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Posted Date: 25 October 2023

doi: 10.20944/preprints202310.1530.v1

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Article

Network Inference from Cancer and Epithelial Cell Co-Culture Reveal Microenvironmental Configurations That Drives Population Dynamics

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Abstract: Cellular Automata and Boolean Networks are generalizations of one another because algorithms to compute the preimage of cellular automata reveal the underlying network, i.e., the global dynamics in terms of the basins of attraction. Therefore, we hypothesize we can unveil the local dynamics of cellular automata from the basin of attraction of an inferred boolean network. Our motivation was the observation that human keratinocytes and melanoma can stick together to form clusters after eight days in co-culture. This cluster formation would be the attractor of population dynamics, and cell seeding would be the initial condition of the basin of attraction. Hence, we propose a method to estimate the rules of cellular automata, which consist of comparing the density states within each state transition and reaching a consensus among state transitions belonging to a basin of attraction. Therefore, we aim: (1) to infer a boolean network from the *in vitro* co-culture growth curve; (2) to estimate the rules of cellular automata; and (3) to implement a cellular automata for spatial dynamics simulations. The binarization of the growth curve shows high population density after four days; the estimated cellular automata rules were compatible with cell proliferation and migration in agreement with experimental observations. Spatial dynamics shows that: (1) keratinocytes exhibit higher density in neighborhoods where melanoma is present; (2) the chance of keratinocyte migration increases until the fourth day, but the probability of survival increases; and (3) space is freed for cells with maximum proliferation capacity through proliferation compensated by death. Our approach suggests that the attractor state of cell co-culture would be induced by an increase in keratinocyte migration and survival, as well as the balance of proliferation and death concerning melanoma. Our approach has the potential to offer valuable clues about microenvironmental interactions or configurations that drive population dynamics.

Keywords: cells co-culture;; population dynamic; cellular automata; boolean network; basin of attraction

1. Introduction

In cells co-culture experiments, after eight days, keratinocytes surround the melanoma clusters, limiting the growth of the melanoma cells. To be more specific, the keratinocytes are denser near the melanoma clusters and less dense farther away [1]. It's interesting that when keratinocytes are cultured with melanoma, they become less differentiated and connect with the melanoma instead of other keratinocytes nearby [2] [3]. These observations may explain biologically the different levels of tolerance to high density, which is called allelophilia, between melanoma and keratinocytes [1].

As in concepts about discrete dynamic system modeling, this spatial characteristic observed on the eighth day of the experiment would correspond to an equilibrium state or attractor of the population dynamics. Conversely, the initial distribution of cells on the cell culture plate would be the

initial condition of the basin of attraction. Thus, the system moves from the initial condition towards its attractor through state transitions like proliferation, migration, survival, and cell death, etc, causing changes in the spatial pattern of the population.

Herein, discrete dynamic system modeling approaches enables the modeling and analysis of the mechanisms governing the regulation of population density. Cellular Automata (CA) is a computational model where the cells are placed on the lattice and the state of each cell is updated at time $t + 1$, based on logical rules that consider the state of the neighborhood cells at time t such that the outcome is the emergence of spatial patterns at any given moment [4] [5] [6]. Boolean networks (BN), in turn, are models commonly used in gene expression or signaling pathway studies. They consist of a collection of variables (the nodes), with binary states determined by other variables in the network through Boolean functions [7] [8]. There are algorithms that calculate the preimage of the CA trajectory in a backward direction, thereby unveiling the network of CA dynamics also called global dynamics [9][10]. For this reasons, CA and BN are considered instances of discrete dynamical systems composed of parallel-acting components [10,11]. The dynamics of both are driven by transition functions that determine the state transitions culminating in the system's trajectory which is a path within the basin of attraction. Hence, CA are generalizations of BN one another in terms of basins of attraction [9–15].

The Square-Lattice Cellular Automata (SLCA) and a two-dimensional version of Elementary Cellular Automata (ECA) has been successfully used to simulate avascular and vascular tumor growth [16–23] and invasion [24–27], tumor interactions with environmental cues [28–34], stromal composition [34–38] and tissue architecture [39–42]. Similarly, BN has been applied in many fields, including quantitative System Pharmacology [43], competition in microbiome [44], stability of apoptosis [45,46], epithelial-mesenchymal transition [47] and target therapy [48].

Here, we argue that once one can find the global dynamics of an CA in terms of basin of attraction, through an inferred boolean network (IBN) basin of attraction, one could discover the local dynamics of a CA to model a given population dynamic. The primary dimensional unit in a cell culture plate is the cell density ($cells/mm^2$), and one can qualitatively characterize it as "high" or "low," like the confluence in cell plate culture. Given that CA dynamics are based on logical rules about the configuration of neighborhoods, the changing of these qualitative density states ("low" or "high") in the IBN basin of attraction could be the substrate for estimating CA rules. Given the above we aim (1) to infer a BN from the growth curve time series obtained from *in vitro* co-culture, (2) to estimate the rules of CA dynamics from the IBN basin of attraction by comparing the density states within each state transition and reaching a consensus among state transitions, and (3) to implement a CA to simulate spatial dynamics.

We observed that the binarization of the time series points to a high population density after the fourth day. The IBN functions let us choose a regulator for each node that is supported by the literature, and the CA rules estimated from the IBN basin suggest a condition for cells to migrate or proliferate that is consistent with what has been seen experimentally [1][3]. The migration of keratinocytes increases smoothly until the fourth day of the simulation, making them denser near melanoma clusters and limiting melanoma proliferation, so that the melanoma survival probability curve shape looks like a normal distribution depending on the proliferation capacity. Therefore, the simulation suggests that the attractor state of cell co-culture population dynamics would be induced by an increase in keratinocyte migration and survival, as well as the balance of proliferation and death concerning melanoma.

2. Material and Methods

In this section, we present the methodological approach used for data inference and modeling. We shall highlight that the modeling process is contingent upon the outcomes derived from the inference. Figure 1 illustrates the workflow for data analysis and modeling. The source code link can be found at S5 Source code.

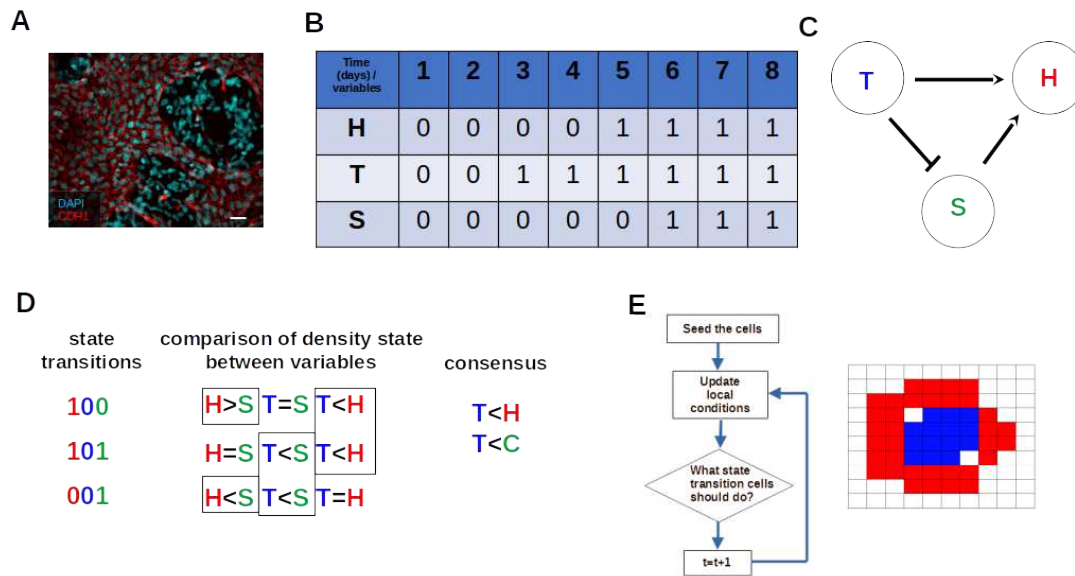


Figure 1. Workflow of data inference and population dynamic modeling. (A) Cell co-culture model of keratinocytes and melanoma cells Scale bar: 50 μm [1]. (B) Population growth time-series binarization. Number 0 indicates a low density of cells (OFF state), and number 1 indicates a high density of cells (ON state). (C) Inferred Boolean Network Topology Arrows indicate activation and blunt arrows indicate inhibition. (D) Estimating cellular automata rules from a state graph or transition table. (E) Implementation of simulation through cellular automata.

