

Optimal structure of heterogeneous stem cell niche: The importance of cell migration in delaying tumorigenesis

Leili Shahriyari^{1,‡} and *Ali Mahdipour–Shirayeh*²

¹ *Mathematical Biosciences Institute, The Ohio State University, OH, USA*

² *Biomedical Research Group, Applied Mathematics Department, University of Waterloo, ON N2L 3G1, Canada*

[‡] *shahriyari.1@mbi.osu.edu*

Abstract

Studying the stem cell niche architecture is a crucial step for investigating the process of oncogenesis and obtaining an effective stem cell therapy for various cancers. Recently, it has been observed that there are two groups of stem cells in the stem cell niche collaborating with each other to maintain tissue homeostasis. One group comprises the border stem cells, which is responsible to control the number of non-stem cells as well as stem cells. The other group, central stem cells, regulates the stem cell niche. In the present study, we develop a bi-compartmental stochastic model for the stem cell niche to study the spread of mutants within the niche. The analytic calculations and numeric simulations, which are in perfect agreement, reveal that in order to delay the spread of mutants in the stem cell niche, a small but non-zero number of stem cell proliferations must occur in the central stem cell compartment. Moreover, the migration of border stem cells to the central stem cell compartment delays the spread of mutants. Furthermore, the fixation probability of mutants in the stem cell niche is independent of types of stem cell division as long as all stem cells do not divide fully asymmetrically. Additionally, the progeny of central stem cells have a much higher chance than the progeny of border stem cells to take over the entire niche.

Introduction

Many studies have been designed to investigate the stem cell dynamics because of their crucial role in many diseases such as cancer. Recently, there have been several studies about employing stem cell therapy for various diseases including cancers [1]. In an in vivo study, it has been observed that the injection of rat umbilical cord stem cells (rUSCs) can completely abolish rat mammary carcinomas with no evidence of metastasis or recurrence hundred days post-tumor cell inoculation [2].

Knowing stem cell dynamics such as their division and death rates, their types of divisions, and the rate of each type of division can assist us to improve stem cell therapies. This knowledge can suggest ways of altering the structure of the stem cell niche to minimize the number of mutant stem cells and control the

cells' growth rate. Understanding the division patterns of stem cells in healthy and malignant tissues can also help us to determine the origins of several diseases like cancer.

Tissue cells are divided in two general categories: Stem cells and non-stem cells, which include transit amplifying (TA) and fully differentiated (FD) cells. The main characteristic of stem cells is their diverse division types. There are two types of stem cell divisions: symmetric and asymmetric. When a stem cell divides asymmetrically, one of the newborn cells is a stem cell and the other one is a non-stem cell. There are two types of stem cell symmetric division: proliferation (both newborn cells are stem cells) and differentiation (both children are TA cells). It has been shown that normal adult human stem cells divide mostly symmetrically in the intestinal crypts [3]. There is also evidence showing that stem cells divide asymmetrically in the response to the injury during tissue regeneration [4]. Since the asymmetric stem cell division leads to a high probability of two-hit mutant productions [5], injuries might increase the chance of tumorigenesis.

There are many experimental observations about the cooperation among different cell types within and around the niche. For instance, in the intestinal crypts, *Lgr5* stem cells receive niche support from Paneth cells, which are located at the bottom of the crypts in the stem cell niche (SC niche) [6]. The cooperation of epithelial hair follicle stem cells (HFSCs) and melanocyte stem cells (MCSCs) in the bulge niche leads to the differentiation and proliferation of MCSCs [7]. There is evidence showing that the haematopoietic stem cell (HSC) niche in the bone marrow is protected and regulated by the HSC progeny and mesenchymal stem cells [8–10]. Additionally, two distinct groups of dormant (d-HSCs) and activated (a-HSCs) stem cells have been identified [11–14].

It has been suggested that stem cells in many tissues are characterized by a bi-compartmental organization [15]. One of the compartments is responsible to regenerate the tissue, while the other compartment controls the SC niche. Such patterns have been observed in several adult SC niches, such as those in hair follicles, blood, intestine, and brain [15]. More details of the stem cell dynamics in the bi-compartmental niche structure have been recently reported by Ritsma et al. [16]. They observed that there are two groups of stem cells in the SC niche: border stem cells (BSCs) and central stem cells (CeSCs). Border stem cells, which are located between central stem cells and transit amplifying cells, mostly differentiate to control the number of non-stem cells. The central stem cells, which are located at the middle of stem cell niche surrounded by BSCs, mostly proliferate to control the number of stem cells [16].

Mathematical models have been developed to understand cell dynamics in both normal and malignant tissue [17–21]. Many of these models investigate the dynamics of stem cells [22–28]. Some of the studies are concentrated on mutations in stem cells, because of their importance in the initiation and progressions of cancers, see e.g. [29–34]. Shahriyari and Komarova [35] developed a stochastic bi-compartmental model for the SC niche to obtain the optimal structure of the SC niche, which delays two-hit mutants' production. They found that the optimal structure corresponds to the stem cell symmetric divisions and most stem cell divisions occurring in BSCs (very small but not zero number of division happening in the CeSCs). Recently, we have developed a 4-compartmental stochastic model, which includes two stem cell groups, to study the movement of mutants in the intestinal and colon crypts [36].

Ritsma et al. [16] observed a small number of migrations from BSCs to CeSCs. However, the model developed in [35] does not consider the possibility of the migration from BSCs towards the CeSC compartment. In this work, by generalizing that model, adding the possibility of migration from BSCs to CeSCs, we obtain the optimal niche's architecture that delays the spread of mutants in the niche. Our model reveals

that the migration from BSCs to CeSCs delays the spread of mutants. The model also shows that the percentage of symmetric and asymmetric divisions does not make a difference in the probability of mutants taking over the entire SC niche as long as all stem cell divisions are not fully asymmetric. The number of mutants does not change if all stem cell divisions are asymmetric, because when a mutant stem cell divides asymmetrically it renews itself and it produces a mutant non-stem cell. Furthermore, the model indicates that it is unlikely that the progeny of mutant BSCs take over the entire niche.

Results

Model set-up

We develop a Moran model for cell dynamics in the SC niche by generalizing the model introduced in [35]. The model contains two stem cell compartments: the BSC compartment (S_b) and the CeSC compartment (S_c). In this model, we only keep track of stem cells and denote the population size of CeSCs and BSCs respectively by $|S_c|$ and $|S_b|$.

In the Moran cell dynamics models, at each updating time step one cell dies and one cell divides to keep the number of cells constant. Here, we assume two non-stem cells die and they are replaced by two stem cells. The missing non-stem cells could be replaced by the divisions of non-stem cells. Since we are modeling the stem cell dynamics, we only consider those that are replaced by stem cell divisions. In other words, we assume at each updating time step, two non-stem cells are required to be produced by some stem cell divisions. We choose two cell divisions only to accommodate stem cells' symmetric divisions and keep the number of cells at each compartment constant, because we are constructing a Moran model to obtain fixation probabilities. In this model, at each updating time step two cells are randomly chosen to divide

Tab. 1: Model parameters.

Symbol	Description
r	Prolif. rate of malignant w.t. and stem cells
σ	Prob. of symmetric division in CeSCs
γ	Prob. of proliferation in CeSC
α	Prob. of migration from BSC to CeSc when a BSC proliferates
$ S_c $	Total number of CeSCs
$ S_b $	Total number of BSCs

based on their fitness. We assume that the normalized fitness of wild-type cells is 1, while the relative fitness of mutants is r . We consider a range of values for the mutants' fitness: $r < 1$ (disadvantageous mutants), $r > 1$ (advantageous mutants), and $r = 1$ (neutral mutants). For instance, if a division occurs in the BSC compartment, with probability $\frac{rb^*}{rb^*+b}$ one BSC mutant divides, and with probability $\frac{b}{rb^*+b}$ one normal cell divides, where b and b^* are respectively the number of normal and mutant BSCs. At each updating time, two stem cell divisions happen based on the following algorithm (See Fig 1).

- With probability $1 - \sigma$, two BSCs divide asymmetrically.
- Or, with probability σ , one BSC differentiates to produce two non-stem cells.

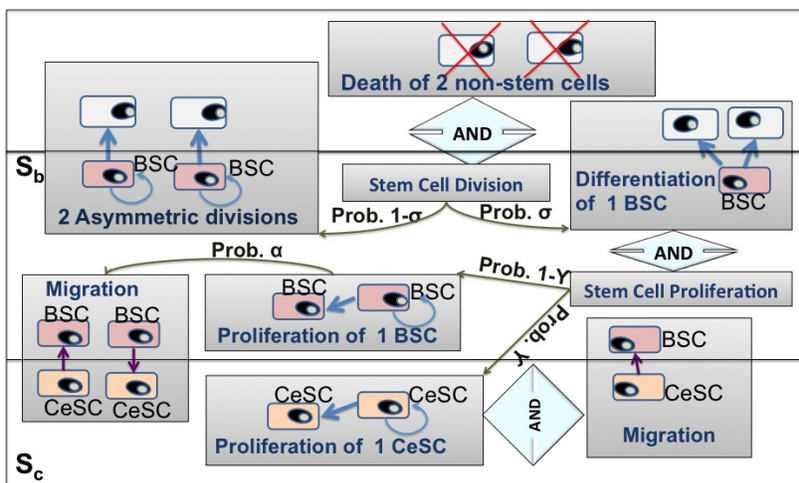


Fig. 1: Schematic representation of the model. When two non-stem cells die, either two BSCs divide asymmetrically or one BSC differentiates to generate two non-stem cells. When a BSC cell differentiates, a proliferation occurs in the SC niche to replace the differentiated BSC cell.

