MATHEMATICAL MODEL OF COLORECTAL CANCER INITIATION

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ABSTRACT. Colorectal cancer (CRC) initiation is driven by the accumulation of critical genetic alterations, including the inactivation of tumor suppressor genes (APC and TP53) and the activation of the oncogene KRAS. This study builds upon the foundational stochastic model of Paterson et al.[1], reformulating it into a matrix-based framework to enhance the analysis of mutation dynamics and crypt expansion. By leveraging transition and growth matrices, we model the progression of 32 genotypic states, capturing the selective growth advantages of APC and KRAS mutations. Comparative results demonstrate that while the matrix approach aligns with tau-leaping simulations, it consistently underestimates malignancy probabilities at early ages, highlighting discrepancies attributable to outflow terms and double mutation assumptions. Our findings underscore the utility of matrix methods in elucidating cancer evolution and provide insights into CRC risk modeling, aligning theoretical predictions with observed epidemiological data.

1. Introduction

Cancer development is a complex evolutionary process driven by the accumulation of genetic alterations. In colorectal cancer (CRC), this progression typically involves specific mutations in key genes that regulate cell growth and division. By reviewing references [2, 3], we gained a basic understanding of colorectal cancer, learning about the critical role of three driver gene mutations in the development of lung and colorectal cancers. This forms the basis for recent research[1], which has established that just three driver mutations are often sufficient for malignant transformation: the inactivation of two tumor suppressor genes (TSGs) and the activation of one oncogene.

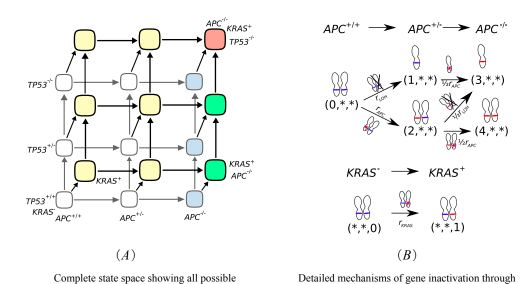


Figure 1. State transition diagram of colorectal cancer evolution¹

mutation and loss of heterozygosity.

Paterson et al.[1], in their seminal work, developed a mathematical model of CRC initiation that focuses on three critical genetic events: the inactivation of tumor suppressors APC and TP53,

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combinations of mutations in APC, TP53, and KRAS.

¹Figure 1 is adapted from the paper by Paterson et al[1].

and the activation of the KRAS oncogene. Their model, illustrated in Figure 1, presents the complex network of possible evolutionary pathways to cancer. Figure 1(A) shows the full state space of genetic alterations, where each node represents a unique combination of mutation states, and arrows indicate possible transitions between states. Figure 1(B) details the specific mechanisms of gene inactivation, showing how APC and TP53 can be inactivated through either mutation or loss of heterozygosity (LOH), while KRAS activation requires only a single mutation event. This comprehensive model accounts for all possible orders of mutation acquisition, allowing for a more realistic representation of cancer evolution compared to traditional linear models.

In this paper, we first analyze the two mathematical models used in Figure 2—ODE system and Tau-leaping. Then we reproduced the figure using the same method from the paper. Building upon their work, and inspired by [4], we present an alternative mathematical formulation using a matrix-based approach. Our method reformulates the evolutionary dynamics shown in Figure 1 as a system of linear transformations, where a transition matrix captures all possible mutation events between states, and a growth matrix represents the selective advantages conferred by different mutations. Following this theoretical foundation, we present comparative results between our matrix approach and the original methods, analyzing the similarities and differences in their predictions. For the second model, τ -leaping, we drew inspiration from the simulation methods described in [5] and [6]. After comparing the strengths and weaknesses of these two approaches, we developed our own implementation of τ -leaping in PYTHON to better align with our framework.

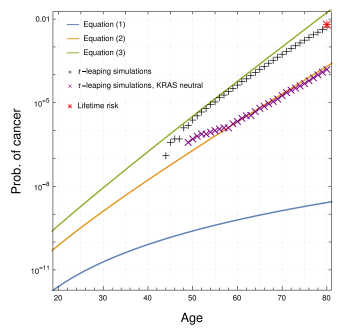


Figure 2. Age-dependent Probability of Cancer ²

2. Mathematical Model

Model Equations

Paterson et al. propose a stochastic mathematical model to describe the dynamics of driver gene mutations and crypt expansion during the initiation of CRC. This study primarily focuses on the probability of disease occurrence in humans, emphasizing the probability of cancerous crypts reaching the malignant state. The model is constructed by deriving the rate of change in the number of crypts for each genotype, combined with experimental parameter estimates from the

²Figure 2 is adapted from the paper by Paterson et al[1].

original study, to create a complete dynamic framework. Below are the detailed derivations and methods of the model.

- 2.1. Transition Rules and Parameter Settings. For the mutation in crypts, the driver events occur as stochastic processes. The inactivation of APC and TP53 requires both alleles to be inactivated, while the activation of KRAS only requires a single mutation. The inactivation of tumor suppressor genes is governed by both mutations and loss of heterozygosity (LOH) which is depicted in Figure 1(B). For example, in the case of APC:
 - If the first allele is inactivated through LOH, the second allele can only be inactivated through mutation at a rate $r_{\rm APC}/2$.
 - If the first allele is inactivated through mutation, the second allele can be inactivated either through LOH at a rate $r_{\text{LOH}}/2$ or through mutation at a rate $r_{\text{APC}}/2$.

A similar mechanism applies to TP53. The activation of KRAS occurs at a rate r_{KRAS} and requires only a single mutation. The mutation rate r_{APC} is calculated as $r_{APC} = u \cdot n_{APC}$, and we can get r_{KRAS} , r_{TP53} in same way. The parameter setting in the model can be found in Table 1.

| Parameter | Value | Unit | Definition | |
|--------------------------|-----------------------|----------|---|--|
| \overline{u} | 1.25×10^{-8} | per year | Base pair mutation rate per year | |
| \overline{t} | 0 - 80 | year | The time that model predict | |
| \overline{n} | 10^{8} | - | Total number of crypts in the colon | |
| $\overline{n_{ m APC}}$ | 604 | - | Number of driver sites in the APC gene | |
| n_{P53} | 73 | - | Number of driver sites in the TP53 gene | |
| $\overline{n_{ m KRAS}}$ | 20 | - | Number of driver sites in the KRAS gene | |
| b_1 | 0.2 | per year | Division rate of APC-/- crypts | |
| b_2 | 0.07 | per year | Division rate of KRAS+ crypts | |
| $r_{ m LOH}$ | 1.36×10^{-4} | per year | Loss of heterozygosity rate | |
| r_{TP53lost} | 0.0 | per year | Division rate of TP53-/- crypts | |
| $r_{ m LD}$ | 0.0 | per year | Division rate of crypts with other mutations | |
| $APC_{\text{mult}1}$ | 2.1 | - | Fixation multiplier for the first APC mutation | |
| $APC_{ m mult2}$ | 2.8 | - | Fixation multiplier for the second APC mutation | |
| $KRAS_{ m mult}$ | 3.6 | - | Fixation multiplier for KRAS mutation | |

Table 1. Model Parameters and Definitions

The genotype of a crypt is represented as a triplet (i, j, k), where:

- $i \in \{0, 1, 2, 3, 4\}$: Activation state of APC.
- $j \in \{0, 1, 2, 3, 4\}$: Activation state of *TP53*.
- $k \in \{0, 1\}$: Activation state of KRAS.

