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Highlights

- A multiscale model for the cell cycle-circadian clock coupling is developed
- The model is based on a transport PDE structured by molecular contents
- A particle-based method is used for resolution
- Impacts of inter and intracellular dynamics on cell proliferation are studied
- Discordance of division rhythms between population and single cell levels is observed
A multiscale modelling approach for the regulation of
the cell cycle by the circadian clock

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Abstract

We present a multiscale mathematical model for the regulation of the cell cycle by
the circadian clock. Biologically, the model describes the proliferation of a population
of heterogeneous cells connected to each other. The model consists of a high dimen-
sional transport equation structured by molecular contents of the cell cycle-circadian
clock coupled oscillator. We propose a computational method for resolution adapted
from the concept of particle methods. We study the impact of molecular dynamics
on cell proliferation and show an example where discordance of division rhythms be-
tween population and single cell levels is observed. This highlights the importance of
multiscale modeling where such results cannot be inferred from considering solely one
biological level.

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1 Introduction

The mammalian cell cycle and the circadian clock are two molecular processes that operate in a rhythmic manner. On one hand, the cell cycle is driven by the rhythmic activity of cyclin-dependent kinases, which dictate the time a cell must engage mitosis and the time it must divide giving birth to two daughter cells. On the other hand, the circadian clock is a network of transcriptional and translational feedback-loops that generate sustained oscillations of different mRNAs and proteins concentration with a period of approximately 24h. It turns out that several components of the circadian-clock regulate various cyclin-dependent kinases at different stages of the cell cycle. This makes the circadian clock a key player in the temporal organization of the cell cycle and makes these two biological processes act as two tightly coupled oscillators.

Many computational models explored the interaction between the cell cycle and the circadian clock. One category of models consists of systems of ordinary differential equations that describe the interaction at the molecular level [14, 33]. The second category consists of physiologically-structured partial differential equations that describe the effect of circadian regulation on the growth rate of cells [8, 9, 2, 7]. In a recent study, we have combined these two approaches by coupling an age-structured PDE to a circadian clock-cell cycle molecular ODE system [12]. Although this model could capture the influence of intracellular dynamics on the growth rate of cells, it lacked a proper multiscale description [1]. It could not take into consideration, for example, intracellular heterogeneity among cells.

Here, we present a multiscale formulation of our mathematical model for the regulation of the cell cycle by the circadian clock and a numerical method for solving it. The multiscale formulation consists of a transport equation structured by the molecular contents of the coupled cell cycle-circadian clock oscillator. In its general form, the equation reads

$$\partial_t \rho(x,t,\lambda) + \nabla_x \cdot [u(x,t,\lambda,\psi)\rho(x,t,\lambda)] = L[x,\lambda](\rho(x,t,\lambda)), \quad (1.1)$$

with $\rho$ representing the density of cells. In the coupled PDE/ODE model, the PDE (equation (11) in [12]) was structured in age. By contrast, partial derivatives in equation (1.1) are structured by the molecular contents (denoted $x$) and the space where it is solved is $d$-dimensional with $d$ the number of molecular components $x_i$; this is why we call it a molecular-structured equation. Since the molecular mechanism of the cell cycle-circadian clock interaction involves an abundant number of components, the model is high-dimensional in nature. Classical numerical methods such as finite
volumes/differences are inappropriate for solving the transport equation. This is the main difficulty that makes molecular-structured models scarcely used in similar applications. We circumvent this difficulty by adapting a particle method to solve the main equation.

Particle methods have arisen as an alternative to classical numerical methods for solving high-dimensional problems. They are used in different applications, like the incompressible Euler equations in fluid mechanics [17, 19, 20], the Vlasov equation in plasma physics [3, 15] and in turbulence models for reactive flows [25, 24]. The computational implementation of particle methods is conceptually simple. At a given time, the solution is represented by a large number of particles, each having its own properties, for example position and weight. These properties evolve in time according to a system of stochastic or ordinary differential equations. The classical solution to the PDE is obtained by reconstructing a smooth density from the particles distribution. Theoretically, the numerical solution is a linear combination of Dirac masses