- And, with probability γ , one CeSC proliferates. Then, one random cell from CeSc migrates to the S_b compartment, to replace the differentiated BSC.
- Or, with probability $1 - \gamma$, one border stem cell proliferates to replace the differentiated BSC cell.
 - * And, with probability α , one random cell from the S_b compartment migrates to the S_c compartment. Then, one random cell from the S_c group migrates to the S_b compartment, to keep the number of cells at each compartment constant.

The parameters introduced in the above algorithm are given in Table 1. Also, the general mechanism, which is illustrated in the algorithm, can be seen in Figure 1, in which the different steps of the procedure are given in detail.

The progeny of mutant CeSCs will take over the CeSC compartment and the entire SC niche with a very high probability

The numerical simulations and analytic calculations illustrated in the Materials and Methods section show that the fixation probability of CeSC mutants in the CeSC compartment is an increasing function of the mutants' fitness r . Furthermore, the probability that the progeny of CeSC advantageous mutants will take over the S_c compartment is very high. Furthermore the analytic results and simulations show that this probability is not very sensitive to α , when γ (the probability that CeSCs proliferate) is large or when the mutants are not advantageous (Figure 2).

The results of simulations, which are in perfect agreement with analytic calculations (Figures 2 and 3), show that with a high probability the progeny of CeSC mutants will take over the entire SC niche. Moreover, this probability is independent of σ (the probability of symmetric division). The time to fixation

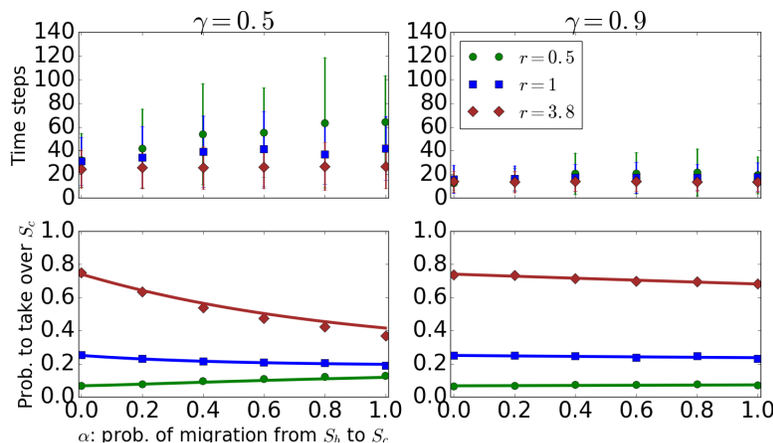


Fig. 2: The probability that the progeny of a CeSC mutant taking over the CeSC compartment.

The bottom sub-figures show the mean and standard deviations of results of simulations (points and bars) and results of formula (7) (solid lines) for the probability that the progeny of a CeSC mutant will take over the CeSCs. The top sub-figures represent the results of simulations for the time that the progeny of one mutant CeSC needs to take over the S_c compartment. The parameters for this figure are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

of advantageous mutants increases and their fixation probability decreases, when the migration rate from S_b to S_c rises (see Figure 2).

It is unlikely that the progeny of a BSC mutant take over the BSC compartment and the entire SC niche

The analytic calculations and simulations, which are in perfect agreement, show that the fixation probability of BSC mutants in BSC and CeSC compartments is an increasing function of α , the probability of migration from S_b to S_c (Figures 5 and 6). When $\alpha = 0$, BSCs do not have a chance to move to the CeSC compartment. Therefore, with a high chance BSC mutants will differentiate and be removed from the niche. Furthermore, advantageous mutants differentiate fast, because with a high probability they will be chosen to divide (i.e. differentiate). Since differentiations only happen in the BSC compartment, the probability that the progeny of a BSC mutant will take over the entire SC niche is very small. For example, the fixation probability of one BSC mutant in the BSC and CeSC compartments are approximately zero when $\alpha = 0$, and these probabilities are around 0.05 when $\alpha = 1$ for $\gamma = 0.5$ (Figures 5, 6, and 7). Although the time that BSC mutants need to take over the niche is increasing when stem cells divide mostly asymmetrically. The fixation probability of BSC mutants in the entire SC niche does not depend on σ , the probability of symmetric division, when $\sigma > 0$ (Figure 7).

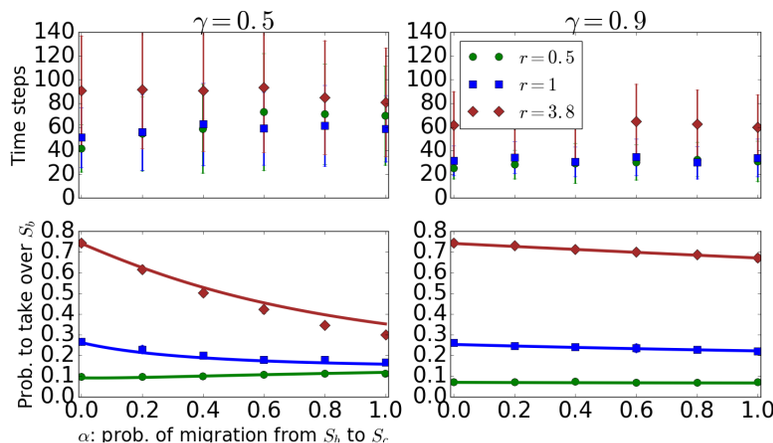


Fig. 3: The probability that the progeny of a CeSC mutant taking over the BSC compartment.

The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars) and results of formula (10) (solid lines) for the probability that the progeny of a mutant CeSC will take over BSCs. The top sub-figures show the results of simulations for the time that the progeny of one mutant CeSC needs to take over the S_b compartment. The parameters are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

The fixation probability of mutants does not depend on the probability of migration from S_b to S_c when γ is close to one

Simulations and analytic calculations show that the fixation probability of mutants in the SC niche is an increasing function of α , the probability of migration from S_b to S_c , when γ is not large. However, the fixation probability of mutants in the niche is not sensitive to α when most of the stem cell proliferations occur in the CeSC compartment, i.e. $\gamma > 0.9$. BSCs have a chance to migrate to the CeSC compartment when proliferations occur in the BSC compartment. Thus, when most of the proliferations happen in the CeSC compartment, then BSCs have a very small chance to migrate from S_b to the CeSC compartment. This leads to the probability of the progeny of a mutant stem cell taking over the niche becoming almost insensitive to α when γ is large (Figures 2,3, 5, and 6).

Migration from S_b to S_c delays tumorigenesis

The fixation probability of neutral and disadvantages CeSC mutants in the entire SC niche is not very sensitive to α , which is the probability of migration from S_b to S_c . However, the fixation probability of advantageous CeSC mutants is decreasing function of α . Moreover, simulations show that the time to fixation of CeSC mutants in the SC niche is an increasing function of α for advantageous mutants (Fig 8). In other words, a higher rate of migration from S_b to S_c maximizes the time that the progeny of advantageous CeSC mutants need to take over the SC niche. Therefore, the optimal value for α in terms of delaying the spread of advantageous CeSC mutants is one.

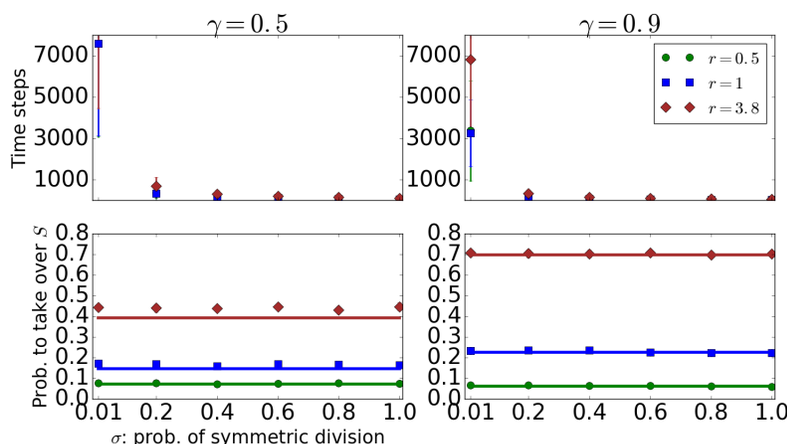


Fig. 4: The time and probability that the progeny of a CeSC mutant taking over the SC niche.