We use numbers to represent different states, with all possible states listed in Table 2. Since APC and TP53 share the same transformation rules, the states represented by the numbers are defined using the same set of rules.

| \overline{i} | Description of APC State | k | Description of KRAS State |
|----------------|--|---|---------------------------|
| 0 | Both APC alleles are wild-type | 0 | KRAS is wild-type |
| 1 | One APC allele lost through LOH (single loss) | 1 | KRAS is mutated |
| 2 | One APC allele inactivated by mutation (single mutation) | | |
| 3 | One APC allele lost (LOH), one inactivated by mutation | | |
| 4 | Both APC alleles inactivated by mutation (double mutation) | | |

Table 2. States of i and k

The model allows a total genotype space of $5 \cdot 5 \cdot 2 = 50$ states, including the malignant genotypes (3,3,1), (4,3,1), (3,4,1), and (4,4,1), which correspond to crypts that have accumulated all three driver events.

In theoretical studies, it is recognized that genetic mutations can lead to malignant transformations, necessitating the simulation of this process in the model. The model simulates crypts carrying APC or KRAS driver mutations gaining a growth advantage, resulting in increased division rates, which further enhance the likelihood of crypt malignant transformation. Specifically, the study sets the division rate of $APC^{-/-}$ crypts as $b_1 = 0.2$ /year, the division rate of $KRAS^+$ crypts as $b_2 = 0.07$ /year, and the division rate of double-mutant $APC^{-/-}KRAS^+$ crypts as $b_{12} = b_1 + b_2 = 0.27$ /year. It is assumed that genotypes without APC inactivation or KRAS activation have a division rate of 0. This assumption is based on experimental data indicating that normal crypts in adult colonic tissue divide very rarely[7].

2.2. **Probabilistic Model and Analysis.** We use the mean-field approximation to construct probabilistic models for the dynamics of crypt mutation and clonal expansion. In this approximation, the probability of double mutations is considered negligible, significantly reducing the complexity of the system. This allows the number of distinct genotypes to be reduced to 32, with the malignant state fully turned on represented by (3,3,1).

The probabilities of crypts with different genotypes transitioning to malignant states are comprehensively modeled through various mutation and LOH pathways. When a specific mutation step involves only one allele (e.g., LOH or single-allele mutation), the likelihood of one allele mutating is factored in, halving the mutation rate to reflect this probability. To determine the probability of the first malignant crypt arising from the non-malignant crypt population, while excluding contributions from already malignant crypts, the factor (1 - P(t)) is included in the ODE. The rate of change of P(t) is given as:

$$P'(t) = \left(\frac{1}{2}r_{\text{LOH}}n_{231} + \frac{1}{2}r_{\text{APC}}n_{131} + \frac{1}{2}r_{\text{LOH}}n_{321} + \frac{1}{2}r_{\text{TP53}}n_{311} + r_{\text{KRAS}}n_{330}\right)(1 - P(t)).$$

The equation for P(t) is derived by solving the differential equation P'(t), which describes the probability that at least one cancerous cryptis present by time:

$$P(t) = 1 - \exp\left(-\int \left(\frac{1}{2}r_{\text{LOH}}n_{231} + \frac{1}{2}r_{\text{APC}}n_{131} + \frac{1}{2}r_{\text{LOH}}n_{321} + \frac{1}{2}r_{\text{TP53}}n_{311} + r_{\text{KRAS}}n_{330}\right)dt\right).$$

This equation forms the foundation for analyzing P(t), representing the cumulative probability of at least one malignant crypt by time t. However, solving P(t) requires knowledge of the expected number of crypts for each genotype $n_{ijk}(t)$, which is derived from a system of differential equations.

The system of differential equations describes the time evolution of $n_{ijk}(t)$, the expected number of crypts with genotype (i, j, k). These equations track the accumulation of driver mutations, starting from the initial state where all crypts are wild-type. The system assumes linear first-order dynamics, with "outflow" terms neglected due to their minimal impact given the small mutation rates $(r_{\text{LOH}} \ll 1)[1]$.

The construction of $n'_{ijk}(t)$ involves two key processes: inflows and clonal expansion. Inflows account for transitions from precursor genotypes via mutation or loss of heterozygosity (LOH), where the inflow rate is proportional to the mutation or LOH rate and the number of crypts in the precursor genotype. The clonal expansion represents the growth advantages provided by mutations, such as $b_1 = 0.2/\text{year}$ for $APC^{-/-}$ crypts and $b_2 = 0.07/\text{year}$ for $KRAS^+$ crypts, which increase the population size of these genotypes over time.

This system provides the input for evaluating P(t), with solutions tailored to the mutation dynamics and growth effects in the simulated system.

2.3. Case Analysis. To understand the contribution of driver mutations to cancer risk, P(t) is analyzed under different cases of driver growth advantages. Each case isolates specific biological scenarios, offering insights into how APC and KRAS mutations influence cancer initiation.

Case 1. All Drivers Neutral

- Assumption: $b_1 = b_2 = b_{12} = 0$, meaning no crypt growth advantage is provided by APC or KRAS mutations.
- n_{000} is constant $(n_{000} \approx N)$, and other solutions $n_{ijk}(t)$ are simple polynomial functions of t, derived by integrating mutation and LOH rates.
- P(t) is computed as:

(1)
$$P(t) \approx N r_{\text{APC}} r_{\text{TP53}} r_{\text{KRAS}} r_{\text{LOH}}^2 \frac{t^5}{4}.$$

Case 2. APC Only Has Advantage

- Assumption: $b_1 > 0$, $b_2 = 0$, and $b_{12} = b_1$.
- The growth advantage of $APC^{-/-}$ crypts introduces exponential terms (e^{b_1t}) in n_{321} , n_{311} , and n_{330} , significantly impacting P(t).
- P(t) is computed as:

(2)
$$P(t) \approx \frac{3}{2} N r_{\text{APC}} r_{\text{TP53}} r_{\text{KRAS}} r_{\text{LOH}}^2 \frac{t^2}{b_1^3} e^{b_1 t}.$$

Case 3. APC And KRAS Have Advantage

- Assumption: $b_1 > 0$, $b_2 > 0$, and $b_{12} = b_1 + b_2$.
- $APC^{-/-}$ and $KRAS^+$ crypts jointly contribute to the growth advantage. Solutions for $n_{231}, n_{131}, n_{321}, n_{311}, n_{330}$ involve b_1, b_2 , and b_{12} , reflecting their combined impact.
- In addition to considering the proliferation rate of mutated crypts, we explicitly introduce the fixation multiplier in the case of double malignant mutations because the cumulative selective advantage and synergistic effects of double malignant mutations significantly influence the fixation probability and expansion capacity of mutated crypts[8]. We use $c = APC_{\text{mult}1} \times APC_{\text{mult}2} \times KRAS_{\text{mult}}$ as the correction to the formula to account for the increased chance of fixation of APC and KRAS mutated in a crypt.
- The P(t) is computed as:

(3)
$$P(t) \approx cNr_{\text{APC}}r_{\text{TP53}}r_{\text{KRAS}}r_{\text{LOH}}^2 \frac{t}{b_{12}^3(b_{12} - b_1)(b_{12} - b_2)}e^{b_{12}t}.$$

TAU-LEAPING SIMULATION

According to Paterson et al., two numerical simulation methods are introduced and implemented to predict the probability of CRC initiation with time: Gillespie's Stochastic Simulation Algorithm[5] and τ -leaping[6]. Considering the expensive computational cost of Gillespie algorithm, we choose to use PYTHON to reproduce the τ -leaping numerical simulation.