$$\rho(x, t) = \sum_{j=1}^{N} \alpha_j(t) \delta(x - x^j)$$

where $\alpha_j$ is the weight of particle $j$, $x^j$ its position (state) and $N$ is the total number of particles. To obtain a classical solution, one has to update the positions and weights of particles and then regularize the Dirac masses. The overriding strength of particle methods is that, for $N$ fixed, the size of the system increases only linearly with the dimensionality of the space. This means that if we have a structured equation with dimension $d$, we have to solve $d \times N$ ODEs/SDEs. Since recovering a classical solution requires a regularization of the Dirac distribution, the performance of particle methods depends also on the quality of the regularization procedure. For generalities about particle methods, one can refer to [23, 27, 10].

In this work, we investigated the influence of molecular dynamics on cell proliferation. Therefore, the novelty of our approach emanate from considering a multiscale design that incorporates three linked biological levels. Hereafter some specificities of our model:

- **Intracellular heterogeneity among cells:** generally speaking, the particle method consists of solving $N$ copies of an ODE/SDE system, each representing the dynamics of a given particle. In our model, the $N$ copies are not the same. Since each particle is associated with a given cell, we considered several sources of variability. For example, we have multiplied the
right-hand side of the ODE system describing cell cycle dynamics by a factor \( \lambda \). This ensures that each cell has a distinct cell cycle period. Another source of variability was introduced with cell division. Once a cell divides, its daughter is attributed a new coefficient \( \lambda \) given by \( \lambda^{N+1} = \lambda^N + D \xi_\lambda \), with \( D \) a “diffusion” coefficient and \( \xi_\lambda \) a normally distributed random number. This implies that the daughter and mother cells will have distinct cell cycle periods.

- **Inter-cellular connection:** we assumed that cells are connected with each other by making Per/Cry mRNA transcription dependent on the average concentration of Per/Cry among cells. Including connectivity between cells is important in certain tissues to maintain a robust synchronized activity.

- **Coupling back from population to molecular level:** we assumed that cell division is dependent on the MPF-WEE1 molecular activity and that the total number of cells has an impact on Per/Cry mRNA activity. This implies a two way coupling between the molecular and population levels. We assumed also that the MPF activation coefficient decreases at an exponential rate proportional to the total number of cells. This ensures a limited growth, which is physiologically more realistic.

The paper is divided mainly into two parts. First, we introduce the model structure, give details about its construction and how the transport equation is solved by adapting a particle method. Then, we illustrate biological properties that can be examined using our model; like the effect of intracellular dynamics on cell proliferation, the synchronization among cells, and how heterogeneity among cells affects growth rate.

2 Description of the model

2.1 General framework and main equation

We consider a large collection of cells with state \((x, \lambda) \in \Omega \times \Lambda \subset \mathbb{R}^d \times \mathbb{R}^p\), where \( x \) is a cellular dynamical state in an open subset \( \Omega \) of dimension \( d \) and \( \lambda \) a cellular parameter state in an open subset \( \Lambda \) of dimension \( p \). In our case, \( x = (x_1, \cdots, x_d) \) with \( d = 10 \) represents the set of proteins/mRNAs concentrations and \( \lambda \in \mathbb{R} \) (\( p = 1 \)) represents the intrinsic cell cycle period of each cell. The distinction between the dynamical and the parameter state is that the dynamical state changes during the lifespan of a cell, while
the parameter state is fixed over the lifespan of the cell, but can vary among cells or
when a cell divides. The population is described by its density $\rho(x, t, \lambda)$ which evolves
according to the following hyperbolic PDE

$$\frac{\partial \rho}{\partial t}(x, t, \lambda) + \nabla_x \cdot [u(x, t, \lambda, \psi) \rho(x, t, \lambda)] =$$

$$+ \frac{1}{2} \Delta_\lambda \cdot [\sigma^2(x, t, \lambda) R(x, \lambda) \rho(x, t, \lambda)] + r(x, \lambda) \rho(x, t, \lambda). \tag{2.2}$$