The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars) and the results of formula (2) (solid curves) for the probability that the progeny of a mutant CeSC will take over the SC niche as a function of σ , the probability of symmetric division. The top sub-figures show the results of simulations for the time that the progeny of one mutant CeSC needs to take over the niche. The parameters are $\alpha = 0.5$, $|S_c| = 4$, and $|S_b| = 7$.

The fixation probability of mutants in the SC niche is independent of the probability of symmetric division (σ) when $\sigma > 0$

Although the symmetric division of stem cells minimizes the probability of double-hit mutant production [5,37], the fixation probability of mutants in the SC niche is independent of the rate of symmetric stem cell division (σ) as long as stem cells do not divide fully asymmetrically ($\sigma \neq 0$) (Figures 4 and 7). When no symmetric proliferation occurs in the SC niche, then the mutant stem cells do not have a chance to spread in the niche. Furthermore, the time that mutant stem cells need to take over the stem cell niche is a decreasing function of the probability of symmetric division σ . In other words, asymmetric division delays the process of spreading mutants in the stem cell niche.

Optimal structure

Figure 10 (a)-(c) represents a contour plot for the probability of a CeSC mutant taking over the SC niche varying α , which is the probability of migration from S_b to S_c , and γ , which is the probability of proliferation in S_c . The fixation probability of CeSC mutants is zero when $\gamma = 0$, because in this case no division will happen in the CeSC compartment. When $\gamma \approx 0$, the minimum fixation probability of a disadvantageous CeSC mutant in the entire SC niche corresponds to $\alpha \approx 0$. However, if $\gamma \neq 0$, then the probability that the progeny of an advantageous or neutral CeSC mutant will take over the entire niche is minimized when $\alpha > 0.2$ and $\gamma \approx 0$. Additionally, when $\gamma \gg 0$, the minimum value of the fixation probability of CeSc mutants in the entire SC niche corresponds to the $\alpha = 1$. Note, when $\alpha = 1$, BSCs have a high chance

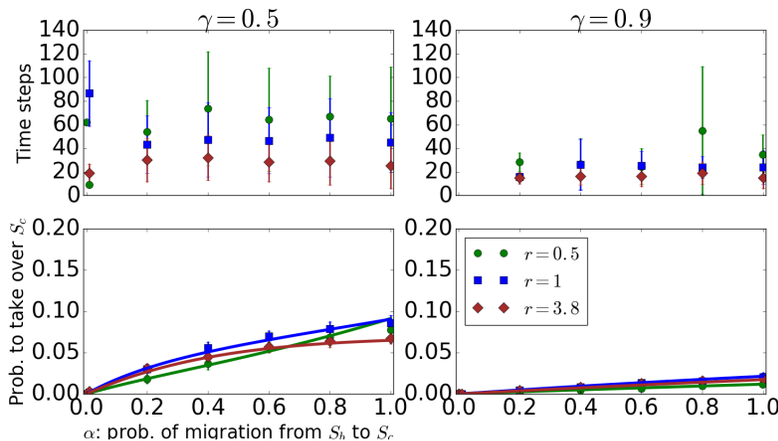


Fig. 5: The probability that the progeny of a BSC mutant taking over the CeSC compartment.

The bottom sub-figures indicate the mean and standard deviations of results of simulations (points and bars) and results of formula (7) (solid lines) for the probability that the progeny of a mutant BSC will take over the CeSCs. The top sub-figures represent the results of simulations for the time that the progeny of one mutant BSC needs to take over the S_c compartment. The parameters for this figure are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

to migrate to CeSCs, therefore CeSC mutants gain a high chance to move to S_b , because each migration from S_b to S_c is coupled with a migration from S_c to S_b . Then, with a high probability, mutants in the S_b compartment will differentiate and will be removed from the SC niche.

Figure 10 (d)-(f) indicates the probability of a BSC mutant taking over the entire SC compartment, which is very small, as a function of α and γ . In this case the minimum fixation probability, which is zero, happens when $\gamma = 1$, because in this case, all stem cell proliferations occur in the CeSC compartment and BSCs only differentiate. When $\gamma < 1$, the fixation probability of BSC mutants in the entire SC niche is an increasing function of α . The maximum of the probability that the progeny of a mutant BSC cell will take over the entire SC, which is around 0.2 for neutral and disadvantageous mutants and around 0.1 for advantageous mutants, happens when $\gamma = 0$ and $\alpha = 1$.

From the figure 11, which shows the summation of the fixation probability of a BSC mutant and the fixation probability of a CeSC mutant within the entire niche, we may conclude that the optimal structure of the SC niche, which minimize the spread of both CeSC and BSC mutants in the entire niche, should have a very small number of proliferations in the CeSC compartment, i.e. $0 < \gamma \approx 0$. Furthermore, from this figure and the results provided above, we may conclude that the optimal stem cell architecture of the niche, which delays and minimizes the spread of mutants, includes a small non-zero number of migrations from BSCs to CeSCs.

When stem cells divide fully asymmetrically ($\sigma = 0$), then no mutants will have a chance to spread in the SC niche, i.e. the probability of fixation is zero. Simulations and analytic calculations show that when $\sigma > 0$, the fixation probability of mutants in the niche is not sensitive to the probability of symmetric

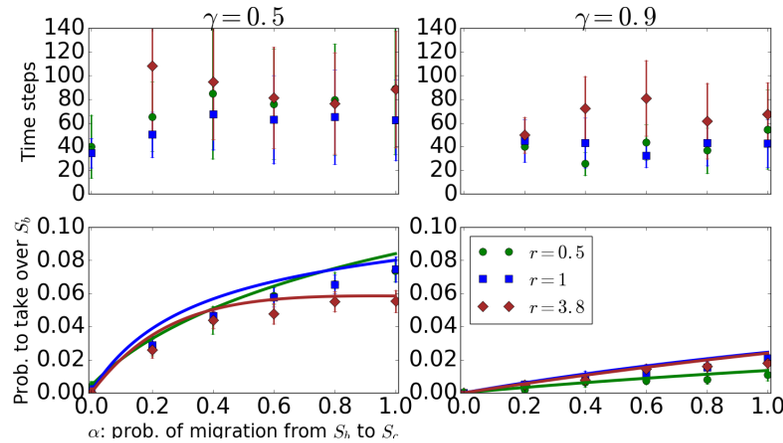


Fig. 6: The time and probability that the progeny of a BSC mutant taking over the BSC compartment. The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars) and results of formula (10) (solid lines) for the probability that the progeny of a mutant BSC will take over the BSCs. The top sub-figures show the results of simulations for the time that the progeny of one mutant BSC needs to take over the S_b compartment. The parameters for this figure are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

division σ . However, the time to fixation is a decreasing function of σ . The maximum time of fixation happens when σ is close to zero. This result is valid for both CeSC mutants and BSC mutants. Thus, the optimal value for σ , which delays and lowers the spread of mutants, is zero.

Discussion

Shahriyari and Komarova [35] have developed a bi-compartmental model for the SC niche to investigate the probability of double-hit mutant production in the niche. They found that the existence of two stem cell groups delays the double-hit mutant production. The current work generalizes that model by adding the possibility of migration from BSCs to CeSCs to study the dynamics of mutants in the SC niche. Furthermore, this work provides a framework to obtain the fixation probability of mutants in a bi-compartmental stochastic model. Analytic formulas have been obtained to calculate the fixation probability of mutants in each of compartments as well as the entire system.

The present study reveals that the small probability of migration from S_b to S_c , which is reported in recent experimental observations [16], causes more delay in the process of tumorigenesis by delaying the fixation of mutants in the niche. The analytic calculations and numerical simulations, which are in perfect agreement, show that the probability of BSC mutants taking over the entire SC niche is increased when the rate of migrations from BSCs to central stem cells is increased. Additionally, the probability of CeSC mutants taking over the entire niche is decreasing when the probability of migration from BSCs to CeSCs, which is denoted by α , is increasing. Moreover, the time that CeSC mutants need to take over the niche is increasing when α is increasing. Note, the probability of mutants' occurrences is higher in the BSC

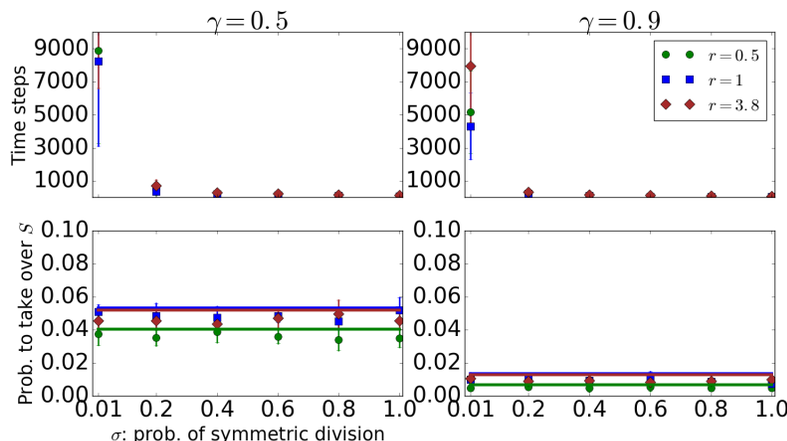


Fig. 7: The time and probability that the progeny of a BSC mutant taking over the SC niche.