2.4. Gillespie's Stochastic Simulation Algorithm. The first method is Gillespie's Stochastic Simulation Algorithm (SSA). SSA provides an exact realization of the underlying master equation by simulating each event individually. CRC initiation probability modeling in Figure 2 of Paterson et al paper, where multiple driver genes (APC, TP53, KRAS) can mutate at low rates (on the order of 10^{-8} per base pair per year) amidst a high background rate of more common events such as crypt fission, the SSA's fidelity comes at the price of immense computational expense. Mathematically, the SSA proceeds by first computing the total event rate $\Gamma = \sum_i n_i \omega_i$, where n_i is the population at site i and ω_i is the sum of rates w_i^j for all reactions j at that site. The next event time Δt is then drawn from an exponential distribution with parameter Γ , $\Delta t \sim \text{Exp}(\Gamma^{-1})$,

and the specific event is selected with probability proportional to its rate. However, as Γ becomes very large—dominated by frequent fission events rather than rare mutations—the required number of steps grows prohibitively large. The flow of the Gillespie Algorithm is shown in Figure 3 below.

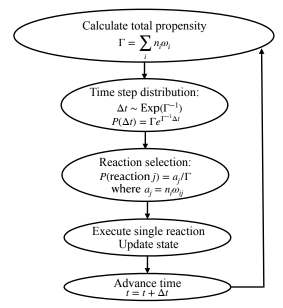


Figure 3. Gillespie Algorithm Scheme Diagram

2.5. τ -leaping. In contrast, the tau-leaping method trades some exactness for greater efficiency. Rather than sampling one event at a time, tau-leaping fixes a time step τ and draws the number of events of each type from appropriate distributions over that interval. For example, mutation events from state i to state j might be incremented by a Poisson-distributed random variable $\Delta n_{i \to j} \sim \text{Poisson}(n_i r_{i \to j} \tau)$. This allows the simulation to "leap" forward by time τ without enumerating each event. Moreover, for strongly proliferative processes like crypt fission, the paper describes using a negative binomial distribution to capture the "contagious" nature of division events more realistically, rather than relying solely on Poisson increments.

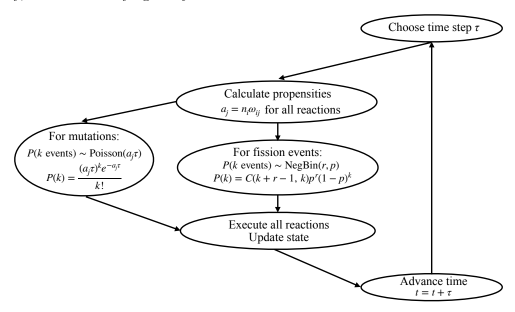


Figure 4. Tau-leaping method Scheme Diagram

2.6. Python Implementation. While the Python code (Appendix B.1.2) we developed attempted to implement τ -leaping foundational ideas, it did not fully replicate the recommended approach—particularly Step 3, which involves drawing the number of fission events from a negative binomial distribution $\Delta n_i \sim \text{NegBinomial}(n_i, p)$ with mean $n_i b_i \tau$. Step 3 involves representing branched paths of mutation and fission events, as illustrated in the tau-leaping scheme diagram in Figure 4 above. Instead, our code employed a mixture of Poisson, binomial, and negative binomial distributions in an ad-hoc manner, failing to precisely follow the protocol and thereby reducing the accuracy of the tau-leaping approximation. The comparison of our Python implementation and the original simulation in the paper is shown in Figure 5.

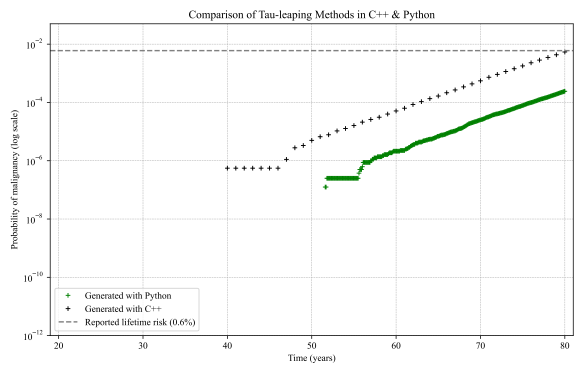


Figure 5. Different Simulation Approaches C++ (paper)/ Python (our implementation)

Additionally, due to Python's interpreted nature and the complexity of handling large populations with numerous events, the run time of our implementation was exceedingly long. Therefore, we resorted to using the C++ implementation provided by the authors of the paper(Appendix B.1.1), as compiled languages and carefully optimized code can handle the computational load more efficiently. By doing so, we achieved simulations that more closely matched the paper's results and could be directly compared with analytically inferred probabilities and rate matrices, thereby ensuring both computational tractability and improved fidelity to the intended tau-leaping formulation.

3. Reproduction of Results

After performing tau-leaping simulations four times for both advantageous scenarios and two times for the KRAS neutral scenario, we used the equations 1, 2 and 3 outlined in Section 2.3 to reproduce the figure from the original paper, as shown in Figure 6. The analytical equations for both advantageous and KRAS neutral cases align well with the two tau-leaping simulation lines, and the resulting plot closely resembles Figure 2, which is the corresponding figure in the original paper. The plotting code we used is from Appendix B.2.

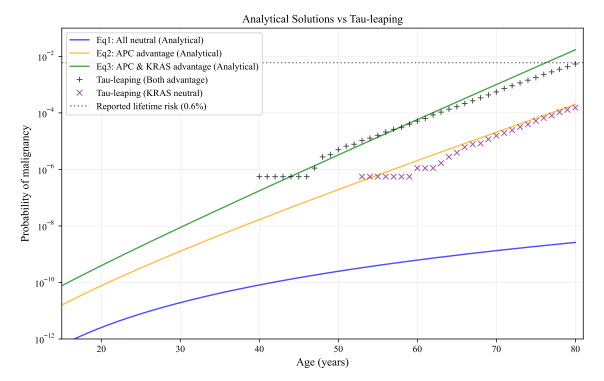


Figure 6. The analytical result with Tau-leaping

4. Novel Results

Building upon the biological framework established by Paterson et al., we develop a matrix-based approach to model colorectal cancer initiation, inspired by the *Matrix Models and Structured Population Dynamics* chapter in [4]. Paterson et al.'s work demonstrates that colonic crypts exhibit two fundamental behaviors: replication through fission and state transitions through genetic mutations. These characteristics naturally lend themselves to representation through a stage-based population model, where crypts can reproduce and transition between discrete genetic states. Our matrix formulation captures these dynamics through three key components: state vectors that track the population distribution across different genetic configurations, a transition matrix that governs the mutation processes between states, and a growth matrix that describes the selective proliferation of crypts with specific mutations. The setup of the linear model mainly involves three parts: The state variables that record the population compositions, the transition matrix that directs the mutation process, and the growth matrix that directs the cryptic expansion process.