We assumed that the cellular state dynamics $x$ is governed by a deterministic system of
ODEs, represented by the term $u(x, t, \lambda, \psi)$, which depends on the cellular state $(x, \lambda)$
and on $m$ population-level statistics $\psi : \Omega \times \mathbb{R} \times \Lambda \to \mathbb{R}^m$ where

$$\psi_i = \langle \rho, \Phi_i \rangle = \int_{\Omega \times \Lambda} \rho(x, t, \lambda) \Phi_i(x, t, \lambda) dxd\lambda, \quad i = 1, \cdots, m \tag{2.3}$$

with $\Phi_i : \Omega \times \mathbb{R} \times \Lambda \to \mathbb{R}, \quad i = 1, \cdots, m$ taken such that $\psi_i$ is finite. The operator $L[x, \lambda]$
describes the population relative growth rate. The terms $\sigma$, $R$ and $r$ are diffusion in
parameter state, differentiation and growth rates of the population, respectively. For
the sake of simplicity, we omitted the full description of the derivation of equation (2.2).
The reader could refer to Supplementary materials for more details. We adopted a more
computational and biological description of the problem.

### 2.2 Intracellular dynamics

The evolution of cells in the state space depicts the cell cycle-the circadian clock cou-
pling mechanism. Therefore, intracellular dynamics were described according to the
following deterministic ODE system presented in [12]:

$$\frac{dx}{dt} = u(x, t, \lambda, \psi). \tag{2.4}$$
where

\[
\begin{align*}
    u_1 &= \frac{d x_1}{dt} = \frac{\nu_1(x_7 + c)}{k_{1b}(1 + (\frac{x_7}{k_{1a}})^p)} + x_7 + c - k_{1d} x_1 + k_s \psi_1, \\
    u_2 &= \frac{d x_2}{dt} = k_2 b x_1^q - k_2 d x_2 - k_2 h x_2 + k_3 x_3, \\
    u_3 &= \frac{d x_3}{dt} = k_2 i x_2 - k_3 i x_3 - k_3 d x_3, \\
    u_4 &= \frac{d x_4}{dt} = \frac{\nu_4 x_5}{k_{r_b} + x_3^p} - k_{1d} x_4, \\
    u_5 &= \frac{d x_5}{dt} = k_5 b x_4 - k_5 d x_5 - k_5 i x_5 + k_6 i x_6, \\
    u_6 &= \frac{d x_6}{dt} = k_5 i x_5 - k_6 i x_6 - k_6 d x_6 + k_7 a x_7 - k_6 a x_6, \\
    u_7 &= \frac{d x_7}{dt} = k_6 a x_6 - k_7 a x_7 - k_7 d x_7, \\
    u_8 &= \frac{d x_8}{dt} = \lambda \left( \frac{k_{imp} + k_{impf} \exp^{-q_2}}{k_{imp} + x_8 + x_{10}} \right) k_{impf} (1 - x_8) - d_{wee} x_8 x_9, \\
    u_9 &= \frac{d x_9}{dt} = \lambda \left( \frac{k_{act} w}{k_{act} + d_{w1}} (c_w + C x_7) + \left( \frac{k_{act}}{k_{act} + d_{w1}} \right) \frac{k_{inact} w x_8 x_9}{k_{inact} w + x_8} - d_{w2} x_9 \right), \\
    u_{10} &= \frac{d x_{10}}{dt} = \lambda \left( k_{act} x_8 - x_{10} \right).
\end{align*}
\]

Equations (2.5-2.11) describe the circadian oscillator and equations (2.12-2.14) the cell cycle. Circadian dynamical variables are \( x_1, \) \( Per2 \) or \( Cry \) mRNA and proteins; \( x_2, \) PER2/CRY complex (cytoplasm); \( x_3, \) PER2/CRY complex (nucleus); \( x_4, \) \( Bmal1 \) mRNA; \( x_5, \) BMAL1 cytoplasmic protein; \( x_6, \) BMAL1 nuclear protein and \( x_7, \) active BMAL1. Cell cycle dynamical variables are \( x_8, \) active MPF; \( x_9, \) active WEE1 and \( x_{10}, \) active MPF inhibitor. The coupling between these two oscillators is taken into consideration in the term \( C x_7 \) of equation (2.13) through the regulation of WEE1 by BMAL1/CLOCK. The coefficient \( C \) is the coupling strength. Details about this model can be found in [12]. The cellular dynamics is now part of a multiscale system, some changes were added to the original system to reflect that cells interact with each other. These changes are detailed in the coming paragraphs.