2.1. Experimental Data

We use previously published cell proliferation data measured in co-culture [1]. The data are the density (cells/mm^2) of normal human keratinocytes (HaCaT), metastatic melanoma (SK-MEL-147), at co-culture with the initial proportion of 1:10 (SK-MEL-147:HaCaT) [49] measured along 8 days in triplicate. To quantify the cell density necessary to achieve the carrying capacity, we introduced the variable **S** to express the difference between the carrying capacity (**K**) and the population density at each specific time point. This additional variable is justified because the space has a significant impact on the population dynamics. Hence, the more cells need to reach carrying capacity, less space is available, and the fewer cells need to reach carrying capacity, more space is available. Consequently, this variable serves as an initial arithmetic approximation of the space availability. Throughout this paper **T** will represent melanoma, **H** keratinocyte, and **S** space availability.

2.2. Boolean Network Inference

BN inference involves two steps: 1) binarization, which transforms cell counts per day in binary discrete data, and 2) network learning. We evaluated three different binarization methods: (1) *k-means*, (2) *iterative k-means* [50] [44] and (3) *BASC* [51]. The binarization outcome chosen to network learning shall be the one which point highest density after confluence onset in agreement with Morais et al [1]. We used *BoolNet* 2.1.8 from R 4.2 [52] to implement the binarization by *BASC* and *k-means*, and Python 3.10.2 to implement the binarization by *iterative k-means*, adapted from Steinway et al [44]. In the context of this work, the number 0 denotes low density, and the number 1 denotes high density. We performed BN learning using the *BestFit* algorithm [53] through the *BoolNet* package 2.1.8 from R 4.2 [52].

2.3. Estimating the Cellular Automata dynamic rules from BN basins through consensus of state transition comparison

Here propose a simple method to estimate CA rules from BN basin of attraction. The proposed approach involves three steps. First, we compare the density states among variables within all individual transition belonging to a given basin of attraction. Second, we obtain a consensus of all comparison among state transitions. For instance, while analyzing six state transitions, it is observed that a specific variable $\mathbf{A} = 1$ and variable $\mathbf{B} = 0$ ($A \geq B$) in three of them, while $\mathbf{A} = 0$ and $\mathbf{B} = 1$ ($A \leq B$) in one of them, and $\mathbf{A} = 0$ and $\mathbf{B} = 0$ ($A = B$) in two of them. The most observed inequalities are $A \geq B$ and $A = B$, and we define the CA rule as $A \geq B$. We determine absence of regulation between variables when $>$, $=$ and $<$, have the same frequencies such in case.

2.4. Implementation of Square-Lattice Cellular Automata

The Square-Lattice Cellular Automata (SLCA) was built based on the estimation of CA dynamic rules from the BN basin (Figure 1). The model consists of a tissue scale, resulting in a single discrete variable, namely the cells. The model encompasses a dynamic process involving cell proliferation, death, migration, and survival. This process runs on a two-dimensional lattice ($n \times n$ with $n = 200$). The criteria for determining lattice size was based on the spatial model developed by Morais et al [1]. Each vertex can accommodate only one tumor cell. During each iteration, all cells undergo the actions outlined in Figure 5. The cells were randomly distributed over the lattice in a ratio of 1 **T** to 10 **H** according to [1]. We assume a time-step of 12 hours resulting in a total of 16 iteration, corresponding to 8 days-culture, for each simulation. Section 3.3 describes in detail the state transition rules.

2.5. Spatial configuration, parameter estimation, and population dynamics process probabilities

We applied a convolutional Gaussian filter of size $k = 5 \times 5$ and standard deviation σ^2 to estimate the relative local density of **H** cells (arbitrary units) in the simulation domain. To calculate the growth curve of both cell lines we fitted the logistic model (Eq. 1) using the *curve fit* function of the *optimize* module of the Scipy library (version 1.7.1) in Python (version 3.10.2)

$$N(t) = \frac{K}{1 - e^{-\rho(t-\tau)}}, \quad (1)$$

where K denotes carrying capacity, ρ denotes proliferation rates and τ denotes time for half asymptote.

Throughout the simulations, we calculate the probability of the cells going through those tumor growth dynamic processes described in section 2.4. Denote $i = 1, \dots, n$ as an element of the list $\omega =$ (Proliferation, Death, Migration, Survive) and f the frequency of occurrence of one of the processes in the list ω , thus

$$Pr(\omega_i) = \frac{f_{\omega_i}}{\sum_{i=1}^n f_{\omega_i}}, \quad (2)$$

for both **T** and **H**.

3. Results

3.1. The iterative k-means binarization method demonstrated that both populations are in high density after 4 days

As described in Section 2.2, the time series was binarized, and then we learned a BN. The binarization by iterative k-means showed a high density of **H** and **T** and a low availability of **S**, after four days (Figure 1 and in S1 Figure).

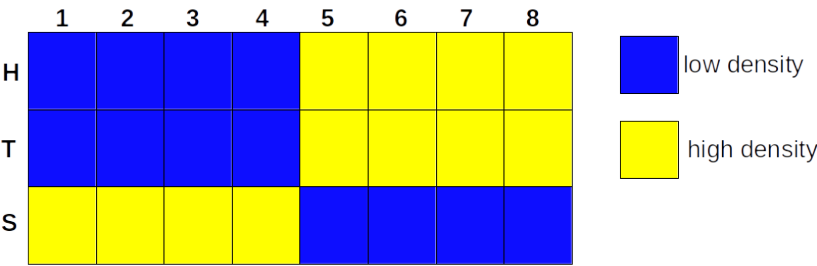


Figure 2. The heatmap represents the binarization of co-culture time series by iterative k-means. The yellow cells represent 1, which means high density, and the blue cells represent 0, which means low density. Each row represents the binarized trajectories of **H**, **T**, and **S**, and each column represents the state transition of the network over eight days.

Table 1 shows the possible Boolean functions for each variable. The possible function for each variable is highlighted in yellow with its justification. The intention of showing two different biologically reasonable networks would be to ascertain possible plasticity in the estimated rules for CA as the BN basin topologies change once a BN has 2^n possible state transitions, here $n=3$ (**H**, **T**, and **S**).

Table 1. Inferred boolean functions through the Best Fit algorithm.

Inferred Boolean Network 1		
Node	Possible functions	Justification
T	<div><div>T</div><div>H</div><div>S</div></div>	Cancer cells have autonomous proliferation [54–57]
H	<div><div>T</div><div>H</div><div>S</div></div>	HaCaT density is higher next to SKMEL-147 and lower far from SKMEL-147 [1]. HaCaT form connections with melanoma cell line rather than adjacent HaCaT [3]
S	<div><div>T</div><div>H</div><div>S</div></div>	Cancer cells do not respond to contact inhibition and other anti-proliferative processes [54–57]
Inferred Boolean Network 2		
Node	Possible functions	Justification
T	<div><div>T</div><div>H</div><div>S</div></div>	Cancer cells have autonomous proliferation [54–57]
H	<div><div>T</div><div>H</div><div>S</div></div>	When no space is available keratinocytes do not increase density
S	<div><div>T</div><div>H</div><div>S</div></div>	Cancer cells do not respond to contact inhibition and other anti-proliferative processes [54–57]

All functions inferred showed non-null error rates. The highlighted functions set the state graph in Figures 3A-3B. The bar over the node symbol denotes the NOT logical operator. All the inferred boolean functions have error rates equal to 1. The highlighted functions set the state graph in Figures 3A-3B. The bar over the node symbol denotes the NOT logical operator. All the inferred boolean functions have error rates equal to 1.