- 4.1. **State Variables.** To model the progression of colorectal cancer, we define state variables as a combination of three key genes: APC, TP53, and KRAS. These state variables represent the mutational states of each gene and are denoted as a triplet (x, y, z), following the convention in Section 2.1, where:
 - $x \in \{0, 1, 2, 3\}$: Denotes the state of the APC gene.
 - x = 0: The APC gene is fully functional $(APC^{+/+})$.
 - x = 1: Loss of one APC allele $(APC^{+/})$.
 - x=2: Mutation in the remaining APC allele $(APC^{/+})$.
 - x=3: Complete functional loss of the APC gene (APC).
 - $y \in \{0, 1, 2, 3\}$: Denotes the state of the *TP53* gene.
 - y = 0: The TP53 gene is fully functional.
 - y = 1: Loss of one TP53 allele (TP53 LOH).

y = 2: Mutation in the remaining TP53 allele (TP53 mutation).

y = 3: Complete loss of TP53 functionality (TP53/).

• $z \in \{0,1\}$: Denotes the state of the KRAS gene.

z=0: The KRAS gene is in its wild-type form (KRAS⁻).

z = 1: The KRAS gene has undergone an activating mutation (KRAS⁺).

Note that we only consider states 0,1,2, and 3 for APC and TP53. This is because, according to the original paper[1], losing both chromosomes for a given crypt is too unlikely to happen that the probability do not affect the calculation results. Therefore, the combination of these variables results in a total of $4 \times 4 \times 2 = 32$ distinct states. Each state is represented as a unique triplet (x,y,z), corresponding to a specific mutation status for the three genes. For example, (0,0,0) represents the initial state where all three genes are fully functional, (0,1,0) indicates the loss of heterozygosity (LOH) in the TP53 gene, and (3,3,1) signifies the complete loss of APC and TP53 with an activating mutation in KRAS. A complete listing of state variables can be found in the Appendix A for further reference. The state vector is initialized as:

$$\mathbf{v}_0 = \begin{bmatrix} 10^8 \\ 0 \\ 0 \\ 0 \\ \vdots \end{bmatrix},$$

where 10^8 crypts are assumed to start in the state (0,0,0) in Section 2.1, and all other states have an initial cell population of 0. The state vector evolves over time according to a transition matrix, where each entry represents the mutation rates between different states, and the final probability of cancer can be extracted from specific states at later times.

4.2. **Transition Matrix.** The transition matrix T is a key component in modeling the progression of mutations in colorectal cancer. It represents mutation rates between different states, capturing the dynamics of transitions between combinations of APC, TP53, and KRAS gene mutations. Table 3 detailing the transition rules and their corresponding rates for mutations in the APC, TP53, and KRAS genes.

| Transition | Rate |
|------------------------------------|----------------------|
| $APC^{+/+} \rightarrow APC^{+/}$ | $r_{ m LOH}$ |
| $APC^{+/+} \rightarrow APC^{/+}$ | $r_{ m APC}$ |
| $APC^{+/} \to APC^{/}$ | $r_{\mathrm{APC}}/2$ |
| $APC^{/+} \rightarrow APC^{/}$ | $r_{\rm LOH}/2$ |
| $TP53^{+/+} \to TP53^{+/-}$ | $r_{ m LOH}$ |
| $TP53^{+/+} \rightarrow TP53^{/+}$ | r_{TP53} |
| $TP53^{+/} \rightarrow TP53^{/}$ | $r_{{ m TP53}}/2$ |
| $TP53^{/+} \rightarrow TP53^{/}$ | $r_{ m LOH}/2$ |
| $KRAS^- \to KRAS^+$ | $r_{ m KRAS}$ |

Table 3. Rules for Transition Rates

The 32×32 transition matrix T is constructed as follows:

- Each row corresponds to a **From State** (x, y, z), and each column corresponds to a **To State** (x', y', z').
- The entry a_{ij} in the *i*-th row and *j*-th column represents the **rate of transition** from state i (or (x, y, z)) to state j (or (x', y', z')). For example:

$$T[i,j] = r$$

- -i is the index corresponding to the **From State** (x, y, z). States are indexed sequentially based on their triplet representation (e.g., (0,0,0) is index 1, (0,0,1) is index 2, ..., (3,3,1) is index 32).
- -j is the index corresponding to the **To State** (x', y', z'), following the same indexing as i.
- -r is the mutation rate that governs the probability of transitioning from state i to state j. For example, r_{LOH} represents the rate of loss of heterozygosity, r_{APC} represents the mutation rate of APC, and so on.
- If there is no direct transition between (x, y, z) and (x', y', z'), T[i, j] = 0.

We can illustrate this with a few simple examples. Suppose Transition 1 is from state (0,0,0) to (0,1,0), which corresponds to a loss of heterozygosity (LOH) in the TP53 gene. The transition rate for this process is given by: $T[0,2] = r_{\text{LOH}}$. Similarly, Transition 2 is from state (0,0,0) to (0,2,0), which represents a point mutation in the TP53 gene. The transition rate for this process is given by: $T[0,4] = r_{\text{TP53}}$.

4.3. **Growth Rate Matrix.** The growth rate matrix represents the growth advantages conferred by mutations in specific genes. These growth rates account for the selective advantages of different mutational states in the progression of colorectal cancer. The growth rate matrix G is a 32×32 diagonal matrix, defined as follows:

$$G = \begin{bmatrix} b_{1,1} & 0 & 0 & \cdots & 0 \\ 0 & b_{2,2} & 0 & \cdots & 0 \\ 0 & 0 & b_{3,3} & \cdots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \cdots & b_{32,32} \end{bmatrix},$$

where $b_{i,i}$ represents the growth rate for state i. As we discussed in Section 4.1, each state is represented as a triplet: $(APC, TP53, KRAS) \rightarrow \text{state } (x, y, z)$, where $x, y \in \{0, 1, 2, 3\}, z \in \{0, 1\}$. The diagonal entries $b_{i,i}$ are determined by the following sources of growth rates:

- b_{APC} : Growth advantage when the APC gene mutation leads to complete functional loss $(x \ge 3)$,
- b_{KRAS} : Growth advantage when the KRAS gene undergoes an activating mutation (z=1),
- b_{BOTH} : Combined growth advantage when both APC and KRAS mutations occur simultaneously $(x \ge 3 \text{ and } z = 1)$.

Therefore, the growth rates are assigned to the diagonal entries of G based on the state variables:

$$b_{i,i} = \begin{cases} b_{\text{BOTH}}, & \text{if } x \ge 3 \text{ and } z = 1, \\ b_{\text{APC}}, & \text{if } x \ge 3 \text{ and } z = 0, \\ b_{\text{KRAS}}, & \text{if } x < 3 \text{ and } z = 1, \\ 0, & \text{otherwise.} \end{cases}$$

This matrix is used in conjunction with the transition matrix to simulate the evolution of cancer probabilities over time.

4.4. Computing the Probability. The Leslie-like matrix combines the transition matrix T and the growth rate matrix G to model the progression of mutations in colorectal cancer over time. This approach allows us to calculate the probability of malignancy at a specific age N. The final probability is computed using the following structure:

$$\mathbf{v}_N = (G \cdot T)^N \cdot \mathbf{v}_0,$$

where:

• G: The growth rate matrix (diagonal matrix).

- T: The transition matrix, representing mutation rates between states.
- \mathbf{v}_0 : The initial state vector, where all 10^8 crypts start in state (0,0,0).
- \mathbf{v}_N : The state vector after N years, containing probabilities for all 32 states.

The resulting state vector \mathbf{v}_N after N years contains the expected cell population for each of the 32 states. The probability of malignancy is the last entry in \mathbf{v}_N corresponding to the cancerous state (3,3,1), denoted as $\mathbf{v}_N[32]$.