### 2.3 Population synchronization

Including connectivity between cells is important in certain tissues to maintain a robust synchronized activity. It is known, for example, that neuronal clocks within the the suprachiasmatic nucleus SCN form a heterogeneous network that must synchronize to
maintain time keeping activity. The coherent output of the SCN is established by intracellular signaling factors, such as vasointestinal polypeptide [13]. A simple way to induce synchronization in our model is to make Per/Cry mRNA transcription depends on the average concentration of Per/Cry among cells. For that, we included in equation (2.5) the term $k_s \psi_1 := k_s \langle \rho, \Phi_1 \rangle$ with $k_s$ a coefficient that represents the connectivity strength and $\Phi_1 = x_1$. Although peripheral tissues, as modeled here, would run out-of-phase without the SCN, there are so many factors that can synchronize clocks that it is relevant to study the possibility for peripheral tissues to be connected.

2.4 Variability among cells

Variability among cells arises naturally from the difference in their molecular contents. This was taken into consideration in our model through the initial states of cells, which can be chosen randomly. We added another source of variability through the parameter $\lambda$. We assumed that each cell has a distinct cell cycle period. This was modeled by multiplying equations (2.12-2.14) by a scaling factor $\lambda$. Cell division can also induce variability by assigning to each new born cell an intrinsic cell cycle period that is different from its mother cell. (This is accomplished by taking a non trivial distribution $p(\lambda|z)$, see Supplementary materials equations (1.4), (1.5)).

2.5 Cell division

A central aspect of our model is that it takes into account cell division. Cell division timing is based on the antagonistic relationship between the mitosis promoting factor MPF and the protein WEE1. It is assumed that a cell enters mitosis once the activity of MPF surpasses that of WEE1 and then divides once MPF activity shuts down abruptly. Based on this mechanism, we considered two division types, one deterministic and one stochastic. The deterministic division occurs exactly every time MPF activity rises above WEE1 activity and then shuts down. This was taken into consideration with a growth rate $r(x, \lambda) := r_d(x, \lambda) = \delta_{x \in \Gamma}$. For the stochastic division, we considered that a cell divides with a certain probability $\Delta t \times r(x, \lambda) + o(\Delta t)$ for a small time step $\Delta t$. The division rate $r$ is a function of $x$ and $\lambda$ that mimics the deterministic case.

For example, a switch-like function that takes small values on one side of $\Gamma$ and large values on the other side could be used (see Supplementary materials for more details about the growth rates and $\Gamma$). Here, we used the Koshland-Goldbeter function [16]
given by
\[
K(y,z) = \frac{2yJ_i}{z - y + zJ_a + yJ_i + \sqrt{(z - y + zJ_a + yJ_i)^2 - 4yJ_i(z - y)}}.
\]  
(2.15)

This function generates a switching behavior [21]. If the ratio \( y/z \) (\( y \) and \( z \) are dummy variables here) becomes larger than one, the function switches to the upper state and the transition occurs. \( J_a \) and \( J_i \) are two constants that determines the stiffness of the switch, if they converge to zero, the switch converges to a step function.

### 2.6 Limited growth

To make sure the growth is bounded (for physiological and computational reasons), we assumed that cell proliferation slows down when the total number of cells reaches a threshold value. We assumed that the activation coefficient of MPF decreases at an exponential rate proportional to the total number of cells. This was taken into consideration in equation (2.12) by multiplying MPF activation coefficient \( k_{\text{mpf}} \) by \( \exp^{-\eta \psi_2} \) where \( \psi_2 = \langle \rho, \Phi_2 \rangle \) with \( \Phi_2 = 1 \) (\( \psi_2 = N(t) \)). When cell number is large enough, MPF activity cannot increase above that of WEE1, and cell division is blocked.

