1. By the assumptions, the threshold is again exceeded as the cluster enters and crosses $S$. While crossing $S$ the cluster will de-synchronize, but all the cells will remain in $S$ for at least $t_1 = |S|/(1 + \epsilon)$ and no longer than $|S|/(1 - \epsilon)$. Thus, while the first cluster crosses $S$ at least $t_1d$ or $d|S|/(1 + \epsilon)$ cell accumulate at $s$. This group we call the second cluster and from the assumption $q(0, x) > 2\tau + 2\epsilon$ it is critical. Its crossing time $t_2$ will again be bounded below by $|S|/(1 + \epsilon)$ and above by $|S|/(1 - \epsilon)$, and so this process of cluster formation will continue for as long as a density of cells at least $d$ is arriving at $s$.

By a straight-forward calculation, the gap between any two adjacent clusters formed in this process is bounded below by

$$|S| \frac{1 - \epsilon}{1 + \epsilon},$$

and obviously the gap is bounded above by $|S|$. If the first of two adjacent clusters travel at the minimal speed $1 - \epsilon$ while outside of $S$, all the cells in the first cluster must arrive again at $s$ no later than

$$t^* = \frac{1}{1 - \epsilon} - \frac{|S|}{1 + \epsilon},$$

after cluster 2 leaves $S$. During the same time period, the second of the adjacent clusters can travel at a rate at most $1 + \epsilon$ and thus can travel a distance of no more than

$$d^* = t^*(1 + \epsilon) = \frac{1 + \epsilon}{1 - \epsilon} - |S| < 1.$$  

Therefore, the entirety of the first cluster must reach $s$ before any of the second cluster does so. The gap at that time is at least

$$1 - d^* = |S| - \frac{2\epsilon}{1 + \epsilon} > \frac{2|S|}{3}.$$  

Next consider the relative motion between the cells that began in $S$ and the cells in cluster 1. Note that the initial gap will be at least $|S|/2$. Since $\epsilon < |S|/4$, the last of the cells will reach $s$ before the first cluster returns.
When cluster 2 leaves $S$, it is easily shown that the gap between clusters 1 and 2 is no wider than
\[ |S| \frac{1 + \epsilon}{1 - \epsilon} < |S|(1 + 2\epsilon). \]

At the same time, the gap between clusters 1 and 3 is no more than the gap between clusters 1 and 2 plus $|S|$. That is,
\[ 2|S|(1 + \epsilon). \]

When cluster 3 reaches $t$, the gap between clusters 1 and 3 is no more than
\[ 2|S|(1 + \epsilon) + |S| \frac{1 + \epsilon}{1 - \epsilon} - |S| < 2(1 + 2\epsilon)|S|. \]

By induction, one can show that the first cluster can be no more than
\[ (j - 1)(1 + 2\epsilon)|S| \]
ahead of the $j$th cluster when the $j$ cluster is at $t$. From this, we can show that at least $\lfloor |S|^{-1} \rfloor$ clusters will form.

On the other hand, when cluster 2 leaves $|S|$ the gap between the clusters 1 and 2 must be at least
\[ \frac{2}{1 + \epsilon} |S|, \]
and cluster 1 must be at least
\[ \frac{j - 1 - \epsilon}{1 + \epsilon} |S| \]
ahead of the $j$ cluster. It is clear then that $j$ must be less than $\lceil |S|^{-1} \rceil + 1$.

Now to show that the clusters persist. Clearly, while cluster 1 is passing through the cycle the first time, there is always a critical cluster in $S$, at least until the cells that began in $S$ again reach $S$. Note that the cells that were inside $S$ when the first cluster was released from $s$ are critical in number. We will call this group of cells the tail. When the tail begins to arrive back at $S$, its leading edge can be no more than
\[ \frac{|S|(1 - |S|)}{1 - \epsilon} \]
ahead of cluster 1. At that moment, there will either be (1) a critical in the interior of $S$, or (2) there are critical clusters at both ends of $S$. Let us consider case (2) first. In this case, the cluster at the beginning of $S$ will be critical and as it enters $S$ the cells from tail will be blocked at $s$. Since the original width of the tail is less than $|S|/2$, its width when it reaches $S$ will be less than $2|S|/3$ and it will all reach $S$ before the cluster ahead of it leaves $S$. Now two things can happen: both of which continue the process, either all or part of cluster 1 reaches $S$ while the tail is blocked there, or it will be blocked by the tail. In the former situation, a part or all of cluster 1 will merge with the tail and be blocked and in the later the entirely of cluster will be blocked by the tail. Either way cluster 1 is blocked and will be inside $S$ when cluster 2 arrives back. Now in case (1) all or a part of the tail will reach $S$ and be blocked while the previous cluster clears $S$. It will merge with other cells blocked at $S$ and they will be a critical cluster when they are released. Thus when cluster 1 arrives at $S$, the last cluster including a part or all of the tail will still be in $S$. Again in this case, cluster 1 is blocked before reentering $S$ and will be inside $S$ when cluster 2 arrives. Now in either case, cluster 1 is not only blocked, but resynchronized as it is blocked. The same happens for the subsequent clusters and the process continues.
3.3. **Perturbed advancing model**