The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars) and results of formula (2) (solid lines) for the probability that the progeny of a mutant BSC will take over the SC niche as a function of σ , the probability of symmetric division. The top sub-figures show the results of simulations for the time that the progeny of one mutant BSC needs to take over the niche. The parameters are $\alpha = 0.5$, $|S_c| = 4$, and $|S_b| = 7$.

compartment comparing to the CeSC compartment [38]. Moreover, the probability that a BSC cell spreads over the SC niche is much smaller than the probability of the progeny of a CeSC mutant taking over the SC niche. Thus, we may conclude that the optimal value for α should be a small non-zero number.

Furthermore, the optimal structure of the SC niche must delay the spread of mutants as well as their production. It has been shown that the symmetric division of stem cells delays the production of mutants [5, 35, 37]. Moreover, the model introduced in [38] indicates that the type of stem cell divisions (symmetric and asymmetric) is not as important as the rate of divisions at each location in the tissue in terms of delaying mutants production. That model also reveals that the natural division rate of cells in the colon and intestinal crypts delays the production of mutants in the CeSC compartment. This present study shows that the location of mutants is very important in the probability that their progeny will take over the entire crypt. Mutants in the CeSC compartment have a higher chance to take over the entire crypt. Therefore, one may conclude that the natural architecture of the SC niche is the optimal structure in terms of minimizing the probability of formation of tumors.

Materials and Methods

Transition Probabilities

In order to calculate the fixation probabilities, we obtain all non-zero transition probabilities, which are the probabilities of moving from the state (e^*, b^*) , which includes e^* number of S_c mutant and b^* number of S_b mutants, to another state in an updating time step:

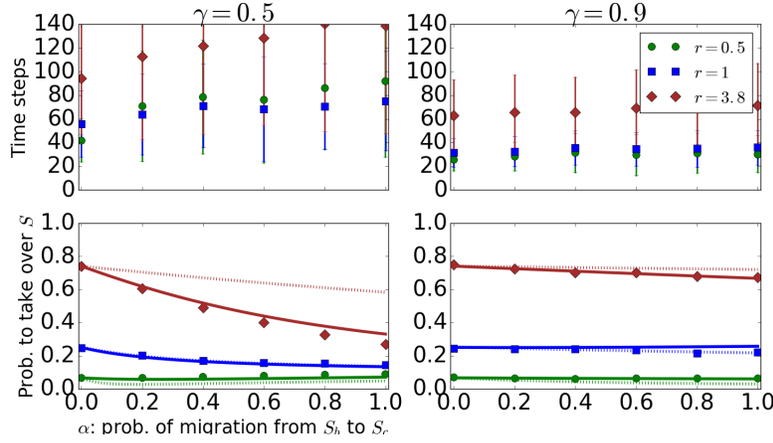


Fig. 8: The time and probability that the progeny of a CeSC mutant taking over the entire SC niche. The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars), results of formula (26) (solid lines), and approximated formula (12) (dashed lines) for the probability that the progeny of a mutant CeSC will take over the niche. The top sub-figures show the results of simulations for the time that the progeny of one mutant CeSC needs to take over the entire niche. The parameters for this figure are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

$$\begin{aligned}
 & \text{Migrations: a mutant BSC to } S_c \text{ and a w.t. } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*)} &= \underbrace{\sigma \frac{b}{rb^* + b}}_{\text{Diff. of a w.t. } S_b} \left[\underbrace{(1-\gamma) \frac{rb^*}{rb^* + (b-1)}}_{\text{Proliferation of a mutant BSC}} \underbrace{\alpha \frac{b^*-1}{b+b^*} \frac{e}{e+e^*}}_{\text{Prolif. of a } S_c \text{ mutant and migration of a w.t. } S_c \text{ to } S_b} + \underbrace{\gamma \frac{re^*}{\mathcal{R}_c} \frac{e}{e+e^*}}_{\text{Prolif. of a w.t. } S_c \text{ mutant and migration of a w.t. } S_c \text{ to } S_b} \right], \\
 & \text{Migrations: a w.t. BSC to } S_c \text{ and a mutant } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*-1, b^*)} &= \underbrace{\sigma \frac{rb^*}{rb^* + b}}_{\text{Diff. of a w.t. } S_b} \left[\underbrace{(1-\gamma) \frac{b}{r(b^*-1) + b}}_{\text{Prolif. of a w.t. BSC}} \underbrace{\alpha \frac{b+1}{b+b^*} \frac{e^*}{e+e^*}}_{\text{Prolif. of a w.t. } S_c \text{ and migration of a } S_c \text{ mutant to } S_b} + \underbrace{\gamma \frac{e}{\mathcal{R}_c} \frac{e^*}{e+e^*}}_{\text{Prolif. of a w.t. } S_c \text{ mutant and migration of a w.t. } S_c \text{ to } S_b} \right], \\
 & \text{Migrations: a w.t./mutant BSC to } S_c \text{ and a w.t./mutant } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*, b^*+1)} &= \underbrace{\sigma \frac{b}{rb^* + b}}_{\text{Diff. of a } S_b \text{ mutant}} \left[\underbrace{(1-\gamma) \frac{rb^*}{rb^* + (b-1)}}_{\text{Prolif. of a mutant BSC}} \underbrace{[(1-\alpha) + \alpha \frac{(b-1)e + (b^*+1)e^*}{(b+b^*)(e+e^*)}]}_{\text{No migration}} + \underbrace{\gamma \frac{re^*}{\mathcal{R}_c} \frac{e^*}{e+e^*}}_{\text{Prolif. \& mig. of a mutant } S_c} \right], \\
 & \text{Migrations: a w.t./mutant BSC to } S_c \text{ and a w.t./mutant } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*, b^*-1)} &= \underbrace{\sigma \frac{rb^*}{rb^* + b}}_{\text{Diff. of a } S_b \text{ mutant}} \left[\underbrace{(1-\gamma) \frac{b}{r(b^*-1) + b}}_{\text{Prolif. of a mutant BSC}} \underbrace{[(1-\alpha) + \alpha \frac{(b+1)e + (b^*-1)e^*}{(b+b^*)(e+e^*)}]}_{\text{No migration}} + \underbrace{\gamma \frac{e}{\mathcal{R}_c} \frac{e}{e+e^*}}_{\text{Prolif. \& mig. of a w.t. } S_c} \right],
 \end{aligned}$$

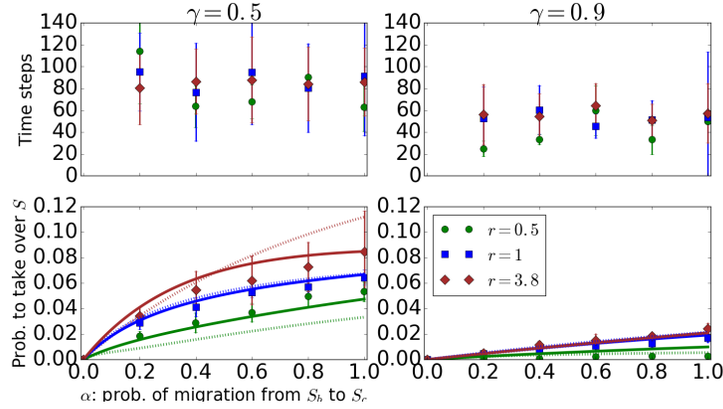


Fig. 9: **The time and probability that the progeny of a BSC mutant taking over the entire SC niche.** The bottom sub-figures represent the mean and standard deviations of results of simulations (points and bars), results of formula (26) (solid lines), and approximated formula (12) (dashed lines) for the probability that the progeny of a mutant BSC will take over the niche. The top sub-figures show the results of simulations for the time that the progeny of a mutant BSC needs to take over the entire niche. The parameters for this figure are $\sigma = 1$, $|S_c| = 4$, and $|S_b| = 7$.