### 2.7 Method of resolution

The main difficulty for solving equation (2.2) comes from the high dimension of the dynamical state space (10 in our case). For that, we adapted a particle method that is more appropriate in high dimensions than classical methods such as finite differences/volumes.

The intuition behind the particle method is to start with a distribution of particles that approximates the initial condition and then follow the evolution of these particles in time in the dynamical and parameter states space. Mathematically, the particle solution is a discrete measure function, which means it is irregular. To obtain a solution in the usual classical sense at a given time \( T \), one has to recover the classical solution with regularization techniques. The method is mainly divided into two steps

#### 2.7.1 Step 1: approximation of the initial condition

Given an initial condition \( \rho^0(\mathbf{x}, \lambda) \in C^0(\mathbb{R}^d \times \mathbb{R}) \), we take an initial set of particles \( \mathbf{x}^j \) with weights \( \alpha_j \) such that \( \rho^0_h = \sum_j^{N} \alpha_j \delta[(\mathbf{x}, \lambda) - (\mathbf{x}^j, \lambda^j)] \) approximates \( \rho^0 \). This should
be understood as an approximation in the sense of measures, which means that one looks for a test function \( \phi \in C^0_0(\mathbb{R}^d \times \mathbb{R}) \) such that

\[
\langle \rho^0, \phi \rangle = \int_{\mathbb{R}^d} \rho^0 \phi dx d\lambda \quad \text{and} \quad \langle \rho^0_h, \phi \rangle = \sum_j \alpha_j \phi(x^j, \lambda^j).
\]

This yields a typical numerical quadrature problem where the integral \( \int_{\mathbb{R}^d} \rho^0 \phi \) is approximated by \( \sum_j \alpha_j \phi(x^j, \lambda^j) \). For biological purposes, in our study we do consider directly an initial condition of the form (\( \alpha_j = 1 \))

\[
\rho^0_h = \sum_j^N \delta[(x, \lambda) - (x^j, \lambda^j)].
\]

### 2.7.2 Step 2: particle evolution in time

In the dynamical state space the particles positions \( X^j(t) \) (not to confuse between \( X^j \) and \( x^j \), our notations implies that \( X^j(0) = x^j \)) can be computed at a given time \( t \) by solving the following ordinary differential equation system

\[
\frac{d}{dt} X^j(t) = u(X^j(t), t). \quad (2.16)
\]

To take into account the source term in equation (2.2), we examined the intracellular state of each cell (position of \( X \) in the state space) after each time step \( \Delta t \) and looked if the biological conditions are met for a cell to divide. This means that if for a given particle, MPF activity rises above that of WEE1 and then shuts down under a threshold level, we add a particle \( X^{N+1} \) with intracellular state similar to the mother cell or randomly chosen. If division occurs, a new parameter \( \lambda^{N+1} = \lambda^N + D \xi_\lambda \) is assigned to the particle of number \( N + 1 \), with \( D \) a diffusion coefficient and \( \xi_\lambda \) a normally distributed random number. Hence the measure solution at a given time \( T \), is given by

\[
\rho_h(x, T) = \tilde{N} \sum_j \delta[(x, \lambda) - (X^j(T), \lambda^j(T))]. \quad (2.17)
\]

with \( \tilde{N} \) the new number particles.

Commonly, with particle method one needs to obtain a solution in the classical sense, which means that the particle solution (2.17) should be
regularized. This is usually accomplished by taking the convolution product of $\rho_h$ with a smoothing kernel $\zeta_\epsilon$. In our case, we did not need to obtain a smooth solution, since there is no explicit dependence of the quantities of interest on the density of cells $\rho$. We focused on studying statistic-like properties of the distribution of cells. Hence, we did not perform any regularization procedure on the particle solution. For more information about particle methods the reader could refer to [24, 27].