3.2. The network dynamics exhibit two singleton attractors

After we binarized the time series and applied a method of network learning, we generated the state graph, and observed two singletons (fix points) attractors in both IBN 1 (Figure 3A) and 2 (Figure 3B). Before evaluating the attractors and proceeding with the estimation of CA dynamics rules, we labeled each transition state as $B_1, B_2, B_3, B_4, B_5, B_6, A_1, A_2$. These IDs belongs to the same state transition in both IBN 1 and 2. The attractor A_1 shows **T** and **H** in low density, while **S** is in high density, which ultimately means high space availability or the population is scattered. The attractor A_2 shows **H** and **T** in high density and **S** in low density, which ultimately means low space availability or that the population is in high density. The IBN 1 has no predecessor state transition from initial condition among basins of attraction, while the IBN 2 present an small sequence of state transition not.

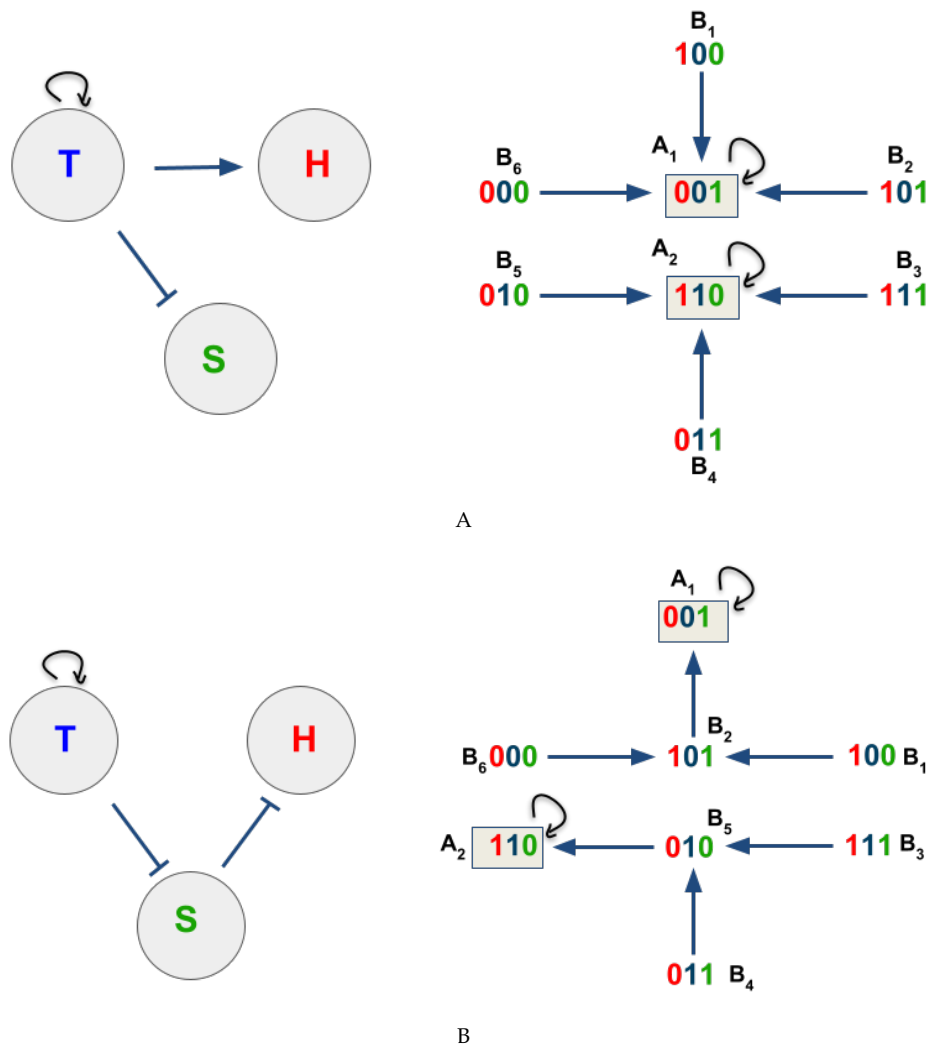


Figure 3. The dynamics of IBN 1 and 2. (A) the topology of the network is shown on the left, and the state graph of IBN 1 on the right. (B) The topology of the network is shown on the left, and the state graph of IBN 2 on the right. A_1 and A_2 denote singleton attractors 1 and 2, respectively, and the letter B denotes state transitions. The red number represents the density state of H, the blue number represents the density state of T, and the green number, the space availability state of S.

3.3. Boolean Network Basin provide cellular automata dynamic rules for proliferation and migration

To estimate the CA dynamic rules from BN basins, we analyze the state transitions in terms of density comparison. We ignore the state transitions that do not have a reasonable biological meaning. B_3 presents a low cell density and low space availability, and B_6 presents a high cell density and high space availability. When we compare the density states between T and S among state transitions B_1 , B_2 , and A_1 , $T < S$ in two-thirds of these state transitions (orange-dash rectangle at Figure 4). Additionally, $T < H$ in 2/3 of the aforementioned state transitions (orange-dash rectangle at Figure 4). Thus, from the consensus of the density state comparisons among these state transitions, we obtained two inequalities: $T \leq S$ and $T \leq H$ (Figure 4). In contrast, upon comparing the density states between T and S among transition states B_4 , B_5 , and A_2 , $T \geq S$ in two-thirds of these state transitions (purple-dash rectangle at Figure 4). Furthermore, $T \geq H$ in 2/3 of those state transitions (purple-dash rectangle at Figure 4), so by consensus among these state transitions, we obtained two new inequalities: $T \geq S$ and $T \geq H$ (Figure 4).