5. Conclusion

5.1. **Matrix Approach.** Based on the new method introduced in Section 4, the matrix approach, combined with the tau-leaping simulation results, we were able to generate Figure 7 using the code provided in Appendix B.2. By comparing it with Figure 2, we observed that the matrix approach performs exceptionally well.

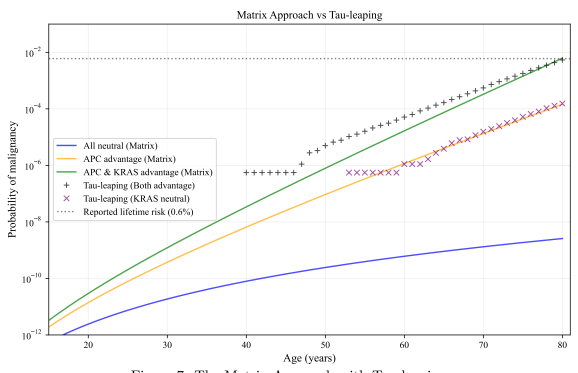


Figure 7. The Matrix Approach with Tau-leaping

- 5.2. Comparison of two approaches. To evaluate whether our matrix approach can truly replace the three equations introduced in Section 2.3, we plotted the probability curves from both the matrix and analytical methods together, as shown in Figure 8. Although the differences are very slight, they do exist. Therefore, we calculated the relative difference between the two methods, as illustrated in Figure 9.
- 5.3. Error Analysis. Based on the results shown in the figures, several key observations and potential explanations emerge regarding the differences between our matrix approach and the original analytical solutions.

The matrix approach consistently underestimates the probability of malignancy compared to the analytical solutions, as shown in Figure 8. This discrepancy is most pronounced in the "APC & KRAS advantage" case, where both mutations confer selective growth advantages. According to Paterson et al.'s supplementary materials[1], this difference may arise from their analytical approximations, which "assume that double mutations in tumor suppressor genes do not occur."

They note that this assumption is valid when "the rate of loss of heterozygosity (LOH) is much lower than the point mutation rate for each gene."

The relative difference plot (Figure 9) reveals intriguing patterns in the discrepancies between the methods. The difference is largest for the "APC & KRAS advantage" case and decreases over time in the "APC advantage only" scenario. This observation aligns with Paterson et al.'s claim that "the correction for double mutation in APC in our analytic approximation will be at most on the order of 2-3%." Our matrix method, which explicitly accounts for all possible transition pathways, including double mutations, demonstrates this effect.

Furthermore, Paterson et al. mention in their supplementary materials[1] that their analytical solutions neglect "outflow terms" in their system of equations. They state that "since the largest mutation rate, $r_{\rm LOH} = 1.36 \times 10^{-4} \, \rm yr^{-1}$, and the largest time $t = 80 \, \rm yr$, it will always be the case that $r_{\rm LOH} t < 1.1 \times 10^{-2} \ll 1$." However, our matrix approach inherently includes these outflow terms, which could contribute to the observed differences.

The decreasing relative difference over time in the neutral case suggests that both methods converge in their predictions over longer time periods, particularly when no selective advantages are present. This supports the validity of both approaches while underscoring the importance of accounting for the full complexity of mutation pathways in cancer evolution modeling.

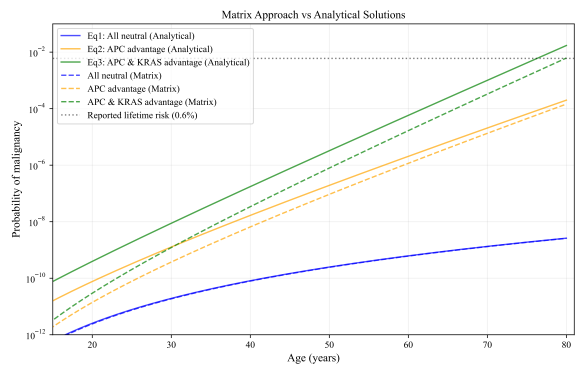


Figure 8. Matrix Approach vs Analytical Approach

5.4. Conclusion. The original paper developed a stochastic mathematical model to describe the initiation of CRC, we reproduced the result and did an alternative matrix approach based on the Leslie Model to explain and predict CRC with observed epidemiological data. The original paper integrates mathematical modeling with experimental data to advance understanding of CRC initiation and provides a foundation for assessing therapeutic and diagnostic strategies. By comparing the results, as the method converges over a long time while it still exists underestimating, we want to emphasize the complicated of the cancer and underscore that people should treat the probability of cancer initiation seriously.

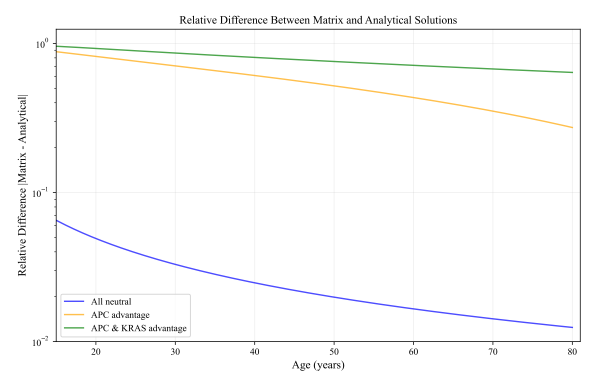


Figure 9. Relative Difference Between Matrix and Analytical Approach

The matrix approach we introduced in this paper provides several advantages. By encoding the complete state space and transition rules into matrix operations, we create a more streamlined computational framework that maintains the biological complexity of the system. Our comparative analysis shows that while the matrix method produces results that generally align with the original approach, there are notable differences that arise from the explicit inclusion of all possible transition pathways and the treatment of outflow terms.

5.5. Future Work. Looking forward, our matrix formulation opens up new avenues for analyzing cancer evolution using tools from linear algebra. Future work could explore the eigenvalue spectrum of the transition and growth matrices, potentially revealing characteristic timescales of cancer development and identifying critical pathways in the evolutionary process. The eigenvectors of these matrices might provide insights into the stable distributions of genetic states and the most likely paths to malignancy. Additionally, matrix perturbation theory could be applied to understand how small changes in mutation rates or growth advantages affect the overall system dynamics, with potential implications for therapeutic interventions. Furthermore, this matrix-based approach could be extended to model other types of cancer or more complex genetic networks. The framework's flexibility allows for the incorporation of additional mutations, different types of genetic alterations, or more sophisticated growth dynamics while maintaining computational traceability.