3 Results

3.1 Regulation of the cell cycle by the circadian clock

We solved equation (2.2) with $L = 0$, i.e. without division. We assumed that each cell has an intrinsic cell cycle period included in the interval $[20 \text{ h}, 28 \text{ h}]$. This was done by assigning to each particle a different coefficient $\lambda$. We let all cells have the same initial molecular state and considered two cases; one without coupling to the circadian clock ($C = 0$), and one with coupling to the circadian clock ($C = 1.2$). We plotted a projection of the solution in the subspaces $(x_7, x_9)$ and $(x_7, x_8)$, which are the subspaces of molecular concentrations (BMAL1/CLOCK, WEE1) and (BMAL1/CLOCK, MPF). We obtained a random distribution of particles for the non-coupled case and a limit cycle distribution for the coupled case (Figure 1). This means that the circadian clock is forcing all cells to oscillate with the same period. This is in agreement with results obtained earlier [12]; it was shown that a population composed of cells with cell cycle periods between $20$ h and $28$ h are brought to oscillate with a unique cell cycle period of $24$ h for large values of $C$.

3.2 Cell division

We took four cells initially, two of them with $\lambda = 230$, and two with $\lambda = 180$ (intrinsic cell cycle lengths $\approx 10$ h and $12.7$ h respectively). These values were taken randomly. We solved equation (2.2) with a source term and considered three cases, one with $r = r_d$ (deterministic growth) and the two other ones with $r = r_s$ (stochastic growth). For the stochastic growth, we used a stiff and a non-stiff Koshland-Goldbeter function. First, we fixed the value of $\eta$ to 0, which means that MPF activity does not depend on the total cells number. We assumed that newborn cells retain the cell cycle length of their mother cells at the division time, hence $p(\lambda|z) = \delta(\lambda - z)$ which leads the growth term to ne $L = r(x, \lambda)\rho$. This ensures that MPF cycle does not change along birth. Our
Figure 1: Solution of equation (2.2) without cell division. (A,B) Projection on the subspace \((x_7, x_9)\); (C,D) projection on the subspace \((x_7, x_8)\). (A,C) Without coupling to the circadian clock, \(C = 0\). (B,D) With coupling to the circadian clock \(C = 1.2\).
simulations showed that the total number of cells after four days was equal to 1152 for the first case, 900 for the second one with a stiff Koshland-Goldbeter function and 480 with a non stiff one. This is natural and is justified by the fact that the division rate $r_d$ is a Dirac-type division which means that division depends deterministically on MPF activity. Looking at MPF activity, we see that it peaks 9 times for $\lambda = 230$, and 6 times for $\lambda = 180$ every 4 days (Figure 2A,B). If division occurs exactly once MPF activity accomplishes a normal cycle, the total number of cells should be $2 \times 2^9 + 2 \times 2^6 = 1152$, which is the result obtained with $r = r_d$ (Figure 2C). The division rate $r_s$ introduces stochasticity in the decision for division. In this case, division does not depend only on MPF-WEE1 activity, but also on the probability $r_s \times \Delta t$ at each time step $\Delta t$ Figure (2C).

Second, we assumed that proliferation depends on the total number of cells, and set $\eta = 2 \times 10^{-3}$ (increasing the value of $\eta$ will make the factor $\exp^{-\eta \psi_2}$ less than one and hence decreases the value of MPF activation coefficient $k_{mpf}$). We took initially 100 cells with different intrinsic cell cycle periods and followed the total number of cells over 15 days. We remarked that division stopped after 8 days, and the growth curve reached a steady state (Figure 2D). Increasing the value of $\eta$ decreases the activity of MPF with increasing number of cells. This means that at a certain time, MPF activity will decrease below a threshold that does not allow the cell to start mitosis.