$$\begin{aligned}
 & \text{Migrations: a mutant BSC to } S_c \text{ and a w.t. } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*-1)} &= \underbrace{\sigma \frac{b}{rb^*+b}}_{\text{Diff. of a w.t. } S_b} \underbrace{(1-\gamma) \frac{b-1}{rb^*+b-1}}_{\text{Prolif. of a w.t. BSC}} \alpha \frac{b^*}{b+b^*} \frac{e}{e+e^*} \\
 & \text{Migrations: a mutant BSC to } S_c \text{ and a w.t. } S_c \text{ to } S_b \\
 & + \underbrace{\sigma \frac{rb^*}{rb^*+b}}_{\text{Diff. of a } S_b \text{ mutant}} \left[\underbrace{(1-\gamma) \frac{r(b^*-1)}{r(b^*-1)+b}}_{\text{Prolif. of a mutant BSC}} \alpha \frac{b^*+1}{b+b^*} \frac{e}{e+e^*} + \underbrace{\gamma \frac{re^*}{\mathcal{R}_c} \frac{e}{e+e^*}}_{\text{Prolif. of a mutant } S_c \text{ \& mig. of a w.t. } S_c \text{ to } S_b} \right], \\
 & \text{Prolif. of a mutant BSC} \\
 P_{(e^*, b^*) \rightarrow (e^*-1, b^*+1)} &= \underbrace{\sigma \frac{rb^*}{rb^*+b}}_{\text{Diff. of a } S_b \text{ mutant}} (1-\gamma) \frac{r(b^*-1)}{r(b^*-1)+b} \alpha \frac{b}{b+b^*} \frac{e^*}{e+e^*} \\
 & \text{Migrations: a w.t. BSC to } S_c \text{ and a mutant } S_c \text{ to } S_b \\
 & \text{Migrations: a w.t. BSC to } S_c \text{ and a mutant } S_c \text{ to } S_b \\
 & + \underbrace{\sigma \frac{b}{rb^*+b}}_{\text{Diff. of a w.t. } S_b} \left[\underbrace{(1-\gamma) \frac{b-1}{rb^*+b-1}}_{\text{Prolif. of a w.t. BSC}} \alpha \frac{b}{b+b^*} \frac{e^*}{e+e^*} + \underbrace{\gamma \frac{e}{\mathcal{R}_c} \frac{e^*}{e+e^*}}_{\text{Prolif. of a w.t. } S_c \text{ \& mig. of a mutant } S_c \text{ to } S_b} \right], \\
 & \text{Migrations: a w.t. BSC to } S_c \text{ and a mutant } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*-1, b^*+2)} &= \underbrace{\sigma \frac{b}{rb^*+b}}_{\text{Diff. of a w.t. } S_b} (1-\gamma) \frac{rb^*}{rb^*+b-1} \alpha \frac{b-1}{b+b^*} \frac{e^*}{e+e^*}, \\
 & \text{Prolif. of a mutant BSC} \\
 & \text{Migrations: a mutant BSC to } S_c \text{ and a w.t. } S_c \text{ to } S_b \\
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*-2)} &= \underbrace{\sigma \frac{rb^*}{rb^*+b}}_{\text{Diff. of a mutant } S_b} (1-\gamma) \frac{b}{r(b^*-1)+b} \alpha \frac{b^*}{b+b^*} \frac{e}{e+e^*}. \\
 & \text{Prolif. of a w.t. BSC}
 \end{aligned}$$

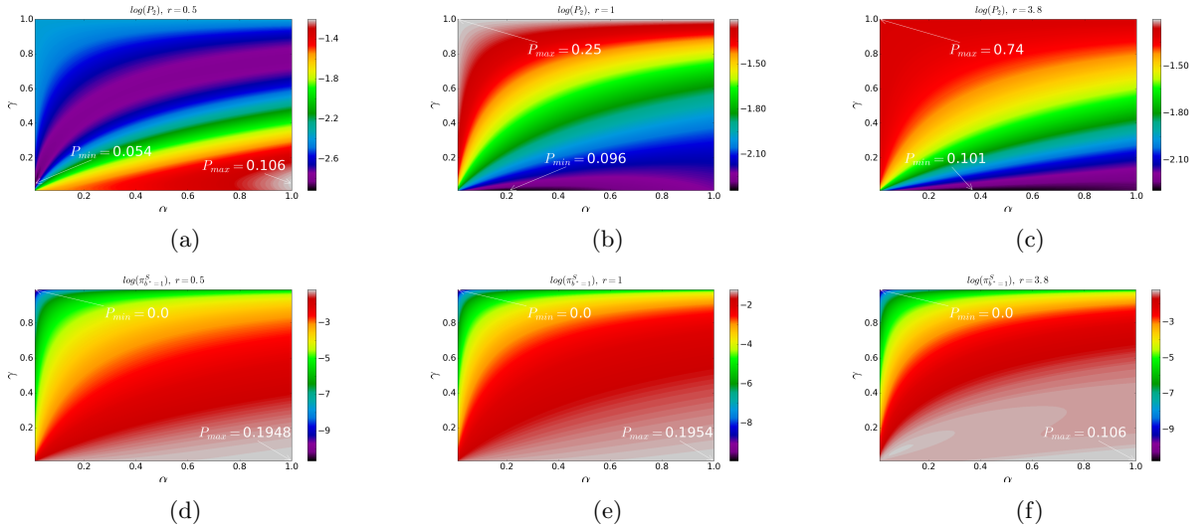


Fig. 10: Contour plot of the fixation probability of an initial mutant in the entire SC niche. Sub-figures (a)-(c) show the probability of the progeny of a S_c mutant in taking over the entire SC niche. Plots (d)-(f) indicate the probability of the progeny of a S_b mutant taking over the entire SC niche. Parameters are $|S_c| = 4$, $|S_b| = 7$, (a,d): $r = 0.5$, (b,e): $r = 1$, and (c,f): $r = 3.8$.

Fixation probability of mutants in the entire SC niche

We denote the probability of the progeny of e^* number of S_c mutants and b^* number of S_b mutants taking over the entire SC niche by $\pi^S(e^*, b^*)$. The fixation probability $\pi^S(e^*, b^*)$, which is the probability of moving from the state (e^*, b^*) to the state $(|S_c|, |S_b|)$, i.e. all cells become mutants, satisfies the following system of equations.

$$\begin{aligned}
 \pi^S(e^*, b^*) &= \sum_{(m,n)} P_{(e^*, b^*) \rightarrow (m,n)} \pi^S(m, n) \\
 &= \sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)} \pi^S(m, n) + \left[1 - \sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)} \right] \pi^S(e^*, b^*), \\
 \pi^S(0, 0) &= 0, \quad \pi^S(|S_c|, |S_b|) = 1.
 \end{aligned} \tag{1}$$

We can rewrite the above system of equations (1) in the form of

$$P^S \Pi^S = b^S, \tag{2}$$

where P^S , Π^S , and b^S are $((|S_c| + 1)(|S_b| + 1) - 1) \times ((|S_c| + 1)(|S_b| + 1) - 1)$, $((|S_c| + 1)(|S_b| + 1) - 1) \times 1$, and $((|S_c| + 1)(|S_b| + 1) - 1) \times 1$ dimensional matrices. We eliminate the state $(0, 0)$, because $\pi^S(0, 0) = 0$. The transition probability $P_{(e^*, b^*) \rightarrow (m,n)}$ is located in the $e^*(|S_b| + 1) + b^*$ row and $m(|S_b| + 1) + n$ column of the

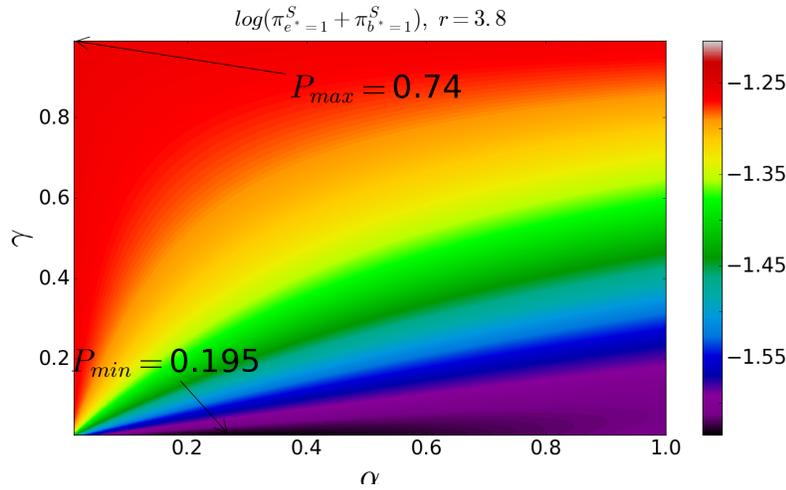


Fig. 11: **Contour plot of the summation of fixation probability of a CeSC mutant and fixation probability of a BSC mutant in the entire SC niche.** This figure shows the summation of the probability of the progeny of a S_c mutant taking over the entire SC niche and the probability of the progeny of a S_b mutant taking over the entire SC niche. Parameters are $|S_c| = 4$, $|S_b| = 7$, and $r = 3.8$.

matrix P if $(e^*, b^*) \neq (m, n)$. If $(e^*, b^*) = (m, n)$, the element in the $e^*(|S_b| + 1) + b^*$ row and $e^*(|S_b| + 1) + b^*$ column of the matrix P is $-\sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)}$, i.e.