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APPENDIX A. STATE VARIABLES

| State | Description | | | |
|-----------|--|--|--|--|
| (0, 0, 0) | Normal crypt, no mutations. | | | |
| (0, 0, 1) | KRAS activated, others normal. | | | |
| (0, 1, 0) | TP53 lost one allele, others normal. | | | |
| (0, 1, 1) | TP53 lost one allele $+$ KRAS activated. | | | |
| (0, 2, 0) | TP53 mutated one allele, others normal. | | | |
| (0, 2, 1) | TP53 mutated one allele + KRAS activated. | | | |
| (0, 3, 0) | TP53 mutated+lost, others normal. | | | |
| (0, 3, 1) | TP53 mutated + lost + KRAS activated. | | | |
| (1, 0, 0) | APC lost one allele, others normal. | | | |
| (1, 0, 1) | APC lost one allele $+$ KRAS activated. | | | |
| (1, 1, 0) | APC lost one allele $+$ TP53 lost one allele. | | | |
| (1, 1, 1) | APC lost one allele $+$ TP53 lost one allele $+$ KRAS activated. | | | |
| (1, 2, 0) | APC lost one allele + TP53 mutated one allele. | | | |
| (1, 2, 1) | APC lost one allele + TP53 mutated one allele + KRAS activated. | | | |
| (1, 3, 0) | APC lost one allele + TP53 mutated+lost. | | | |
| (1, 3, 1) | APC lost one allele $+$ TP53 mutated $+$ lost $+$ KRAS activated. | | | |
| (2, 0, 0) | APC mutated one allele, others normal. | | | |
| (2, 0, 1) | APC mutated one allele + KRAS activated. | | | |
| (2, 1, 0) | APC mutated one allele $+$ TP53 lost one allele. | | | |
| (2, 1, 1) | APC mutated one allele + TP53 lost one allele + KRAS activated. | | | |
| (2, 2, 0) | APC mutated one allele + TP53 mutated one allele. | | | |
| (2, 2, 1) | APC mutated one allele + TP53 mutated one allele + KRAS activated. | | | |
| (2, 3, 0) | APC mutated one allele + TP53 mutated+lost. | | | |
| (2, 3, 1) | APC mutated one allele $+$ TP53 mutated $+$ lost $+$ KRAS activated. | | | |
| (3, 0, 0) | APC mutated+lost, others normal. | | | |
| (3, 0, 1) | APC mutated+lost + $KRAS $ activated. | | | |
| (3, 1, 0) | APC mutated+lost + $TP53$ lost one allele. | | | |
| (3, 1, 1) | APC mutated+lost + $TP53$ lost one allele + $KRAS$ activated. | | | |
| (3, 2, 0) | APC mutated+lost + TP53 mutated one allele. | | | |
| (3, 2, 1) | APC mutated+lost + TP53 mutated one allele + KRAS activated. | | | |
| (3, 3, 0) | APC mutated+lost + $TP53 $ mutated+lost. | | | |
| (3, 3, 1) | APC mutated+lost + TP53 mutated+lost + KRAS activated (Malignant state). | | | |

APPENDIX B. MAIN CODE

B.1. tau-leaping simulations.

B.1.1. Code for tau-leaping simulations in C++.

The C++ code used for tau-leaping simulations was provided by the original paper and can be found in the Data Availability section on the website.[1].