First we note the lack of rigorous results for this model. We are not able to prove anything similar to Theorem 3.1. This lack of results, however, is suggestive that the advancing model does not form clusters as naturally as does the blocking model.

For this model, we can make the following observation: ‘In the advancing model with small random perturbations and $|R| = |S|$, an initial single critical cluster will have “forward leakage.”’ Wherever a cluster begins in the cell cycle, when its edge reaches $s$ it will have a positive width. As some of the cells enter $S$ the threshold will be exceeded, and the rest of the cells will be promoted to $s$. The cluster will then proceed around the cell cycle. When its edge reaches $S$ again, its width will be larger and the leading edge flatter than during the previous cycle. Thus the front edge will reach deeper into $S$ before the advance takes place. Thus, the front edge of the cluster will grow wider and wider on each pass until some of the cells are able to pass through $S$ completely before advance occurs. At this point, these cells have essentially escaped from the cluster.

Note that the cells that leak forward may eventually slow down and drop back into the cluster, or, if they continue at a faster rate then the cluster will eventually be caught in the next cluster (or into the original cluster if it is the only one). This implies that there may be steady states that have persistent clustering, but the cells in the clusters may not actually be synchronized in the strict sense.

4. **Linear feedback models and simulations**

4.1. **Graduated but unstratified model**

Next we consider more realistic models. Rather than strict advancing and blocking we consider the possibility that the number of cells in $S$ either act to slow down or speed up cell growth in the preceding region $R$. The population of cells in the $S$ phase is hypothesized to effect the growth of cells in a preceding phase $R = [r, s]$ via a continuous function. This influence will be assumed to be a function of the number of cells in the $S$ phase. Thus, we have the following equations of motion:

$$
\begin{align*}
    dx_i &= \begin{cases} 
        dt + \sqrt{\sigma} \, dW_i & \text{if } x_i \not\in R \\
        (1 + F(\#\{\text{cells in } S\}) \, dt + \sqrt{\sigma} \, dW_i & \text{if } x_i \in R.
    \end{cases}
\end{align*}
$$

Note that this form is quite general since we might consider any functional dependence $F$ and $R$ could be large or small.

We report simulations of this model where we will assume $F$ to be a linear function $F(x) = ax$, with $a < 0$ for the negative feedback model and $a > 0$ for the positive model. For example, a nonlinear model would be obtained if $F$ was a sigmoidal Hill function $F(x) = A/(1 + (x/\tau)^n)$. In the negative case in the limit of Hill coefficient $n \to \infty$, we obtain a strict blocking model described in Section 2.

In the simulations, we also incorporate a small diffusive term (white noise) with a variation $\sigma$. The programme tracked the trajectories of 5000 individual cells, initially uniformly distributed, through 20 unperturbed cell cycles (20 units of time in Equation (7)). The simulation used the Euler method to integrate the SDE. In each of Figures 3–6, we show (a) a histogram of the final distribution of cells within the cell cycle and (b) a time series of the fraction of cells inside $S$ over the final two periods.

The figures clearly show clustering for all the specified parameter values.

We observe the following things from the simulations:

- The negative feedback produces more clusters than positive.
Figure 3. Simulations of the graduated model with positive feedback. Here $R = [0.1, 0.2], S = [0.2, 0.3], \sigma = 0.02$ and the linear factor was 5. (a) Histogram of the final distribution of cells within the cell cycles; (b) Time series of the final two time frames. One unit of time corresponds to one unperturbed cell cycle.

Figure 4. Simulations of the graduated model with negative feedback. Here $R = [0.1, 0.2], S = [0.2, 0.3], \sigma = 0.02$ and the linear inhibition factor was $-5$. (a) Histogram of the final distribution of cells; (b) Time series of the final two time frames.

Figure 5. Simulations of the graduated model with positive feedback. Here $R = [0.15, 0.2], S = [0.2, 0.25], \sigma = 0.01$ and the linear factor was 10. (a) Histogram of the final distribution of cells; (b) Time series of the final two time frames.
The number of clusters formed is inversely proportional to the width of $S$ or $|R| + |S|$. Positive feedback causes sharper clustering, i.e. the clusters appear to be completely separated. Too much noise can destroy the clustering effect. The amount of noise needed to do so was smaller for larger numbers of clusters.