$$\begin{aligned}
 P[e^*(|S_b| + 1) + b^*, m(|S_b| + 1) + n] &= P_{(e^*, b^*) \rightarrow (m,n)}, & \text{if } (e^*, b^*) \neq (m, n) \\
 P[e^*(|S_b| + 1) + b^*, e^*(|S_b| + 1) + b^*] &= -\sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)}, & \text{if } (e^*, b^*) \neq (|S_c|, |S_b|) \\
 P[e^*(|S_b| + 1) + b^*, e^*(|S_b| + 1) + b^*] &= 1 & \text{if } (e^*, b^*) = (|S_c|, |S_b|) \\
 \Pi^S[e^*(|S_b| + 1) + b^*] &= \pi^S(e^*, b^*) \\
 b^S[e^*(|S_b| + 1) + b^*] &= 0 & \text{if } (e^*, b^*) \neq (|S_c|, |S_b|) \\
 b^S[e^*(|S_b| + 1) + b^*] &= 1 & \text{if } (e^*, b^*) = (|S_c|, |S_b|)
 \end{aligned} \tag{3}$$

In other words,

$$P^S = \begin{bmatrix} -\sum_{(m,n) \neq (0,1)} P_{(0,1) \rightarrow (m,n)} & P_{(0,1) \rightarrow (0,2)} & \cdots & P_{(0,1) \rightarrow (|S_c|, |S_b| - 1)} & P_{(0,1) \rightarrow (|S_c|, |S_b|)} \\ & \cdot & & & \\ & & \cdot & & \\ & & & \cdot & \\ P_{(|S_c|, |S_b| - 1) \rightarrow (0,1)} & P_{(|S_c|, |S_b| - 1) \rightarrow (0,2)} & \cdots & -\sum_{(m,n) \neq (|S_c|, |S_b| - 1)} P_{(|S_c|, |S_b| - 1) \rightarrow (m,n)} & P_{(|S_c|, |S_b| - 1) \rightarrow (|S_c|, |S_b|)} \\ & 0 & 0 & 0 & 1 \end{bmatrix}, \quad (4)$$

$$\Pi^S = \begin{bmatrix} \pi^S(0,1) \\ \cdot \\ \cdot \\ \cdot \\ \pi^S(|S_c|, |S_b| - 1) \\ \pi^S(|S_c|, |S_b|) \end{bmatrix}, \quad b^S = \begin{bmatrix} 0 \\ \cdot \\ \cdot \\ \cdot \\ 0 \\ 1 \end{bmatrix}. \quad (5)$$

Fixation probability of mutants in the S_c compartment

Following the same technique, we calculate the fixation probability in the CeSc compartment and denote the fixation probability of e^* number of CeSc mutants and b^* number of BSC mutants in the CeSc compartment by $\pi^{S_c}(e^*, b^*)$. The $\pi^{S_c}(e^*, b^*)$ satisfies the following equations.

$$\begin{aligned} \pi^{S_c}(e^*, b^*) &= \sum_{(m,n)} P_{(e^*, b^*) \rightarrow (m,n)} \pi^{S_c}(m, n) \\ &= \sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)} \pi^{S_c}(m, n) + \left[1 - \sum_{(m,n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m,n)} \right] \pi^{S_c}(e^*, b^*), \\ \pi^{S_c}(0,0) &= 0, \quad \pi^{S_c}(|S_c|, b^*) = 1 \quad 0 \leq b^* \leq |S_b| \end{aligned} \quad (6)$$

In order to find the matrix representation of the equations (6), we combine all states $(|S_c|, b^*) \quad 0 \leq b^* \leq |S_b|$ to one state. Therefore, the equations can be rewritten in the form of

$$P^{S_c} \Pi^{S_c} = b^{S_c}, \quad (7)$$

where P^{S_c} , Π^{S_c} , and b^{S_c} are $(|S_c|(|S_b| + 1) - 1) \times (|S_c|(|S_b| + 1) - 1)$, $(|S_c|(|S_b| + 1) - 1) \times 1$, and $(|S_c|(|S_b| + 1) - 1) \times 1$.

1) $-1) \times 1$ dimensional matrices, and

$$\begin{aligned}
 P^{S_c}[e^*(|S_b| + 1) + b^*, m(|S_b| + 1) + n] &= P_{(e^*, b^*) \rightarrow (m, n)}, & \text{if } (e^*, b^*) \neq (m, n) \ \& \ e^*, m \neq |S_c| \\
 P^{S_c}[e^*(|S_b| + 1) + b^*, e^*(|S_b| + 1) + b^*] &= -\sum_{(m, n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m, n)}, & \text{if } e^*, m \neq |S_c| \\
 P^{S_c}[e^*(|S_b| + 1) + b^*, m(|S_b| + 1)] &= \sum_n P_{(e^*, b^*) \rightarrow (|S_c|, n)}, & \text{if } e^* \neq |S_c|, m = |S_c| \\
 P^{S_c}[e^*(|S_b| + 1) - 1, m(|S_b| + 1) - 1] &= 1 & \text{if } e^* = m = |S_c| \\
 \Pi^{S_c}[e^*(|S_b| + 1) + b^*] &= \pi^{S_c}(e^*, b^*) & \text{if } e^* \neq |S_c| \\
 \Pi^{S_c}[e^*(|S_b| + 1) - 1] &= \pi^{S_c}(|S_c|, b^*) = 1 & \text{if } e^* = |S_c|, \text{ for every } 0 \leq b^* \leq S_b \\
 b^{S_c}[e^*(|S_b| + 1) + b^*] &= 0 & \text{if } e^* \neq |S_c| \\
 b^{S_c}[e^*(|S_b| + 1) - 1] &= 1 & \text{if } e^* = |S_c|
 \end{aligned} \tag{8}$$

Note, the elements in the locations $|S_c|(|S_b| + 1) + b^*$ for $0 \leq b^* \leq S_b$ in the P^S , π^S , and b^S matrices have been summed as one entry in the P^{S_c} , π^{S_c} , and b^{S_c} matrices.

$$P^{S_c} = \begin{bmatrix} -\sum_{(m, n) \neq (0, 1)} P_{(0, 1) \rightarrow (m, n)} & P_{(0, 1) \rightarrow (0, 2)} & \cdots & P_{(0, 1) \rightarrow (|S_c| - 1, |S_b|)} & \sum_{b^*} P_{(0, 1) \rightarrow (|S_c|, b^*)} \\ & \cdot & & & \\ & & \cdot & & \\ & & & \cdot & \\ P_{(|S_c| - 1, |S_b|) \rightarrow (0, 1)} & P_{(|S_c| - 1, |S_b|) \rightarrow (0, 2)} & \cdots & -\sum_{(m, n) \neq (|S_c| - 1, |S_b|)} P_{(|S_c| - 1, |S_b|) \rightarrow (m, n)} & \sum_{b^*} P_{(|S_c| - 1, |S_b|) \rightarrow (|S_c|, b^*)} \\ 0 & 0 & 0 & 0 & 1 \end{bmatrix}. \tag{9}$$

Fixation probability in the S_b compartment

Here, we obtain the fixation probability, $\pi^{S_b}(e^*, b^*)$ of e^* number of CeSC mutants and b^* number of BSC mutants in the BSC compartment. To calculate the fixation probability $\pi^{S_b}(e^*, b^*)$, which satisfies the following equation, in the BSC compartment, we combine all $(e^*, |S_c|)$ $0 \leq e^* \leq |S_c|$ to one state.

$$P^{S_b} \Pi^{S_b} = b^{S_b}, \tag{10}$$

where P^{S_b} , Π^{S_b} , and b^{S_b} are $((|S_c| + 1)|S_b| - 1) \times ((|S_c| + 1)|S_b| - 1)$, $((|S_c| + 1)|S_b| - 1) \times 1$, and $((|S_c| + 1)|S_b| - 1) \times 1$ dimensional matrices, and

$$\begin{aligned}
 P^{S_b}[e^*|S_b| + b^*, m|S_b| + n] &= P_{(e^*, b^*) \rightarrow (m, n)}, & \text{if } (e^*, b^*) \neq (m, n) \ \& \ b^*, n \neq |S_b| \\
 P^{S_b}[e^*|S_b| + b^*, e^*|S_b| + b^*] &= -\sum_{(m, n) \neq (e^*, b^*)} P_{(e^*, b^*) \rightarrow (m, n)}, & \text{if } b^*, n \neq |S_b| \\
 P^{S_b}[e^*|S_b| + b^*, m|S_b| + n] &= \sum_m P_{(e^*, b^*) \rightarrow (m, |S_b|)}, & \text{if } b^* \neq |S_b|, n = |S_b| \\
 P^{S_b}[|S_c||S_b| + b^* - 1, |S_c||S_b| + n - 1] &= 1 & \text{if } b^* = n = |S_b| \\
 \Pi^{S_b}[e^*|S_b| + b^*] &= \pi^{S_b}(e^*, b^*) & \text{if } b^* \neq |S_b| \\
 \Pi^{S_b}[(|S_c| + 1)|S_b| - 1] &= \pi^{S_b}(e^*, |S_b|) = 1 & \text{if } b^* = |S_b|, \text{ for every } 0 \leq e^* \leq S_c \\
 b^{S_b}[e^*|S_b| + b^*] &= 0 & \text{if } b^* \neq |S_b| \\
 b^{S_b}[|S_c||S_b| + b^* - 1] &= 1 & \text{if } b^* = |S_b|
 \end{aligned} \tag{11}$$

Note, the elements in the locations $e^*|S_b| + |S_b|$ for $0 \leq e^* \leq S_c$ in the P^S , π^S , and b^S matrices have been summed up as one entry in the P^{S_b} , π^{S_b} , and b^{S_b} matrices.