B.1.2. Code for tau-leaping simulations in Python.

```
import numpy as np
   import matplotlib.pyplot as plt
  from dataclasses import dataclass
   from typing import Dict, Set, Tuple
4
   import random
5
6
   @dataclass
   class SimParams:
8
       # Base mutation rate per base pair per year
9
       u: float = 1.25e-8
10
11
       # Number of driver positions in each gene
12
       n_APC: int = 604
                           # Number of APC driver positions
13
       n_TP53: int = 73
                            # Number of TP53 driver positions
14
15
       n_KRAS: int = 20
                            # Number of KRAS driver positions
16
       # Derived mutation rates per year
17
       r_APC: float = 604 * 1.25e-8
                                         # APC mutation rate
18
       r_TP53: float = 73 * 1.25e-8
                                        # TP53 mutation rate
19
       r_KRAS: float = 20 * 1.25e-8
                                       # KRAS mutation rate
20
21
       r_LOH: float = 1.36e-4
                                         # Rate of loss of heterozygosity
22
       # Division rates per year
23
       b_APC: float = 0.2
                              # Division rate for APC-/- crypts
24
       b_KRAS: float = 0.07
                               # Division rate for KRAS+ crypts
25
       b_BOTH: float = 0.27
                               # Division rate for APC-/-, KRAS+ crypts
26
                               # (b\_APC + b\_KRAS)
27
28
       # Time parameters
29
       dt: float = 0.01
                               # Time step size (tau)
30
       t_max: float = 80.0
                               # Maximum simulation time
31
32
       # Initial conditions
       N_crypts: int = 10**8 # Number of initial crypts
34
35
   class State:
36
       """Represents the genetic state of a crypt"""
37
       def __init__(self, APC: int, TP53: int, KRAS: int):
38
           # APC states:
39
           # 0: Wild type
40
           # 1: First hit LOH
41
           # 2: First hit mutation
42
           # 3: Second hit after LOH or mutation
43
           # 4: Double mutation
44
           self.APC = APC
```

```
46
           # TP53 states: similar to APC
47
           self.TP53 = TP53
48
49
           # KRAS states:
           # 0: Wild type
           # 1: Activated
           self.KRAS = KRAS
53
54
55
       def __hash__(self):
           return hash((self.APC, self.TP53, self.KRAS))
56
57
       def __eq__(self, other):
58
           return (self.APC, self.TP53, self.KRAS) ==
                   (other.APC, other.TP53, other.KRAS)
60
61
       def is_malignant(self):
62
           """Check if state represents a malignant crypt"""
63
           # Malignant states: (3,3,1), (3,4,1), (4,3,1), (4,4,1)
64
           return ((self.APC >= 3) and (self.TP53 >= 3) and (self.KRAS == 1))
65
66
   def get_transition_rate(source: State, target: State, params: SimParams)
67
      -> float:
       """Calculate transition rate between states based on Suppl. Mater."""
68
69
       # APC transitions
       if target.APC > source.APC:
70
           if source.APC == 0:
71
                # First hit can be LOH or mutation
72
                return params.r_LOH if target.APC == 1 else params.r_APC
73
           elif source.APC == 1:
74
75
                # After LOH, only mutation possible
                return params.r_APC / 2
76
           elif source.APC == 2:
77
                # After mutation, both LOH and mutation possible
78
                return params.r_LOH / 2 if target.APC == 3 else params.r_APC/2
79
80
81
       # TP53 transitions (similar to APC)
       if target.TP53 > source.TP53:
82
           if source.TP53 == 0:
83
                return params.r_LOH if target.TP53 == 1 else params.r_TP53
84
           elif source.TP53 == 1:
85
                return params.r_TP53 / 2
86
           elif source.TP53 == 2:
87
                return params.r_LOH / 2 if target.TP53==3 else params.r_TP53/2
88
89
       # KRAS activation
90
       if target.KRAS > source.KRAS:
91
92
           return params.r_KRAS
93
       return 0.0
94
95
96
   def get_division_rate(state: State, params: SimParams) -> float:
       """Get division rate for a given state based on Suppl. Mater."""
97
       if state.APC >= 3:
                                  # APC-/-
98
```

```
if state.KRAS == 1: # Also KRAS+
99
                return params.b_BOTH
100
            return params.b_APC
101
        elif state.KRAS == 1:
                                  # Only KRAS+
102
103
            return params.b_KRAS
        return 0.0
104
105
106
   def get_neighbors(state: State) -> Set[State]:
        """Get all possible next states from current state"""
107
108
        neighbors = set()
109
        # APC transitions
110
        if state.APC == 0:
111
112
            neighbors.add(State(1, state.TP53, state.KRAS)) # LOH
            neighbors.add(State(2, state.TP53, state.KRAS)) # Mutation
113
        elif state.APC == 1:
114
            neighbors.add(State(3, state.TP53, state.KRAS)) # Mutation after LOH
115
        elif state.APC == 2:
116
            neighbors.add(State(3, state.TP53, state.KRAS)) # LOH after
117
                mutation
            neighbors.add(State(4, state.TP53, state.KRAS)) # Second mutation
118
119
        # TP53 transitions
120
        if state.TP53 == 0:
121
122
            neighbors.add(State(state.APC, 1, state.KRAS))
                                                                # LOH
            neighbors.add(State(state.APC, 2, state.KRAS))
                                                               # Mutation
123
        elif state.TP53 == 1:
124
            neighbors.add(State(state.APC,3,state.KRAS)) # Mutation after LOH
125
        elif state.TP53 == 2:
126
            neighbors.add(State(state.APC, 3, state.KRAS))
127
                                                               # LOH after
            neighbors.add(State(state.APC, 4, state.KRAS)) # Second mutation
128
129
        # KRAS activation
130
        if state.KRAS == 0:
131
            neighbors.add(State(state.APC, state.TP53, 1))
132
133
        return neighbors
134
135
   def tau_leaping_simulation(params: SimParams, n_runs: int = 1000) -> np.
136
       ndarray:
        """Run multiple tau-leaping simulations following Suppl. Mater."""
137
        time_points = np.arange(0, params.t_max + params.dt, params.dt)
138
        malignant_counts = np.zeros(len(time_points))
139
140
        for run in range(n_runs):
141
            if run % 100 == 0:
142
143
                print(f"Run {run}/{n_runs}")
144
            # Initialize population dictionary
145
            population = {State(0,0,0): params.N_crypts}
146
147
            had_malignant = False
148
            for t_idx, t in enumerate(time_points):
149
```

```
if had_malignant:
150
                     malignant_counts[t_idx:] += 1
151
152
153
                 # Process each populated state
154
                 new_events = {}
155
                 for state, count in list(population.items()):
156
                     if count == 0:
157
                          continue
158
159
                     #Check for division events(negative binomial distribution)
160
                     division_rate = get_division_rate(state, params)
161
                     if division_rate > 0:
162
163
                          p = np.exp(-division_rate * params.dt)
                         n_divisions = np.random.negative_binomial(count, p)
164
                          if n_divisions > 0:
165
                              new_events[state] = new_events.get(state, 0) +
166
                                  n_divisions
167
168
                     # Check for transitions to neighbor states
                     for neighbor in get_neighbors(state):
169
                          rate = get_transition_rate(state, neighbor, params)
170
                          if rate > 0:
171
                              # Use Poisson approximation for small rates
172
173
                              if rate * params.dt < 0.01:</pre>
                                  n_transitions = np.random.poisson(rate *
174
                                      params.dt * count)
                              else:
175
                                  # Use binomial for larger rates
176
                                  p = 1 - np.exp(-rate * params.dt)
177
178
                                  n_transitions = np.random.binomial(count, p)
179
                              if n_transitions > 0:
180
                                  new_events[state] = new_events.get(state, 0) -
181
                                       n_{transitions}
                                  new_events[neighbor] = new_events.get(neighbor
182
                                      , 0) + n_transitions
183
                                  if neighbor.is_malignant():
184
                                       had_malignant = True
185
                                       break
186
187
                 # Update population
188
                 for state, delta in new_events.items():
189
                     population[state] = population.get(state, 0) + delta
190
                     if population[state] < 0: # Sanity check</pre>
191
                          population[state] = 0
192
193
                 if had_malignant:
194
                     malignant_counts[t_idx:] += 1
195
                     break
196
197
        return malignant_counts / n_runs
198
199
```

```
def plot_results(time_points: np.ndarray, probabilities: np.ndarray):
200
        """Plot the probability of malignancy over time"""
201
202
        plt.figure(figsize=(10, 6))
        plt.semilogy(time_points, probabilities, 'b-', label='Tau-leaping
203
           simulation')
        plt.xlabel('Age (years)')
204
        plt.ylabel('Probability of malignancy')
205
        plt.title('Probability of Colorectal Cancer Development')
206
        plt.grid(True)
207
        plt.legend()
208
        plt.show()
209
210
    from tqdm import tqdm
211
212
    def tau_leaping_simulation_with_progress(params: SimParams, n_runs: int =
213
       1000, checkpoints: list = None) -> np.ndarray:
        """Run multiple tau-leaping simulations with intermediate
214
        probability output and a progress bar"""
215
        time_points = np.arange(0, params.t_max + params.dt, params.dt)
216
217
        malignant_counts = np.zeros(len(time_points))
218
219
        if checkpoints is None:
            checkpoints = [] # Default: no checkpoints
220
221
222
        # Initialize tqdm progress bar
        with tqdm(total=n_runs, desc="Simulation Progress") as pbar:
223
            for run in range(n_runs):
224
                # Update progress bar
225
                pbar.update(1)
226
227
228
                # Checkpoints output
                if (run + 1) in checkpoints:
229
                     probabilities = malignant_counts / (run + 1)
230
                     print(f"Checkpoint {run + 1}/{n_runs}:")
231
                     print(f"Probabilities: {probabilities}")
232
233
234
                # Initialize population dictionary
                population = {State(0, 0, 0): params.N_crypts}
235
                had_malignant = False
236
237
                for t_idx, t in enumerate(time_points):
238
239
                     if had_malignant:
                         malignant_counts[t_idx:] += 1
240
241
                         break
242
                     # Process each populated state
243
                     new_events = {}
244
245
                     for state, count in list(population.items()):
                         if count == 0:
246
                              continue
247
248
249
                         # Check for division events (negative binomial distri.)
                         division_rate = get_division_rate(state, params)
250
                         if division_rate > 0:
251
```

```
p = np.exp(-division_rate * params.dt)
252
                              n_divisions = np.random.negative_binomial(count, p
253
                              if n_divisions > 0:
254
                                  new_events[state] = new_events.get(state, 0) +
255
                                       n_divisions
256
                         # Check for transitions to neighbor states
257
                         for neighbor in get_neighbors(state):
258
259
                              rate = get_transition_rate(state, neighbor, params)
                              if rate > 0:
260
                                  # Use Poisson approximation for small rates
261
                                  if rate * params.dt < 0.01:</pre>
262
263
                                       n_transitions = np.random.poisson(rate *
                                          params.dt * count)
                                  else:
264
                                       # Use binomial for larger rates
265
                                       p = 1 - np.exp(-rate * params.dt)
266
                                       n_transitions=np.random.binomial(count,p)
267
268
                                  if n_transitions > 0:
269
                                       new_events[state] = new_events.get(state,
270
                                          0) - n_transitions
                                       new_events[neighbor] = new_events.get(
271
                                          neighbor, 0) + n_transitions
272
                                       if neighbor.is_malignant():
273
                                           had_malignant = True
274
                                           break
275
276
277
                     # Update population
                     for state, delta in new_events.items():
278
                         population[state] = population.get(state, 0) + delta
279
                          if population[state] < 0: # Sanity check</pre>
280
                              population[state] = 0
281
282
283
                     if had_malignant:
                         malignant_counts[t_idx:] += 1
284
                         break
285
286
        return malignant_counts / n_runs
287
288
    # Run the simulation with a progress bar
289
    params = SimParams(
290
        dt = 0.1,
                            # Time step size
291
        t_max=80.0,
                            # Maximum time
292
        N_{crypts}=10**8
                            # Number of initial crypts
293
294
295
    n_runs = 8000000
296
    checkpoints = [500000, 1000000, 1500000, 2000000, 2500000, 3000000,
297
                    3500000, 4000000, 4500000, 5000000, 5500000, 6000000,
                    6500000, 7000000, 7500000, 8000000]
299
300
```

```
time_points = np.arange(0, params.t_max + params.dt, params.dt)
probabilities = tau_leaping_simulation_with_progress(params, n_runs=n_runs
    , checkpoints=checkpoints)

# Plot the results
plot_results(time_points, probabilities)
```