### 3.3 Synchronization

To study the synchronization of cells, we compared the molecular concentrations of a cell chosen randomly with the average molecular concentration of all cells. We let the rate of production of Per2/Cry mRNA of each cell to be dependent on the average value for Per2/Cry mRNA. This was done by taking $k_s = 0.05$ in equation (2.5). We took a population of 100 cells with autonomous cell cycle periods randomly distributed between 20 and 28 hours. Initial molecular concentrations were chosen randomly between 0 and 1 and each cell has a distinct initial molecular state. We did not consider division hence $L = 0$, but we considered coupling between the cell cycle and the circadian clock ($C = 1.2$). We followed the evolution of three components, Per2/Cry mRNA, BMAL1/CLOCK and WEE1, over 20 days. Our simulations showed that for $k_s = 0$, the average molecular concentration had small oscillations which are asynchronous with those of the random cell concentration (Figure 3 A,B,C). Whereas for $k_s = 0.05$, we obtained that the average molecular concentration tends to coincide with that of the random cell (Figure 3 D,E,F). This indicates that all cells are oscillating in a similar manner (same period and phase), and indicates synchronization of all popu-
Figure 2: (A) MPF-WEE1 antagonistic activity for $\lambda = 180$: division occurs if MPF activity rises above that of WEE1 and then shuts down. (B) Impact of $\lambda$ on MPF activity. (C) Total number of cells with different source terms with no dependence of MPF activity on the total number of cells ($\eta = 0$). Initial number of cells = 4, two cells with $\lambda = 230$ and two with $\lambda = 180$. (D) Impact of the dependence of MPF activity on the total number of cells ($\eta = 2 \times 10^{-3}$).
lation cells. In the case $k_s = 0$, even though cells are coupled in the same manner to their circadian clock, cells keep oscillating in an asynchronous manner. The circadian clock regulates all cells to oscillate in a similar period which is 24 h in this case but due to the randomness in the initial molecular concentrations, each cell oscillates with a different phase.

### 3.4 Heterogeneity of cells and growth rate

We studied the growth rate of a cell population where each cell had a different cell cycle period. We took 100 cells initially and conferred to each of them a coefficient $\lambda$ chosen randomly between 128 and 193. These coefficients scale the intracellular cell cycle system so that periods range randomly from 12 h to 18 h. The circadian control strength value $C$ was fixed to 1.6. We did not consider connection between cells ($k_s = 0$) and we did not consider dependence of the intracellular dynamics on any population level statistics. Simulations were done over 20 days and with a net death rate to mainly reduce the computation time. Simulations showed a growth rate with two daily peaks, suggesting that there were two division rounds every day (Figure 4A). To explain this bimodal behavior for the growth rate, we looked at the average activity of WEE1 and MPF. Simulations showed that MPF activity overcomes that of WEE1 once a day (Figure 4B). Based on average MPF activity alone, division should occur only once a day, and cannot explain the two daily peaks. We then followed a subpopulation of cells and examined at which time each cell divided. We identified three regimes for division: a single division per day, two divisions per day and three divisions every two days (Figure 4C,D). These regimes can be explained by the entrainment results obtained previously (for autonomous cell cycle periods between 12 and 18 h) showing large phase-locking regions entrainment with ratios 1:1, 2:1 and 3:2 [12]. The double peaks for the growth rate can be then explained by the fact that all cells are dividing at least once a day and some of them will undergo a second round of division (Figure 4D).

### 4 Discussion and conclusion

We presented in this work a multiscale mathematical model for the regulation of the cell cycle by the circadian clock and its influence on cell growth. The model consisted of a master transport partial differential equation of high dimension structured by the molecular contents of the coupled cell cycle-circadian clock oscillator. We proposed a
Figure 3: Comparison between population average molecular concentrations randomly chosen and single cell molecular concentrations. (A,B,C) No synchronization between cells: $k_s = 0$; (E,F,G) Synchronized activity: $k_s = 0.05$. 
Figure 4: (A) Growth rate indicating two peaks of division each day. (B) MPF-WEE1 average activity for all cells. (C) Counting the number of times each cell divides per day. Each square indicates a division. (D) Similar to figure C but simulation shows the division times for 100 cells instead of 10.
computational approach for resolution adapted from the concept of particle methods.