We use the matrix representation of the fixation probability equations to obtain the fixation probability of mutants at each of stem cell compartments S_c and S_b as well as the entire SC niche.

Approximated formula

To gain more insight about the fixation probability of mutants in the entire niche, we do more analytic analysis on the equation (1). For simplicity, we assume $\mathcal{R}_c \approx |S_c|$ and $\mathcal{R}_b \approx |S_b|$, and $\left(\frac{e^*}{|S_c|}\right)^2 \approx 0$, $\frac{e^* b^*}{|S_c||S_b|} \approx 0$, $\left(\frac{b^*}{|S_b|}\right)^2 \approx 0$. Under such a restriction of keeping the number of mutants small compared with $|S_c|$ and $|S_b|$ we can rewrite the transition probabilities as following

$$\begin{aligned}
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*)} &\approx \frac{\sigma r \gamma}{|S_c|} e^*, & P_{(e^*, b^*) \rightarrow (e^*-1, b^*)} &\approx 0, \\
 P_{(e^*, b^*) \rightarrow (e^*, b^*+1)} &\approx \frac{\sigma r (1 - \gamma)}{|S_b|} b^*, & P_{(e^*, b^*) \rightarrow (e^*, b^*-1)} &\approx \frac{\sigma r}{|S_b|} b^*, \\
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*-1)} &\approx \frac{\sigma (1 - \gamma) \alpha}{|S_b|} b^*, & P_{(e^*, b^*) \rightarrow (e^*-1, b^*+1)} &\approx \sigma \frac{\gamma + (1 - \gamma) \alpha}{|S_c|} e^*, \\
 P_{(e^*, b^*) \rightarrow (e^*+1, b^*-2)} &\approx 0, & P_{(e^*, b^*) \rightarrow (e^*-1, b^*+2)} &\approx 0.
 \end{aligned} \tag{12}$$

To obtain the fixation probability of mutants in the entire SC niche, we define the following generating function:

$$G(z_1, z_2; t) = \sum_{e^*, b^*} \phi(e^*, b^*; t) z_1^{e^*} z_2^{b^*}, \tag{13}$$

where $\phi(e^*, b^*; t)$ is the probability of being at the state (e^*, b^*) at time t , and it satisfies the following equation.

$$\frac{\partial \phi}{\partial t} \approx \phi(e^*, b^*; t+1) - \phi(e^*, b^*; t) = \sum_{(m,n)} \left\{ P_{(m,n) \rightarrow (e^*, b^*)} \phi(m, n; t) - P_{(e^*, b^*) \rightarrow (m,n)} \phi(e^*, b^*; t) \right\}, \quad (14)$$

From the equations (13) and (14), the transition probabilities (12), and using the substitutions $\langle e^* z_1^{e^*} z_2^{b^*} \rangle = z_1 \frac{\partial}{\partial z_1}$ and $\langle b^* z_1^{e^*} z_2^{b^*} \rangle = z_2 \frac{\partial}{\partial z_2}$, where $\langle \cdot \rangle$ is the summation over states (e^*, b^*) (see [39] for more details), we get

$$\begin{aligned} \frac{\partial G(e^*, b^*; t)}{\partial t} \approx & \left[r \gamma z_1 (z_1 - 1) + (\gamma + (1 - \gamma) \alpha) (z_2 - z_1) \right] \frac{\partial G}{|S_c| \partial z_1} \\ & + \left[r (z_2 - 1) \left((1 - \gamma) z_2 - 1 \right) + (1 - \gamma) \alpha (z_1 - z_2) \right] \frac{\partial G}{|S_b| \partial z_2}, \end{aligned} \quad (15)$$

At the steady state, we can derive the following system of equations, which can provide the fixed points (in fact, the pseudo-fixed points of the system) (z_1^*, z_2^*) of the generating function:

$$\begin{cases} r \gamma z_1^* (z_1^* - 1) + (\gamma + (1 - \gamma) \alpha) (z_2^* - z_1^*) = 0, \\ r (z_2^* - 1) \left((1 - \gamma) z_2^* - 1 \right) + (1 - \gamma) \alpha (z_1^* - z_2^*) = 0. \end{cases} \quad (16)$$

The latter system has two different solutions. The trivial solution is $(z_1^*, z_2^*) = (1, 1)$, and the second solution (z_1^*, z_2^*) satisfies in the following equations (for $\alpha \neq 0, \gamma < 1$):

$$z_1^* = \frac{1}{\alpha(1-\gamma)} \left\{ -r(1-\gamma)(z_2^*)^2 + (\alpha(1-\gamma) + r(2-\gamma))z_2^* - r \right\}, \quad (17)$$

$$\mathcal{A}_3 (z_2^*)^3 + \mathcal{A}_2 (z_2^*)^2 + \mathcal{A}_1 z_2^* + \mathcal{A}_0 = 0, \quad (18)$$

where

$$\mathcal{A}_0 = r \gamma \alpha (\gamma - 1) - \alpha^2 (1 - \gamma)^2 - \alpha \gamma (\gamma - 1) - r^2 \gamma, \quad (19)$$

$$\mathcal{A}_1 = -2r^2 \gamma^2 + \alpha \gamma^3 + 3r^2 \gamma + \alpha \gamma^3 r - 2\alpha^2 \gamma + \alpha^2 + 3r \gamma \alpha + \alpha \gamma - 4r \gamma^2 \alpha + \alpha^2 \gamma^2 - 2\alpha \gamma^2, \quad (20)$$

$$\mathcal{A}_2 = -2\alpha \gamma^3 r - r^2 \gamma^3 + 4r \gamma^2 \alpha - 3r^2 \gamma - 2r \gamma \alpha + 4r^2 \gamma^2, \quad (21)$$

$$\mathcal{A}_3 = r^2 \gamma (1 - \gamma)^2 \quad (22)$$

Note that when $\alpha = 0$ then the equation (16) results in the non-trivial solutions $z_1^* = 1/r$ and $z_2^* = 1/(1-\gamma)$. It can be easily checked that the boundary and initial conditions for the derived generating function respectively are as follows

$$G(z_1 = 1, z_2 = 1, t) = 1, \quad \forall t > 0, \quad (23)$$

$$G(z_1, z_2, t = 0) = z_1^{i_1} z_2^{i_2}, \quad (24)$$

where i_1 and i_2 are assumed as the initial number of mutant CeSC(s) and BSC(s) respectively. The system has two distinct absorption states: the progeny of the initial mutant either fixated or eliminated from the SC population. More precisely at the steady state, we have

$$G(z_1, z_2; t \rightarrow \infty) = p\left(e^* = 0, b^* = 0; t \rightarrow \infty\right) + p\left(e^* = |S_c|, b^* = |S_b|; t \rightarrow \infty\right) z_1^{|S_c|} z_2^{|S_b|}. \quad (25)$$

Therefore, the fixation probability of e^* number of CeSC mutants and b^* number of BSC mutants in the entire SC niche is approximately

$$\pi_{(e^*, b^*)}^{SC} \approx \frac{1 - (z_1^*)^{e^*} (z_2^*)^{b^*}}{1 - (z_1^*)^{|S_c|} (z_2^*)^{|S_b|}}. \quad (26)$$

Author Contributions

L.S. designed research; L.S. and A.M.S. performed research; L.S. and A.M.S. contributed analytic tools; L.S. performed numerical simulations; L.S. and A.M.S. analyzed data; L.S. and A.M.S. wrote the manuscript.

Acknowledgment

This research has been supported in part by the Mathematical Biosciences Institute and the National Science Foundation under grant DMS 1440386. Authors declare that they have no competing interests.