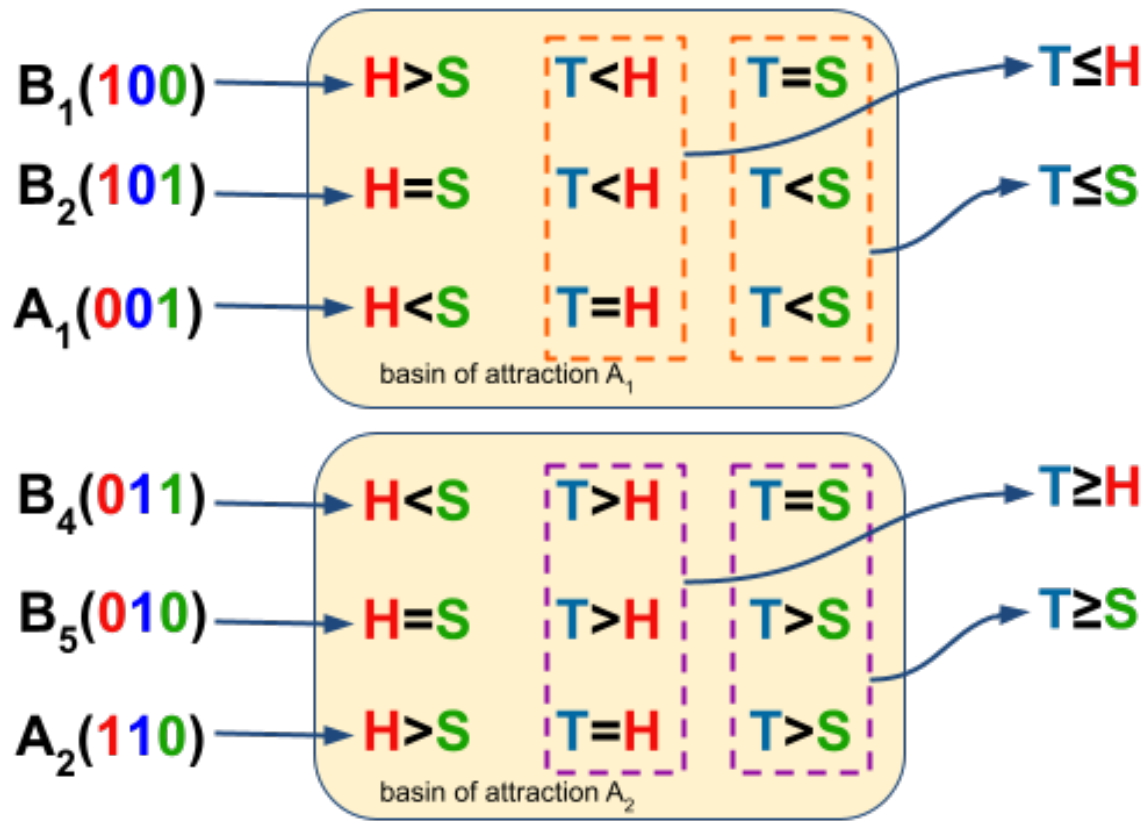


Figure 4. Estimate of the CA dynamic rule from the BN basin. One can observe the comparison within each state transition (B_1, B_2, B_4, B_5, A_1 and A_2) where the number in red indicating the density state of **H**, blue indicating **T**, and green indicating **S**. The dashed rectangles marks the comparisons among state transitions belonging to a basin of attraction, and the arrow points to the comparison consensus among state transitions. The orange-dash rectangle concerns the basin of attraction A_1 , while the purple-dash rectangle to A_2 .

The inequalities $T \leq S$ and $T \leq H$ represent a condition in which there is a low amount of **T** and a high amount of **H** or **S**, which do not favor an increase in density, and according to [3], we determine that it favors migration. Thus, we set eq. 3 where $M_{T,H}$ denotes migration.

$$M_{T,H} = (T \leq S) \vee (T \leq H) \quad (3)$$

The inequalities $T \geq S$ and $T \geq H$ represent a condition where a higher density of **T** and a lower density **H** or **S**, favors density increase. Thus, P_T and P_H denotes proliferation of **T** and **H** in equations 4 and 5, respectively. Due to the lack of data about the resources present in the cell culture media, we decided to augment equations 4 and 5 with the probability of instantaneous proliferation $1 - \frac{N}{K}$ [58], since the logistic model fitted to the growth curves present in Morais et al [1]. If the probability of instantaneous proliferation is satisfied ($1 - \frac{N}{K} \geq r$, where r is a random number) then the cells can proliferate. Another important condition to simulate cell dynamics is cell death, represented by the death term \bar{D}_T in eq. 4 and 5, and the survive term $\bar{S}\bar{V}_H$ that blocks **H** cell proliferation, as explained in section 3.4.

$$P_H = \left(1 - \frac{N}{K} \vee ((T \geq S) \vee (T \geq H))\right) \wedge \bar{S}\bar{V}_H \quad (4)$$

$$P_T = \left(1 - \frac{N}{K} \vee ((T \geq S) \vee (T \geq H))\right) \wedge \bar{D}_T \quad (5)$$

3.4. Data from cell lines and growth curves allow us to propose a simple rule to describe survival and death dynamics

Here, we only could estimate CA rules that are compatible with migration and proliferation. So to describe the survival and death of cells, we turn to Hayflick's limit [59–61] concept to **H** and cancer stem cell hypothesis [62–67] to **T** because both can embrace the so-called parameter proliferation capacity (ρ_{max}). Thus, with each cell division, the proliferation capacity of **T** and **H** will decrease by one unit, $\rho_{max} = \rho_{max} - 1$. It is well documented that **H** undergoes a limited number of duplications in the cell culture plate and does not always reach 100% confluence; thus, once **H** reaches $\rho_{max} = 0$, they survive (SV_H), as proposed in eq. 6. We assume that when **T** reaches $\rho_{max} = 0$ and in circumstances where there is no space (D_T), the crowding situations might yield cell de-adherence and death [68], as in eq. 7.

$$SV_H = \rho_{max} = 0 \quad (6)$$

$$D_T = (\rho_{max} = 0) \wedge (S = 0) \quad (7)$$

Once **H** does not migrate, proliferate, or survive, they die, as in eq. 8. Alternately, once **T** does not migrate, proliferate, and die, they survive, eq. 9.

$$D_H = \overline{M}_H \wedge \overline{P}_H \wedge \overline{SV}_H \quad (8)$$

$$SV_T = \overline{M}_T \wedge \overline{D}_T \wedge \overline{P}_T \quad (9)$$

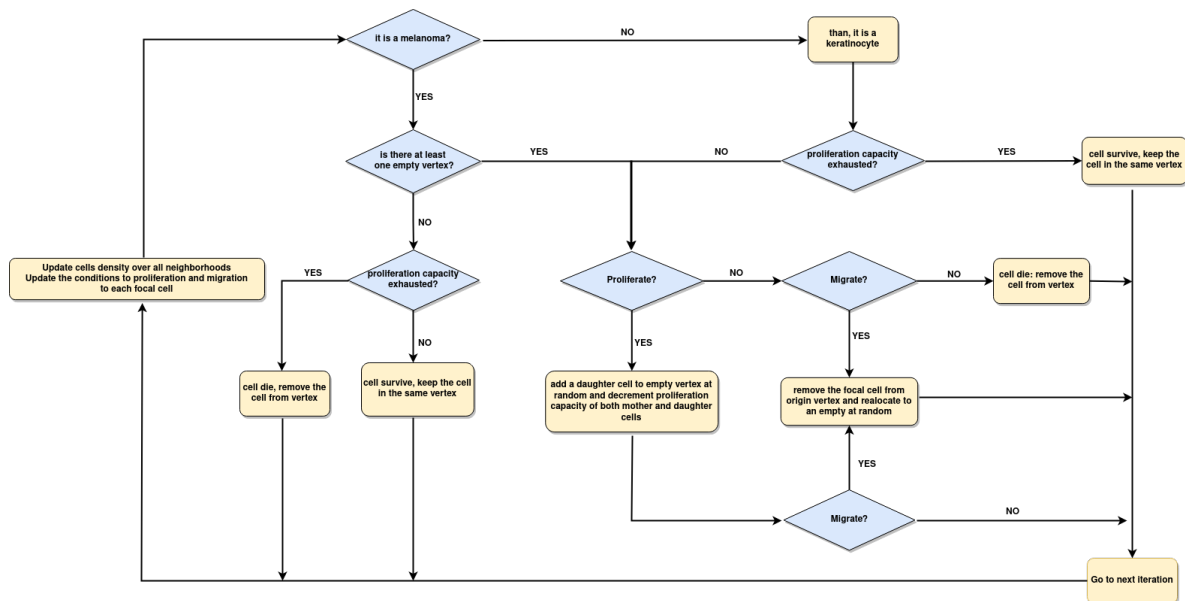


Figure 5. Flowchart of the CA implementation. First, we set the initial density of cells and a random initial distribution of them over Lattice. At each iteration, the algorithm updates the amount of **T**, **H**, and empty vertex (**S**) in all focal cell neighborhoods. The conditions described in eqs. 3 to 9 are updated in each cell. When there is at least one empty vertex, the cells proliferate or migrate, but when the proliferation capacity is exhausted, the cells survive or die.