The first two of these observations are consistent with the results concerning the idealized advancing and blocking models. The third observation is somewhat in contrast to the lack of rigorous results in Section 3.3. The stochastically perturbed positive feedback model is seen to form clusters just as readily and even more markedly than the negative model. These simulations with more realistic models reinforce the hypothesis that clustering is a robust phenomenon in cell cycle dynamics with any form of feedback.

### 4.2. Stratified model

When yeast cells divide, one is the mother and the other the daughter. The mother carries a scar from the bud and the daughter does not. The number of daughters a cell has had can be determined by counting the bud scars. We will refer to a cell’s generation as the number of daughters it has produced, starting the count from 0 for the daughters themselves in the first cell cycle.

It is known that after division the mother’s volume is slightly larger than that of a daughter. Further, the later generations have slightly shorter times to budding and slightly shorter cell cycle times.

Leslie proposed a general stratified population model that takes into account the differences in generations that we adapt to yeast in [44]. As before, we will simplify the model by considering logarithmic coordinates, normalized by the cell cycle length of the 0-th generation. In these coordinates, the cell cycle of the 0 generation is the unit interval $I_0 = [0, 1]$. We will denote the S phase and R phases of the 0 generation by $R = [r_0, s_0]$ and $S = [s_0, t_0]$. The successive generations in these coordinates are represented by the intervals $I_k = [0, D_k]$ with $R_k = [r_k, s_k]$ and $S_k = [s_k, t_k]$. Note first that if $D_k = 1$, $R_k = R_0$ and $S_k = S_0$ for all $k \geq 1$, then this model reduces to the simplified model of the previous sections. Next we note that if these conditions approximately hold, then the above analysis can be applied in the same way as the small random perturbations.

If there are marked differences in the parameters between generations, then we still might be able to repeat a rigorous analysis. For example, suppose that $D_k = 2/3$ for all $k > 0$. Then in the idealized models, we could have three clusters in the 0-th generation and two clusters in the higher generations. In this scenario, at division parents would be synchronized with a cluster of
daughters, which was previously one-third of a cycle ahead of them. As another example, suppose that the ratio above is three-fourth, then the system could support four daughter clusters and three parent clusters. In the same way, rational ratios of cell cycles could produce a variety of clustering combinations.

In real systems, first of all, ratios close to the above idealized ‘resonances’ could still lead to clustering. Second, we may suppose that only the first few generations matter since higher generations are represented in numbers that decay approximately geometrically. Biologically, the successive generations beyond the second have not been found to exhibit marked differences anyway.

5. Discussion

The main conclusion we wish to emphasize is that clustering seems to be a very robust phenomenon; it occurs in all the models studied and for a large spectrum of parameter values. This later point is quite important since the actual systems in question are so complex that many of the parameter values are difficult to accurately determine.

The observed synchronous behaviour here is not driven by the cell cycle itself, but by feedback mechanisms acting on the cell cycle. However, the clusters must necessarily be an integer in number, and so the oscillations produced by clustering would naturally appear with a period that is an integer fraction of the cell cycle period.

We observe two phenomena that possibly make negative feedback a more reasonable explanation of yeast cell-cycle synchrony. First, negative feedback allows for large number of clusters as in some experiments. Second, negative feedback is seen to initiate clustering more naturally, which we observe in analysis of the idealized models. However, either positive or negative feedbacks are possible agents of clustering and conclusive evidence of either cause would need extensive biological modelling confirmed by experiment. Currently, there is a strong interest in modelling the details of the cell cycle.

Given the robustness of clustering, we suspect that it occurs in many other biological systems with cell cycles. Specifically, it could play a role in many types of microbiological systems with cell cycles and some type of signalling, including bacteria [9,12,15,32,33].

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References


Appendix 1. PDE models of the cell cycle

In this appendix, we turn our attention briefly to partial differential equation (PDE) approximations of the models. The PDE models derived are presented to give some context to the ODE, RDE and SDE models that we consider in the rest of the paper. Also, there is some history of such models in the literature.

A.1 General PDE model

Let $U(t,x)$ denote the distribution of cells within the cell cycle as represented by $x \in [0,1)$, i.e.

$$U(t,x) = \frac{1}{N} \sum_{i \in S(t)} \delta_{S_i(t)}(x),$$

where $S(t) \subset S$ is the index set of cells that are living at time $t$ and $N$ might be taken as the average number of active cells. As is customary, we may approximate the point distribution by a more regular distribution $u(t,x)$.