References

- [1] Dah-Ching Ding, Yu-Hsun Chang, Woei-Cherng Shyu, and Shinn-Zong Lin. Human Umbilical Cord Mesenchymal Stem Cells: A New Era for Stem Cell Therapy. *Cell Transplantation*, 24(3):339–347, mar 2015.
- [2] C. Ganta, D. Chiyo, R. Ayuzawa, R. Rachakatla, M. Pyle, G. Andrews, M. Weiss, M. Tamura, and D. Troyer. Rat Umbilical Cord Stem Cells Completely Abolish Rat Mammary Carcinomas with No Evidence of Metastasis or Recurrence 100 Days Post-Tumor Cell Inoculation. *Cancer Research*, 69(5):1815–1820, feb 2009.
- [3] Hugo J Snippert, Laurens G Van Der Flier, Toshiro Sato, Johan H Van Es, Maaïke Van Den Born, Carla Kroon-Veenboer, Nick Barker, Allon M Klein, Jacco Van Rheenen, Benjamin D Simons, et al. Intestinal crypt homeostasis results from neutral competition between symmetrically dividing *lgr5* stem cells. *Cell*, 143(1):134–144, 2010.
- [4] D. Gleason, J.H. Fallon, M. Guerra, J.-C. Liu, and P.J. Bryant. Ependymal stem cells divide asymmetrically and transfer progeny into the subventricular zone when activated by injury. *Neuroscience*, 156(1):81–88, sep 2008.

-
- [5] Leili Shahriyari and Natalia L Komarova. Symmetric vs. Asymmetric Stem Cell Divisions: An Adaptation against Cancer? *PLoS ONE*, 8(10):e76195, oct 2013.
- [6] Toshiro Sato, Johan H van Es, Hugo J Snippert, Daniel E Stange, Robert G Vries, Maaïke van den Born, Nick Barker, Noah F Shroyer, Marc van de Wetering, and Hans Clevers. Paneth cells constitute the niche for Lgr5 stem cells in intestinal crypts. *Nature*, 469(7330):415–8, January 2011.
- [7] Piul Rabbani, Makoto Takeo, Weichin Chou, Peggy Myung, Marcus Bosenberg, Lynda Chin, M Mark Taketo, and Mayumi Ito. Coordinated activation of Wnt in epithelial and melanocyte stem cells initiates pigmented hair regeneration. *Cell*, 145(6):941–55, June 2011.
- [8] Paolo Bianco. Bone and the hematopoietic niche: a tale of two stem cells. *Blood*, 117(20):5281–8, May 2011.
- [9] Stefania Lymperi, Francesca Ferraro, and David T Scadden. The HSC niche concept has turned 31. Has our knowledge matured? *Annals of the New York Academy of Sciences*, 1192:12–8, March 2010.
- [10] Ya-Chieh Hsu and Elaine Fuchs. A family business: stem cell progeny join the niche to regulate homeostasis. *Nature reviews. Molecular cell biology*, 13(2):103–14, February 2012.
- [11] Anne Wilson, Gabriela M Oser, Maike Jaworski, William E Blanco-Bose, Elisa Laurenti, Christelle Adolphe, Marieke A Essers, H Robson MacDonald, and Andreas Trumpp. Dormant and self-renewing hematopoietic stem cells and their niches. *Annals of the New York Academy of Sciences*, 1106(1):64–75, 2007.
- [12] Adlen Foudi, Konrad Hochedlinger, Denille Van Buren, Jeffrey W Schindler, Rudolf Jaenisch, Vincent Carey, and Hanno Hock. Analysis of histone 2b-gfp retention reveals slowly cycling hematopoietic stem cells. *Nature biotechnology*, 27(1):84–90, 2009.
- [13] Ingmar Glauche, Kateri Moore, Lars Thielecke, Katrin Horn, Markus Loeffler, and Ingo Roeder. Stem cell proliferation and quiescence? two sides of the same coin. *PLoS computational biology*, 5(7):e1000447, 2009.
- [14] Marc HGP Raaijmakers and David T Scadden. Divided within: heterogeneity within adult stem cell pools. *Cell*, 135(6):1006–1008, 2008.
- [15] Valentina Greco and Shangqin Guo. Compartmentalized organization: a common and required feature of stem cell niches? *Development*, 137(10):1586–1594, 2010.
- [16] Laila Ritsma, Saskia I J Ellenbroek, Anoeck Zomer, Hugo J Snippert, Frederic J de Sauvage, Benjamin D Simons, Hans Clevers, and Jacco van Rheenen. Intestinal crypt homeostasis revealed at single-stem-cell level by in vivo live imaging. *Nature*, 507(7492):362–365, feb 2014.
- [17] Martin A Nowak. *Evolutionary dynamics*. Harvard University Press, 2006.
- [18] Dominik Wodarz and Natalia L. Komarova. *Dynamics of Cancer : Mathematical Foundations of Oncology*. 2014.

-
- [19] Matthew D Johnston, Carina M Edwards, Walter F Bodmer, Philip K Maini, and S Jonathan Chapman. Mathematical modeling of cell population dynamics in the colonic crypt and in colorectal cancer. *Proceedings of the National Academy of Sciences of the United States of America*, 104(10):4008–13, mar 2007.
- [20] Natalia L Komarova, Leili Shahriyari, and Dominik Wodarz. Complex role of space in the crossing of fitness valleys by asexual populations. *Journal of The Royal Society Interface*, 11(95):20140014–20140014, mar 2014.
- [21] Leili Shahriyari. Cell Dynamics in the Wound Healing Process in Tumor Environment After Treatments. Technical report, oct 2016.
- [22] I P M Tomlinson and W F Bodmer. Failure of programmed cell death and differentiation as causes of tumors : Some simple mathematical models [-2a2No (G)]. *Proc. Natl. Acad. Sci. U.S.A.*, 92(November):11130–11134, 1995.
- [23] Heiko Enderling and Philip Hahnfeldt. Cancer Stem Cells In Solid Tumors: Is 'Evading Apoptosis' A Hallmark Of Cancer? *Progress in Biophysics and Molecular Biology*, 2011.
- [24] Alexandra Jilkine and Ryan N Gutenkunst. Effect of dedifferentiation on time to mutation acquisition in stem cell-driven cancers. *PLoS computational biology*, 10(3):e1003481, March 2014.
- [25] Zheng Sun and Natalia L Komarova. Stochastic modeling of stem-cell dynamics with control. *Mathematical biosciences*, 240(2):231–40, December 2012.
- [26] M. Loeffler and I. Roeder. Tissue stem cells: definition, plasticity, heterogeneity, self-organization and models—a conceptual approach. *Cells Tissues Organs*, 171:8–26, 2002.
- [27] I. Glauche, Cross M., M. Loeffler, and I. Roeder. Lineage specification of hematopoietic stem cells: mathematical modeling and biological implications. *Stem Cells*, 25(53):1791–1799, 2007.
- [28] C. Colijn and M. C. Mackey. A mathematical model of hematopoiesis—I. Periodic chronic myelogenous leukemia. *Journal of Theoretical Biology*, 237:117–132, 2005.
- [29] David Dingli, Arne Traulsen, and Franziska Michor. (A)symmetric stem cell replication and cancer. *PLoS computational biology*, 3(3):e53, March 2007.
- [30] Heiko Enderling, L Hlatky, and Philip Hahnfeldt. Migration rules: tumours are conglomerates of self-metastases. *British journal of cancer*, 100(12):1917–25, June 2009.
- [31] John R Hutchinson, Dan Famini, Richard Lair, Rodger Kram, Steven A Frank, and Martin A Nowak. Developmental predisposition to cancer. *Nature*, 422(April):10570, 2003.
- [32] Martin A Nowak, Natalia L Komarova, Anirvan Sengupta, Prasad V Jallepalli, Ie-Ming Shih, Bert Vogelstein, and Christoph Lengauer. The role of chromosomal instability in tumor initiation. *Proceedings of the National Academy of Sciences of the United States of America*, 99(25):16226–31, December 2002.

- [33] Kamran Kaveh, Mohammad Kohandel, and Siv Sivaloganathan. Replicator dynamics of cancer stem cell: Selection in the presence of differentiation and plasticity. *Mathematical Biosciences*, 272:64–75, feb 2016.
- [34] Daniel Sánchez-Taltavull. Optimal architecture of differentiation cascades with asymmetric and symmetric stem cell division. *Journal of Theoretical Biology*, 407:106–117, oct 2016.
- [35] Leili Shahriyari and Natalia L Komarova. The role of the bi-compartmental stem cell niche in delaying cancer. *Physical Biology*, 12(5):055001, jul 2015.
- [36] Ali Mahdipour Shirayeh and Leili Shahriyari. New Insights into Initiation of Colon and Intestinal Cancer: The Significance of Central Stem Cells in the Crypt. oct 2016.
- [37] Peter T. McHale and Arthur D. Lander. The Protective Role of Symmetric Stem Cell Division on the Accumulation of Heritable Damage. *PLoS Computational Biology*, 10(8):e1003802, aug 2014.
- [38] Leili Shahriyari, Natalia L. Komarova, and Alexandra Jilkine. The role of cell location and spatial gradients in the evolutionary dynamics of colon and intestinal crypts. *Biology Direct*, 11(1):42, dec 2016.
- [39] Bahram Houchmandzadeh and Marcel Vallade. The fixation probability of a beneficial mutation in a geographically structured population. *New Journal of Physics*, 13(7):073020, 2011.