3.5. The parameter estimation are in agreement with experimental data

The values of cells density and the frequency of proliferating, migrating, surviving, and dying cell in the overall simulation were used to evaluate if the growth curve fit to the logistic model (eq. 1).

The growth curves fit the logistic model (Figure 6G) and the estimated values for the parameter K , ρ , and τ are presented at Table 2. Considering each iteration as a $\Delta t = 12$ hours favored the estimation of $\rho_T \approx \rho_H$ and $\tau_T \approx \tau_H$. We shall recall that the results in Figure 6 concern $\rho_{max_T} = 7$ and $\rho_{max_H} = 6$. According to Morais et al [1] the values estimated to ρ_T and ρ_H are equal as well those estimated to τ_T and τ_H .

Table 2. Table of estimated values for the parameters of the logistic model.

Parameters	Melanoma	Keratinocytes
K	0.16	0.7
rho	1.63	1.20
tau	2.7	2.2

These estimated values for K , ρ , and τ refer to simulations with $N_0 = 2000$, $\rho_{max_T} = 6$, and $\rho_{max_H} = 7$.

3.6. The density of **H** indeed is higher close to **T** cluster

We implemented the spatial model using CA (sections 2.4, 3.3 and 3.4) and, starting from a given an initial distribution of cells, on the fourth day we can already see some **T** clusters surrounded by **H**, so this spatial pattern is consolidated on the eighth day (Figures 6A-6C and animation in S3 Video). The heat map shows **H** density in the neighborhood of each focal cell (Figure 6F) points to higher yellow intensity around **T** clusters. This can be explained by the smooth increase in **H** migration probability from the beginning of the simulation (Figure 6H) as well as the increase in **T** survival probability from day 4 onward (Figure 6G). Therefore, as **T** clusters consolidate, more of the **H** neighborhood lacks **T**; hence the probability of migration decrease mostly because the probability of **H** survive increase substantially after four days (Figure 6H).

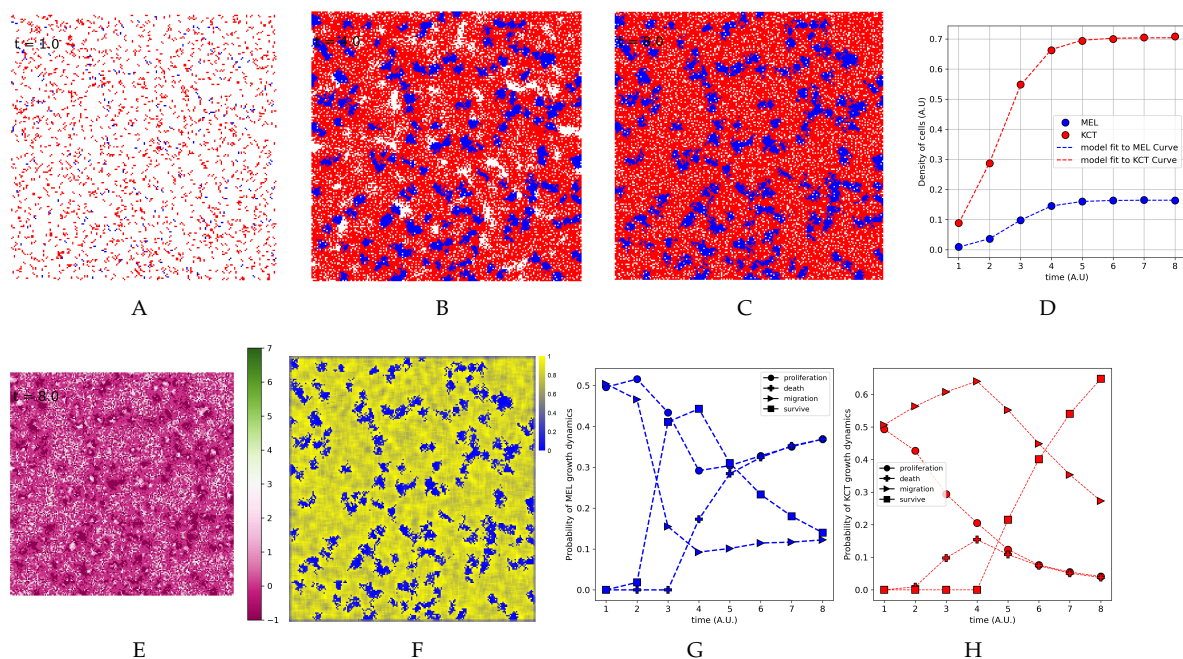


Figure 6. The spatial population and tumor growth dynamics. (A), (B), and (C) frames correspond to days 1, 4, and 8. Blue represents **T**, and red represents **H**. (D) Growth curves of **T** and **H**. (E) Spatial distribution of the cells according to the values of ρ_{max} at day 8. In the bar of color at right, the shades of green to pink represent the ρ_{max} values. (F) Heatmap representing **H** density over all focal cell neighborhoods at day 8. In the bar of color at right, the shades of blue to yellow represent the percentage of occupied vertex by **H** in all focal cell neighborhoods. (G) Probabilities of **T** tumor growth dynamics (H) Probabilities of **H** tumor growth dynamics.

4. Discussion

In this study, we used the time series of the growth curve in the co-culture of **T** and **H** to infer a BN. Through BN inference, we were able to determine the boolean functions that led to network dynamics. These dynamics were shown in a state graph, where we arranged the state transitions according to the basin of attraction and estimated the CA rules. These estimated CA rules were biologically sound; hence, they could represent neighborhood or microenvironmental configurations that drive population dynamics toward the attractor state, as mentioned before.

The binarization of the growth curve data reveals that the population attained a high density after 4 days. This observation aligns with the findings of Morais et al [1], who reported that the population achieves confluence after the same 4-day period. The proposed methodology captured the relationship between the confluence observed on the cell culture plate and the corresponding cell density. A boolean network of n nodes exhibits 2^n state transitions; hence, our time series, consisting of a sequence of 8 data points, can fully define the system. The occurrence of non-null error rates in the inferred boolean network arises due to the similarity of the **H** and **T** binarized trajectories, also known as identifiability; nevertheless, this issue can be solved by prior biological knowledge or by pruning the network nodes according to the context [43].

The BestFit BN learning paradigm searches for possible boolean functions that describe the state of a given variable (node) at time $t + 1$ as a function of the network state (the states of all nodes) at time t . In the heatmap (Figure 2), **S** is ON when both **T** and **H** are OFF, and vice versa, which means that **S** inhibits both **T** and **H**, which in turn inhibits **S**, and finally both **T** and **H** can "activate" each other.

The IBN 1 in Table 1 and Figure 3A shows a canalizing node and a initial condition with no predecessor state transition throughout basin of attraction. The IBN 2 in Table 1 and Figure 3B, on the other hand, shows a small number of states before reaching the attractor. However, both networks (Figures 3A and 3B) have the same attractor. Hence, different BNs can lead to the same attractors. This indicates that the estimation of CA rules was not affected by changes in the topology of the state graph, suggesting a system plastic capacity, as in the genetic networks [69].

Indeed melanoma affects keratinocytes phenotypes [2] [3], our approach only captured an equilibrium state concerning local density of both melanoma and keratinocyte. By comparing the density states of state transitions B_4 , B_5 , and A_2 , we could estimate the logical sentence $(T \geq S) \vee (T \geq H)$, suggesting that **T** will only proliferate where there is more **T**, since experiments with model organisms like zebrafish have shown that melanoma tends to cluster [70]. Likewise, we could estimate the logical sentence $(T \leq S) \vee (T \leq H)$ based on how the density states of states transition B_1 , B_2 , and A_1 , meaning that either **H** or **T** will move away [1] [3]. It is worth highlighting that these Boolean expressions can be concurrently satisfied in a dynamic context. So, the potential for cell proliferation and migration is inherently incorporated, which reduces any challenges in modeling associated with the go-or-grow phenomenon [71–75].