If there is no death or harvesting, then formally a distribution $u(t,x)$ evolves locally under Equation (2) by the Fokker–Planck equation, which takes the form:

$$\frac{\partial u}{\partial t} + \frac{\partial}{\partial x} (a(t,x,[u])u) - \sigma \frac{\partial^2 u}{\partial x^2} = 0. \tag{A1}$$

In this notation, $[u]$ denotes a functional dependence on $u$ and possibly its history, for example via an integral operator. Note that this derivation is not standard since the cells in the distribution are not independent from each other, but coupled through the term $u$ and the resulting equation is inherently nonlinear. Thus for rigour, Equation (A1) requires further justification.

If the culture is well mixed and harvesting is via removal of bulk material, as in a bio-reactor, then all cells are equally likely to be harvested, regardless of the state and the effect of harvesting on the density will be proportional to $u(t,x)$. Similarly, if death of a cell is equally likely at any stage of the cell cycle, then the harvesting and death can be incorporated into the Fokker–Planck equation as a term $-k_u$ on the right-hand side, i.e.

$$\frac{\partial u}{\partial t} + \frac{\partial}{\partial x} (a(t,x,[u])u) - \sigma \frac{\partial^2 u}{\partial x^2} = -k_u. \tag{A2}$$

To take cell division into account in the PDE model, we need to require the boundary conditions:

$$u(t,0) = 2a(t,1^-), \quad a(t,0,[u])u(t,0) + \sigma u_2(t,0) = 2a(t,1^-, [u])u(t,1^-) + 2\sigma u_2(t,1^-). \tag{A3}$$

Finally, in a PDE model we may represent $a$ as an integral operator:

$$a(t,x,[u]) = b(t,x) + \int_0^1 k(t,x,u(z),z) \, dz.$$
the dependence on $t$ is averaged over one period, and that $\gamma$ is small. Then Equation (A2) may be approximated by

$$\frac{\partial u}{\partial t} + \frac{\partial}{\partial x} (a(x) u) = -ku.$$  \hspace{1cm} (A4)

If we make the change of variables

$$u(t, x) = e^{-k \int_0^x 1/a(y) \, dy} z(t, x),$$

then an easy calculation shows that $z(t, x)$ satisfies

$$\frac{\partial z}{\partial t} + \frac{\partial}{\partial x} (az) = 0.$$  \hspace{1cm} (A5)

Further, the balance between the growth and harvesting implies $k = \ln 2/(\int_0^1 1/a)$. This implies that the boundary conditions on $z$ are

$$z(t, 0) = z(t, 1).$$  \hspace{1cm} (A6)

The ODE that generates Equation (A5) is simply

$$\frac{dx}{dt} = a(x),$$

on $[0, 1)$. If we reincorporate the other influences on the growth rate, then the unperturbed and perturbed equations become

$$\frac{dx_i}{dt} = a(t, x_i, \bar{x})$$  \hspace{1cm} (A7)

and

$$\frac{dx_i}{dt} = a(t, x_i, \bar{x}) + g_i(t).$$  \hspace{1cm} (A8)

Note that Equation (A5) with Equation (A6) is a conservative equation in the sense that $\int z \, dx$ is preserved, thus in the ODE (A7) and RDE (A8) the total number of cells considered should not change. It is thus reasonable to take $S$ to be a fixed set \{1, 2, \ldots, N\} and let $x_i(t)$ to be defined for all $t \in [0, T]$. To accommodate division, we may reset $x_i(t)$ to zero each time it reaches one.

**A.3 Notes the PDE models and related work**

The importance of the cell cycle has been recognized in hematopoiesis [13,14], as well as yeast growth [17], where PDE models similar to Equation (A4) are considered with $a$ assumed constant. Thirty years ago, Rotenberg [39] described the PDE model (A4) for yeast growth, but with constant coefficients and an externally applied periodic blocking at cell division. He demonstrated numerically that clustering occurs in the model. Surprisingly, this article has received few citations and little attention. A trivial, but important step, common to our work and that of Rotenberg, is normalization of the cell cycle. This provides a standard domain on which to consider both ODE and PDE models.

In related recent work [50], Zhu et al. have considered population balance models coupled with a substrate balance equation, where $a = a(s)$ with $s$ the effective substrate concentration. In [21], $a = a(s, x)$ is admitted, but the focus is on a survey of possible numerical methods, not analysis of the model, and clustering solutions are not discussed. Note that the models considered in the current manuscript implicitly incorporate the substrate balance equation (and also the density of any other signaling compounds) into the term $a$ via $|u|$. 