Several computational and theoretical studies dealt with modeling cell population connected to molecular systems influencing cell growth. Bekkal Brikci and colleagues, developed a nonlinear model for the dynamics of a cell population divided into a proliferative and quiescent compartments. This model is structured by the time spent by a cell in the proliferative phase and by the Cyclin D CDK4/6 amount [5, 6]. Ribba and colleagues developed a multiscale model of cancer growth based on the genetic and molecular features of the evolution of colorectal cancer [28]. In more recent works, Prokopiou et al. presented a multiscale computational model to study the maturation of CD8 T-cells in a lymph node controlled by their molecular profile [26]. Schluter and colleagues developed a multiscale multicompartment model that accounts for biophysical interactions on both the cell-cell and cell colonies levels [29]. Bratsun and colleagues proposed a multiscale model of cancer tumor development in epithelial tissue. Their model includes mechanical interactions, chemical exchange as well as cell division [4]. Shokhirev and colleagues developed a multiscale model to study the role of NF-κB in cell division. Their approach allows for the prediction of dynamic organ-level physiology in terms of intracellular molecular network.

All of these works emphasize on the importance of taking into account different levels of a biological system by showing results that cannot be inferred by considering solely one level of functioning. We exposed a simulation showing discordance between rhythms observed on the population and signal cell levels. This is important since it underlies the fact that biological function markers which most of the times rely on average-like information can mislead the interpretation in some cases. Our modeling approach differs from the above cited works which can be separated into two categories. One consisted of low dimension structured partial differential equations solved by classical numerical techniques [28, 5]; and the other one consisted of agent-based modeling approaches [4, 29, 30]. Our approach consists of considering a master partial differential equation in high dimension structured by molecular contents for which we propose a computational solution based on a particle method. This allowed us to structure the partial differential equation by a large number of molecular contents, providing a realistic representation of the coupled cell-cycle circadian clock oscillator. By doing so, we were able to describe both population and single cell functioning in one equation. Globally, the model takes into consideration the molecular dynamics inside a cell, connection between heterogeneous cells and the proliferation of cells.

Our results can shed light on the superiority of frequent administration of chemotherapies over the standard maximum tolerated dose (MTD). Scheduling the MTD into
multiple administrations per cycle proved to be more effective and less toxic [18, 22, 31]. We showed in section 3.4 that an apparent population growth rhythm may hide several subpopulation growth rhythms. Biological markers, as the Ki-67 index, are mostly indicative of the growth status at the population level. Therefore, one is tempted to consider that the administration of an anticancer drug during the time of the largest growth peak is the most effective. However, our simulations showed that this largest growth peak is obtained from the crossing of different subpopulation rhythms. Therefore, administrating a drug at the same time every treatment cycle may hit only one subpopulation of cells, leaving another subpopulation without treatment for a long time. This possibly leads to the emergence of resistant subpopulations and make the treatment ineffective.

The aim of this work was to provide a computational multiscale framework for the regulation of the cell cycle by the circadian clock. The use of multiscale computational tools is of growing interest in the area of drug development, especially the area of systems pharmacology. A major challenge for systems pharmacology is the development of mechanistic tools to understand how regulatory networks control variability in drug response at the organismal level [34]. Our model was constructed in the hope to overcome such challenges. It had the specificity of describing the connection between the cell cycle and the circadian clock through the regulation of the protein kinase WEE1 by the complex BMAL1-CLOCK. Irregular activity of the WEE1 kinase has been linked to ovarian and NSCLC cancers. Inhibiting this activity demonstrated interference with DNA damage response within tumor cells, leading to their death. Drugs targeting WEE1 are currently under investigation [11]. In such specific applications, our model can be of great utility. According to Zhao and colleagues, experimental settings in the process of drug development are sometimes not sufficient due to the multitude of interactions between mammalian network proteins. Actions on the desired targets may lead to adverse events occurring at another level due to the wide ramification of molecular interactions. The abundance of such interactions makes it hard to design drugs affecting only the desired targets with controlled therapeutic effects [34]. Therefore it is important to propose modeling frameworks that allow in silico experimentation in the presence of uncertain or incomplete data. In this manuscript, we have provided such an example, and showed why incomplete population level data may lead to a suboptimal treatment.

Multiscale modeling in systems medicine is growing in importance. Wolkenhauer and colleagues recently proposed different recommendations to improve the integration of multiscale models into clinical research [32]. One of their medium-term recommen-
ations is the rising need of developing computational tools and algorithms for efficient multiscale simulations. We hope the modeling approach we propose in this study represents a step forward towards this direction.
References