The spatial dynamic shows **H** more denser nearby **T** clusters than close to themselves (Figures 6A and 6B); moreover, the values estimated for $\rho_{T,H}$ and $\tau_{T,H}$ are very similar, corroborating the results of Morais et al [1] despite the dependence of estimated $\rho_{T,H}$ on the initial condition. The growth curve certainly did not emerge from the dynamics itself, but it is likely hardwired to the instantaneous probability term present in eqs. 5 and 6 that was employed to represent the resources in the culture medium.

We could only estimate CA rules for proliferation and migration, then adopt an intrinsic condition for cells to survive and die. Normal cells like **H** undergo a limited number of duplications in the cell culture plate and do not always reach 100% confluence because at each duplication the telomeres shorten, yielding to the onset of replicative senescence [59–61]. In the case of cancer cells, i.e., **T**, the telomeres shorten with doublings, and the cells will enter a state of crisis or death. Both normal and cancer cells could die whenever there is no space, because *in vitro* crowding situations lead some cells to undergo de-adherence and die [68]. Moreover, crowding-induced elimination would facilitate the

expansion of the malignant clones [76–79]. Due to the duration of the *in vitro* experiment (see Section 2.1), for simplification, we discarded stem cells with $\rho_{max} = \infty$ in the model.

The proliferation capacity parameter (ρ_{max}) as a measure of T death shows that cells with a lower ρ_{max} value tend to be at the edges of the clusters, competing for space with cells with a higher ρ_{max} value (Figure 6E). As soon as the cells on the periphery die, the others proliferate and thus decrease their proliferation capacity. In this respect, the ignition of stem cells by space release and/or their capacity to proliferate more than those with no stemness are in agreement with previously published theoretical models [62–67], experimental evidence [80], and the amalgamation of experimental data with mathematical modeling [81] [82].

Probably, if we had implemented a Lattice Gas Cellular Automata (LGCA), by considering that proliferating T cells could pile up due to the lack of contact inhibition, T proliferation would naturally be counterbalanced by their death because of mass action in the growth dynamics. Even though we used a simple SLCA, which doesn't let us include more than one cell per lattice vertex, this makes it harder to include the lack of contact inhibition, which is typical of cancer cells. However, even if cancer cells present a low contact inhibition degree, mechanical factors could restrict their proliferation [83]. The approach using proliferation capacity as a simpler rule to distinguish death from survival allowed us to observe a decrease in the $\frac{H}{T}$ density ratio, which we hypothesize to be due to death-induced proliferation (Figure 6G). It is biologically sound because pathways involved in contact inhibition might go through a transition towards the regulation of autophagy, which in turn can yield survival and proliferation [84].

Our results suggest that the inference of BN from the co-culture growth curve time series provides a path to estimate specific microenvironmental conditions for proliferation and migration in terms of CA dynamic rules, despite the tumor microenvironment being considered a complex regulatory network that shapes tumor architecture and growth law [57,85–88]. Even though the spatial population dynamics model was made using a mix of estimated data from IBN dynamics and observations from the literature, the results are in line with experimental data. Hitherto, we simulate local dynamics by running the CA forward for space-time patterns and this was enough to reach our goals, but as future perspective our approach could be coupled with the software DDLAB (www.ddlab.org) in order to develop a pipeline to evaluate the dynamic upon or on the network [9,10].

Since either model-driven experimentation or data-driven modeling are important ways to improve and integrate mathematical, biological, and clinical models, unraveling CA rules from BN dynamics could be useful to capture any interaction between cells in terms of density patterns and it deserve room for further refinement. This is mainly because a discrete approach can be linked with a continuous one [89,90], and important parameters beyond the local or global dynamics can be estimated. Cell co-culture is an experimental system that can comprise diverse time points, images of spatial features, and different cell types and densities. This makes it much easier to figure out how networks work at the tissue scale, especially when it comes to figuring out how to reconstruct population dynamics.

5. Conclusion

In this work, we introduce a novel approach to cell proliferation dynamics. We present a approach to estimate the rules of the dynamics of a cellular automaton from the basins of attraction of a Boolean network and, also, align them with experimental conditions. By establishing a direct connection between our theoretical model and experimentally observed spatial patterns, we attain a significant milestone in the field of cellular analysis by effectively connecting computation and experimental methods. This modeling approach can make a significant advancement in our ability to comprehend and predict complex biological phenomena.

Supplementary Materials: The following supporting information can be downloaded at the website of this paper posted on Preprints.org

Acknowledgments: We would like to thank NPAD/UFRN for computational resources. MCCM is supported by São Paulo State Research Foundation (FAPESP process n. 2020/12017-9). ASQ thanks Prof. Dr. Roger Chammas, who read the very first version of the article and assured him that the idea was coherent. The authors are thankful for anonymous reviewers whose comments enhanced manuscript's quality.

References

1. Morais MCC, Stuhl I, Sabino AU, Lautenschlager WW, Queiroga AS, Tortelli Jr TC, et al. Stochastic model of contact inhibition and the proliferation of melanoma in situ. *Scientific reports*. 2017;7(1):8026.
2. Kodet O, Lacina L, Krejčí E, Dvořánková B, Grim M, Štork J, et al. Melanoma cells influence the differentiation pattern of human epidermal keratinocytes. *Molecular cancer*. 2015;14:1–12.
3. Mazurkiewicz J, Simiczyjew A, Dratkiewicz E, Kot M, Pietraszek-Gremplewicz K, Wilk D, et al. Melanoma stimulates the proteolytic activity of HaCaT keratinocytes. *Cell communication and signaling*. 2022;20(1):1–17.
4. Neumann JV. *Theory of self-reproducing automata*. Edited by Arthur W Burks. 1966;.
5. Wolfram S. *Statistical mechanics of cellular automata*. *Reviews of modern physics*. 1983;55(3):601.
6. Wolfram S. *A new kind of science*. vol. 5. Wolfram media Champaign, IL; 2002.
7. Kauffman S. Homeostasis and differentiation in random genetic control networks. *Nature*. 1969;224(5215):177–178.
8. Bornholdt S. Boolean network models of cellular regulation: prospects and limitations. *Journal of the royal society interface*. 2008;5(suppl_1):S85–S94.
9. Wuensche A, Lesser M. *Global dynamics of cellular automata: An atlas of basin of attraction fields of one-dimensional cellular automata*. vol. 1. Colorado: Westview Press; 1992.
10. Wuensche A. *Exploring discrete dynamics*. Luniver Press; 2011.
11. Wuensche A. *Attractor basins of discrete networks*. *Cognitive science research paper*. 1997;461.
12. Wuensche A. *The Ghost in the Machine: Basin of Attraction Fields of Disordered Cellular Automata Networks*. In: *Artificial Life III Conference*; 1992.
13. Wuensche A. *The ghost in the machine: Basins of attraction of random boolean networks*. *Artificial Life III: SFI Studies in the Sciences of Complexity*, vol. VII Addison-Wesley. 1994;.
14. Jeras I, Dobnikar A. Algorithms for computing preimages of cellular automata configurations. *Physica D: nonlinear phenomena*. 2007;233(2):95–111.
15. Soto JMG. Computation of explicit preimages in one-dimensional cellular automata applying the De Bruijn diagram. *Journal of cellular automata*. 2008;3(3):219–230.
16. Dormann S, Deutsch A. Modeling of self-organized avascular tumor growth with a hybrid cellular automaton. *In silico biology*. 2002;2(3):393–406.
17. Spencer SL, Gerety RA, Pienta KJ, Forrest S. Modeling somatic evolution in tumorigenesis. *PLoS computational biology*. 2006;2(8):e108.
18. Enderling H, Hlatky L, Hahnfeldt P. Migration rules: tumours are conglomerates of self-metastases. *British journal of cancer*. 2009;100(12):1917–1925.
19. Enderling H, Anderson AR, Chaplain MA, Beheshti A, Hlatky L, Hahnfeldt P. Paradoxical dependencies of tumor dormancy and progression on basic cell kinetics. *Cancer research*. 2009;69(22):8814–8821.
20. Kansal A, Torquato S, Chiocca E, Deisboeck T. Emergence of a subpopulation in a computational model of tumor growth. *Journal of theoretical biology*. 2000;207(3):431–441.
21. Zhang L, Strouthos CG, Wang Z, Deisboeck TS. Simulating brain tumor heterogeneity with a multiscale agent-based model: linking molecular signatures, phenotypes and expansion rate. *Mathematical and computer modelling*. 2009;49(1-2):307–319.
22. Wcislo R, Dzwinel W, Yuen DA, Dudek AZ. A 3-D model of tumor progression based on complex automata driven by particle dynamics. *Journal of molecular modeling*. 2009;15:1517–1539.
23. Wang Z, Deisboeck TS. Computational modeling of brain tumors: discrete, continuum or hybrid? *Scientific modeling and simulations*. 2009; p. 381–393.
24. Anderson AR, Rejniak KA, Gerlee P, Quaranta V. Microenvironment driven invasion: a multiscale multimodel investigation. *Journal of mathematical biology*. 2009;58:579–624.
25. Zhang L, Wang Z, Sagotsky JA, Deisboeck TS. Multiscale agent-based cancer modeling. *Journal of mathematical biology*. 2009;58:545–559.