B.2. Code for reproducing plots.

```
import numpy as np
   import matplotlib.pyplot as plt
2
  import pandas as pd
  np.set_printoptions(threshold=np.inf)
   plt.rc('font', family='Times New Roman')
   # Define parameters
6
   class Parameters:
       def __init__(self):
8
           # Base mutation rate per base pair per year
9
           self.u = 1.25e-8
11
           # Number of driver positions in each gene
12
13
           self.n_APC = 604
           self.n_TP53 = 73
14
           self.n_KRAS = 20
15
16
           # Mutation rates per year
17
           self.r_APC = self.n_APC * self.u
18
           self.r_TP53 = self.n_TP53 * self.u
19
           self.r_KRAS = self.n_KRAS * self.u
20
           self.r_LOH = 1.36e-4
21
22
           # Growth rates per year
                                # APC-/- growth rate
           self.b_APC = 0.2
24
           self.b_KRAS = 0.07 # KRAS+ growth rate
25
           self.b_BOTH = 0.27 # Combined APC-/-, KRAS+ growth rate
26
           # Initial number of crypts
28
           self.N_crypts = 10**8
30
           # Correction factors
31
           self.c1 = 5.88 # APC fixation advantage
32
                            # KRAS fixation advantage
           self.c2 = 3.6
33
           self.c = self.c1 * self.c2 # Combined correction
```

B.2.1. Matrix Approach.

```
# Matrix Method
def create_transition_matrix(params):
    """Create 32x32 transition rate matrix"""
    n = 32
    T = np.zeros((n, n))

def state_to_idx(apc, tp53, kras):
    return apc * 8 + tp53 * 2 + kras
```

```
9
       # APC transitions
10
11
       for tp53 in range(4):
            for kras in range(2):
                # APC: (0) \rightarrow (1) or (2)
13
                T[state_to_idx(0, tp53, kras), state_to_idx(1, tp53, kras)] =
14
                   params.r_LOH
                T[state_to_idx(0, tp53, kras), state_to_idx(2, tp53, kras)] =
15
                   params.r_APC
16
                # APC: (1) \rightarrow (3)
17
                T[state_to_idx(1, tp53, kras), state_to_idx(3, tp53, kras)] =
18
                   params.r_APC/2
19
                # APC: (2) -> (3)
20
                T[state_to_idx(2, tp53, kras), state_to_idx(3, tp53, kras)] =
21
                   params.r_LOH/2
22
       # TP53 transitions
23
24
       for apc in range (4):
           for kras in range(2):
25
                # TP53: (0) -> (1) or (2)
26
                T[state_to_idx(apc, 0, kras), state_to_idx(apc, 1, kras)] =
                   params.r_LOH
28
                T[state_to_idx(apc, 0, kras), state_to_idx(apc, 2, kras)] =
                   params.r_TP53
29
                # TP53: (1) -> (3)
30
                T[state_to_idx(apc, 1, kras), state_to_idx(apc, 3, kras)] =
31
                   params.r_TP53/2
32
                # TP53: (2) -> (3)
33
                T[state_to_idx(apc, 2, kras), state_to_idx(apc, 3, kras)] =
34
                   params.r_LOH/2
       # KRAS transitions
36
       for apc in range(4):
37
            for tp53 in range(4):
38
                # KRAS: (0) -> (1)
39
                T[state_to_idx(apc, tp53, 0), state_to_idx(apc, tp53, 1)] =
40
                   params.r_KRAS
       return T
42
43
   def create_growth_matrix(params):
44
       """Create 32x32 diagonal growth rate matrix"""
45
       n = 32
46
47
       G = np.zeros((n, n))
48
       def state_to_idx(apc, tp53, kras):
49
50
           return apc * 8 + tp53 * 2 + kras
51
       # Fill diagonal with growth rates
       for apc in range(4):
53
```

```
for tp53 in range(4):
54
                for kras in range(2):
55
                     idx = state_to_idx(apc, tp53, kras)
56
                     if apc >= 3: # APC-/- states
58
                         if kras == 1: \# APC-/- + KRAS+
59
                             G[idx, idx] = params.b_BOTH
60
                         else: # Only APC-/-
61
                             G[idx, idx] = params.b_APC
62
63
                     elif kras == 1: # KRAS+ only
                         G[idx, idx] = params.b_KRAS
64
65
        return G
66
67
   def simulate_evolution(params, t_max=80, dt=0.1):
68
        """Simulate the evolution of crypts using matrix approach"""
69
        # Create matrices
70
       T = create_transition_matrix(params)
71
        G = create_growth_matrix(params)
72
73
        # # Output the transition matrix T and growth matrix G
74
        # print("Transition Matrix (T):")
75
        # print(T)
76
        # print("\nGrowth Matrix (G):")
77
78
        # print(G)
79
        # Time points
80
        times = np.arange(0, t_max + dt, dt)
81
        # Initialize state vector (all crypts in state 000)
83
84
        v = np.zeros(32)
        v[0] = params.N_crypts
85
86
        # Store results
87
        results = np.zeros(len(times))
88
        malignant_idx = 3*8 + 3*2 + 1 # Index for state (3,3,1)
89
90
        # Evolution
91
        for i, t in enumerate(times):
92
            # Store malignant probability
93
            # results[i] = v[malignant_idx] / params.N_crypts
94
            results[i] = v[malignant_idx]
95
96
            # Update state vector
97
            v = v + (np.matmul(v, T) + np.matmul(v, G)) * dt
98
99
        return times, results
100
```

B.2.2. Equation Approach.

```
# equation method
def neutral_solution(t, params):
    """Equation (1): All mutations are neutral"""
    P = (params.N_crypts * params.r_APC * params.r_TP53 *
```

```
params.r_KRAS * params.r_LOH**2 * t**5) / 4
5
       return P
6
   def APC_advantage_solution(t, params):
8
9
       """Equation (2): Only APC provides growth advantage"""
       P = (3 * params.N_crypts * params.r_APC * params.r_TP53 *
10
            params.r_KRAS * params.r_LOH**2 * t**2 *
11
            np.exp(params.b_APC * t)) / (2 * params.b_APC**3)
12
       return P * params.c1
13
14
   def both_advantage_solution(t, params):
15
       """Equation (3): Both APC and KRAS provide advantage"""
16
       b12 = params.b_BOTH
17
       b1 = params.b_APC
18
       b2 = params.b_KRAS
19
20
       P = (params.N_crypts * params.r_APC * params.r_TP53 *
21
            params.r_KRAS * params.r_LOH**2 * t * np.exp(b12 * t) *
22
            (1/(b12**3 * (b12 - b1)) + 1/(b12**3 * (b12 - b2)) +
23
             1/(b12**2 * (b12 - b2)**2)))
24
       return P * params.c
25
```

B.2.3. Tau-leaping.

```
# Consider tau-leaping data
  data1 = pd.read_csv('test1.1.csv', header=None)
   data2 = pd.read_csv('test2.1.csv', header=None)
3
   def process_csv(file_name):
5
       """Read and process a CSV file to calculate cancer probabilities."""
6
       df = file_name
8
       years = df[0]
9
       state_331 = df[38]
                            # (3,3,1) state
       state_341 = df[40] # (3,4,1) state
10
       state_{431} = df[48] # (4,3,1) state
11
       state_441 = df[50] # (4,4,1) state
12
       cancer_prob = state_331 + state_341 + state_431 + state_441
13
14
       print(cancer_prob)
       return years, cancer_prob
```