26. Anderson AR. A hybrid mathematical model of solid tumour invasion: the importance of cell adhesion. *Mathematical medicine and biology: a journal of the IMA*. 2005;22(2):163–186.
27. Sottoriva A, Verhoeff JJ, Borovski T, McWeeney SK, Naumov L, Medema JP, et al. Cancer stem cell tumor model reveals invasive morphology and increased phenotypical heterogeneity. *Cancer research*. 2010;70(1):46–56.
28. Gatenby RA, Smallbone K, Maini PK, Rose F, Averill J, Nagle RB, et al. Cellular adaptations to hypoxia and acidosis during somatic evolution of breast cancer. *British journal of cancer*. 2007;97(5):646–653.
29. Gerlee P, Anderson A. Evolution of cell motility in an individual-based model of tumour growth. *Journal of theoretical biology*. 2009;259(1):67–83.
30. Silva AS, Yunes JA, Gillies RJ, Gatenby RA. The potential role of systemic buffers in reducing intratumoral extracellular pH and acid-mediated invasion. *Cancer research*. 2009;69(6):2677–2684.
31. Zhang L, Chen LL, Deisboeck TS. Multi-scale, multi-resolution brain cancer modeling. *Mathematics and computers in simulation*. 2009;79(7):2021–2035.
32. Gerlee P, Anderson AR. A hybrid cellular automaton model of clonal evolution in cancer: the emergence of the glycolytic phenotype. *Journal of theoretical biology*. 2008;250(4):705–722.
33. Smallbone K, Gatenby RA, Gillies RJ, Maini PK, Gavanagh DJ. Metabolic changes during carcinogenesis: potential impact on invasiveness. *Journal of theoretical biology*. 2007;244(4):703–713.
34. Anderson AR, Hassanein M, Branch KM, Lu J, Lobdell NA, Maier J, et al. Microenvironmental independence associated with tumor progression. *Cancer research*. 2009;69(22):8797–8806.
35. Anderson AR, Weaver AM, Cummings PT, Quaranta V. Tumor morphology and phenotypic evolution driven by selective pressure from the microenvironment. *Cell*. 2006;127(5):905–915.
36. Basanta D, Strand DW, Lukner RB, Franco OE, Cliffl DE, Ayala GE, et al. The role of transforming growth factor- β -mediated tumor-stroma interactions in prostate cancer progression: An integrative approach. *Cancer research*. 2009;69(17):7111–7120.
37. de Pillis LG, Mallet DG, Radunskaya AE. Spatial tumor-immune modeling. *Computational and mathematical methods in medicine*. 2006;7(2-3):159–176.
38. Enderling H, Alexander NR, Clark ES, Branch KM, Estrada L, Crooke C, et al. Dependence of invadopodia function on collagen fiber spacing and cross-linking: computational modeling and experimental evidence. *Biophysical journal*. 2008;95(5):2203–2218.
39. Bankhead A, Magnuson NS, Heckendorn RB. Cellular automaton simulation examining progenitor hierarchy structure effects on mammary ductal carcinoma in situ. *Journal of theoretical biology*. 2007;246(3):491–498.
40. Gevertz J, Torquato S. Growing heterogeneous tumors in silico. *Physical review E*. 2009;80(5):51–91.
41. Anderson AR, Basanta D, Gerlee P, Rejniak KA. Evolution, regulation, and disruption of homeostasis, and its role in carcinogenesis. *Multiscale cancer modeling*. 2011; p. 1–30.
42. Silva AS, Gatenby RA, Gillies RJ, Yunes JA. A quantitative theoretical model for the development of malignancy in ductal carcinoma in situ. *Journal of theoretical biology*. 2010;262(4):601–613.
43. Putnins M, Campagne O, Mager D, Androulakis I. From data to QSP models: a pipeline for using Boolean networks for hypothesis inference and dynamic model building. *Journal of pharmacokinetics and pharmacodynamics*. 2022; p. 1–15.
44. Steinway SN, Biggs MB, Loughran Jr TP, Papin JA, Albert R. Inference of network dynamics and metabolic interactions in the gut microbiome. *PLoS computational biology*. 2015;11(6):e1004338.
45. Schlatter R, Schmic K, Avalos Vizcarra I, Scheurich P, Sauter T, Borner C, et al. ON/OFF and beyond-a Boolean model of apoptosis. *PLoS computational biology*. 2009;5(12):e1000595.
46. Mai Z, Liu H. Boolean network-based analysis of the apoptosis network: irreversible apoptosis and stable surviving. *Journal of theoretical biology*. 2009;259(4):760–769.
47. Steinway SN, Zañudo JG, Ding W, Rountree CB, Feith DJ, Loughran Jr TP, et al. Network modeling of TGF β signaling in hepatocellular carcinoma epithelial-to-mesenchymal transition reveals joint sonic hedgehog and Wnt pathway activation. *Cancer research*. 2014;74(21):5963–5977.
48. Fumia HF, Martins ML. Boolean network model for cancer pathways: predicting carcinogenesis and targeted therapy outcomes. *PloS one*. 2013;8(7):e69008.
49. Cichorek M, Wachulska M, Stasiewicz A, Tymińska A. Skin melanocytes: biology and development. *Advances in dermatology and allergology*. 2013;30(1):30–41.

50. Berestovsky N, Nakhleh L. An evaluation of methods for inferring boolean networks from time-series data. *PloS one*. 2013;8(6):e66031.
51. Hopfensitz M, Mussel C, Wawra C, Maucher M, Kuhl M, Neumann H, et al. Multiscale binarization of gene expression data for reconstructing Boolean networks. *IEEE/ACM transactions on computational biology and bioinformatics*. 2011;9(2):487–498.
52. Müssel C, Hopfensitz M, Kestler HA. BoolNet—an R package for generation, reconstruction and analysis of Boolean networks. *Bioinformatics*. 2010;26(10):1378–1380.
53. Lähdesmäki H, Shmulevich I, Yli-Harja O. On learning gene regulatory networks under the Boolean network model. *Machine learning*. 2003;52:147–167.
54. Hanahan D, Weinberg RA. The hallmarks of cancer. *Cell*. 2000;100(1):57–70.
55. Hanahan D, Weinberg RA. Hallmarks of cancer: the next generation. *Cell*. 2011;144(5):646–674.
56. Weinberg RA. *The biology of cancer*. Garland science, 2013
57. Hanahan D. Hallmarks of cancer: new dimensions. *Cancer discovery*. 2022;12(1):31–46.
58. Benzekry S, Lamont C, Beheshti A, Tracz A, Ebos JM, Hlatky L, et al. Classical mathematical models for description and prediction of experimental tumor growth. *PLoS computational biology*. 2014;10(8):e1003800.
59. Hayflick L, Moorhead PS. The serial cultivation of human diploid cell strains. *Experimental cell research*. 1961;25(3):585–621.
60. Hayflick L. The limited in vitro lifetime of human diploid cell strains. *Experimental cell research*. 1965;37(3):614–636.
61. Blackburn EH, Gall JG. A tandemly repeated sequence at the termini of the extrachromosomal ribosomal RNA genes in Tetrahymena. *Journal of molecular biology*. 1978;120(1):33–53.
62. Poleszczuk J, Hahnfeldt P, Enderling H. Biphasic modulation of cancer stem cell-driven solid tumour dynamics in response to reactivated replicative senescence. *Cell proliferation*. 2014;47(3):267–276.
63. Poleszczuk J, Enderling H. A high-performance cellular automaton model of tumor growth with dynamically growing domains. *Applied mathematics*. 2014;5(1):144.
64. Poleszczuk J, Hahnfeldt P, Enderling H. Evolution and phenotypic selection of cancer stem cells. *PLoS computational biology*. 2015;11(3):e1004025.
65. Poleszczuk J, Enderling H. Cancer stem cell plasticity as tumor growth promoter and catalyst of population collapse. *Stem cells international*. 2016;2016.
66. Morton CI, Hlatky L, Hahnfeldt P, Enderling H. Non-stem cancer cell kinetics modulate solid tumor progression. *Theoretical biology and medical modelling*. 2011;8:1–13.
67. Shyntar A, Patel A, Rhodes M, Enderling H, Hillen T. The Tumor Invasion Paradox in Cancer Stem Cell-Driven Solid Tumors. *Bulletin of mathematical biology*. 2022;84(12):139.
68. Topman G, Sharabani-Yosef O, Gefen A. A method for quick, low-cost automated confluency measurements. *Microscopy and microanalysis*. 2011;17(6):915–922.
69. Martin S, Zhang Z, Martino A, Faulon JL. Boolean dynamics of genetic regulatory networks inferred from microarray time series data. *Bioinformatics*. 2007;23(7):866–874.
70. Campbell NR, Rao A, Hunter MV, Sznurkowska MK, Briker L, Zhang M, et al. Cooperation between melanoma cell states promotes metastasis through heterotypic cluster formation. *Developmental cell*. 2021;56(20):2808–2825.
71. Hatzikirou H, Basanta D, Simon M, Schaller K, Deutsch A. ‘Go or grow’: the key to the emergence of invasion in tumour progression? *Mathematical medicine and biology: a journal of the IMA*. 2012;29(1):49–65.
72. Gallaher J, Anderson AR. Evolution of intratumoral phenotypic heterogeneity: the role of trait inheritance. *Interface focus*. 2013;3(4):20130016.
73. Kimmel GJ, Dane M, Heiser LM, Altrock PM, Andor N. Integrating mathematical modeling with high-throughput imaging explains how polyploid populations behave in nutrient-sparse environments. *Cancer research*. 2020;80(22):5109–5120.
74. Garay T, Juhász É, Molnár E, Eisenbauer M, Czírók A, Dekan B, et al. Cell migration or cytokinesis and proliferation?—revisiting the “go or grow” hypothesis in cancer cells in vitro. *Experimental cell research*. 2013;319(20):3094–3103.
75. Vittadello ST, McCue SW, Gunasingh G, Haass NK, Simpson MJ. Examining go-or-grow using fluorescent cell-cycle indicators and cell-cycle-inhibiting drugs. *Biophysical journal*. 2020;118(6):1243–1247.

76. Kon S, Ishibashi K, Katoh H, Kitamoto S, Shirai T, Tanaka S, et al. Cell competition with normal epithelial cells promotes apical extrusion of transformed cells through metabolic changes. *Nature cell biology*. 2017;19(5):530–541.
77. Hill W, Zaragkoulias A, Salvador-Barbero B, Parfitt GJ, Alatsatianos M, Padilha A, et al. EPHA2-dependent outcompetition of KRASG12D mutant cells by wild-type neighbors in the adult pancreas. *Current biology*. 2021;31(12):2550–2560.
78. Martins VC, Busch K, Juraeva D, Blum C, Ludwig C, Rasche V, et al. Cell competition is a tumour suppressor mechanism in the thymus. *Nature*. 2014;509(7501):465–470.
79. Wagstaff L, Goschorska M, Kozyrska K, Duclos G, Kucinski I, Chessel A, et al. Mechanical cell competition kills cells via induction of lethal p53 levels. *Nature communications*. 2016;7(1):11373.
80. Alowaidi F, Hashimi SM, Alqurashi N, Alhulais R, Ivanovski S, Bellette B, et al. Assessing stemness and proliferation properties of the newly established colon cancer ‘stem’ cell line, CSC480 and novel approaches to identify dormant cancer cells. *Oncology reports*. 2018;39(6):2881–2891.
81. Gao X, McDonald JT, Hlatky L, Enderling H. Acute and fractionated irradiation differentially modulate glioma stem cell division kinetics. *Cancer research*. 2013;73(5):1481–1490.
82. Tang J, Fernandez-Garcia I, Vijayakumar S, Martinez-Ruis H, Illa-Bohaca I, Nguyen DH, et al. Irradiation of juvenile, but not adult, mammary gland increases stem cell self-renewal and estrogen receptor negative tumors. *Stem cells*. 2014;32(3):649–661.
83. Grimes DR, Fletcher AG. Close encounters of the cell kind: The impact of contact inhibition on tumour growth and cancer models. *Bulletin of mathematical biology*. 2020;82:1–13.
84. Pavel M, Renna M, Park SJ, Menzies FM, Ricketts T, Füllgrabe J, et al. Contact inhibition controls cell survival and proliferation via YAP/TAZ-autophagy axis. *Nature communications*. 2018;9(1):2961.
85. West J, Newton PK. Cellular interactions constrain tumor growth. *Proceedings of the national academy of sciences*. 2019;116(6):1918–1923.
86. Hanahan D, Coussens LM. Accessories to the crime: functions of cells recruited to the tumor microenvironment. *Cancer cell*. 2012;21(3):309–322.
87. Quail DF, Joyce JA. Microenvironmental regulation of tumor progression and metastasis. *Nature medicine*. 2013;19(11):1423–1437.
88. Maley CC, Aktipis A, Graham TA, Sottoriva A, Boddy AM, Janiszewska M, et al. Classifying the evolutionary and ecological features of neoplasms. *Nature reviews cancer*. 2017;17(10):605–619.
89. Wittmann DM, Krumsiek J, Saez-Rodriguez J, Lauffenburger DA, Klamt S, Theis FJ. Transforming Boolean models to continuous models: methodology and application to T-cell receptor signaling. *BMC systems biology*. 2009;3(1):1–21.
90. Krumsiek J, Pölsterl S, Wittmann DM, Theis FJ. Odepy—from discrete to continuous models. *BMC bioinformatics*. 2010;11:1–10.

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